# **Equidistribution and Optimal Approximation Class\***

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1 Introduction 10

Local adaptive grid refinement is an important technique in finite element methods. 11 Its study can be traced back to the pioneering work [2] in one dimension. In recent 12 years, mathematicians start to prove the convergence and optimal complexity of the 13 adaptive procedure in multi-dimensions. Dörfler [11] first proved an error reduction 14 in the energy norm for the Poisson equation provided the initial mesh is fine enough. 15 Morin et al. [15, 16] extended the convergence result without the constrain of the initial mesh and they also reveal the importance of data oscillation. But results in 17 [11, 15, 16] only establish the qualitative convergence estimate by a proof of an error 18 reduction property. The number of elements generated by the adaptive algorithm 19 is not under control. A natural theoretical question is if a standard adaptive finite 20 element scheme would give an optimal asymptotic convergence rate in terms of the 21 number of elements. For linear finite element approximation to second order elliptic 22 boundary value problems in two dimensions, for example, an optimal asymptotic 23 error estimate would be something like

$$|u - u_N|_{1,\Omega} \le C(u)N^{-1/2},$$
 (1)

where  $u_N$  is a finite element approximation of the Poisson equation with homogenous 25 Dirichlet boundary condition based on an adaptive grid with at most N elements.

An important progress has been made by Binev et al. [7] concerning the asymp- 27 totic estimate (1). In their algorithm, an additional coarsening step is required to 28 achieve optimal complexity. However in practice the nearly optimal complexity 29

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is obtained without the coarsening step. Such theoretical gap is filled by Steven- 30 son [18] which shows that the practical refinement without a recurrent coarsening 31 will also generate finite element solution with quasi-optimal computational com- 32 plexity. But marking for oscillation and refinement with interior nodes assumptions 33 are still needed. Recently, [8] presented the most standard AFEM and proved a contraction property and quasi-optimal cardinality without any additional assumptions. 35 Their results show that if the solution  $u \in \mathcal{A}_s$ , where  $\mathcal{A}_s$  is the approximation class 36 space of rate s, then  $|u - u_N|_{1,\Omega} \le |u|_{\mathscr{A}_s} N^{-s}$ .

Another important theoretical and practical issue is to characterize the approx- 38 imation class  $\mathcal{A}_{1/2}$  using the smoothness of u. A near characterization of  $\mathcal{A}_{1/2}$  in 39 terms of Besov spaces  $B_{p,q}^k(\Omega)$  in two dimensions can be found in [6, 7] which 40 shows that  $u \in \mathcal{A}_{1/2}$  implies that  $u \in B_{1,1}^2(\Omega)$  and  $u \in B_{n,p}^2(\Omega)$  for p > 1 implies that 41  $u \in \mathcal{A}_{1/2}$ .

In this paper, we shall provide a sharper result: We prove that if  $u \in W^{2,L\log L}(\Omega)$ , i.e.,

$$\int_{\Omega} |D^2 u \log |D^2 u| |dx < \infty, \tag{45}$$

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then  $u\in\mathscr{A}_{1/2}.$  This is an improved result since, when p>1,  $B^2_{p,p}(\Omega)\subset W^{2,L\log L}(\Omega)$ from the Hölder inequality. With the regularity theory of elliptic equations, which 47 ensures  $u \in W^{2,L\log L}(\Omega)$ , we are led to conclude the following practical statement: 48 linear adaptive finite element approximation of second order elliptic equations in two 49 dimensions will achieve optimal rate of convergence.

Our contribution in this paper is further related with recent work on equidistribu- 51 tion and refinement strategies as follows:

- 1. The role of the equidistribution. In Sect. 2 we reveal that the equidistribution 53 principle can be severely violated but asymptoticly optimal error estimates can 54 still be maintained. The result (Theorem 1) is firstly presented in [9] and similar 55 idea can be also found in [8] around the same time.
- 2. The proof of the bound of the pollution of the local mesh refinement in the 57 completion is of its own interest. The estimate (Theorem 2) is a much sharper 58 constant comparing with existing results in [7]. The idea of the proof is borrowed 59 from [1] and the result is generalized from the uniform grids in [1] to compatible 60 divisible unstructured grids.

The rest of the paper is organized as follows. In Sect. 2 we explain the equidistribution principle for the case when the function to be approximated belongs to 63  $W^{2,1}(\Omega)$ . The advantage of our approach is that only standard approximation for the 64 interpolation operator are used, and approximation theory for Besov spaces is not 65 needed. In Sect. 3, we review the newest vertex bisection refinement strategy and 66 provide a sharp estimate for the number of triangle needed for the completion of the 67 mesh after an arbitrary marking and bisection refinement is performed. In Sect. 4, we 68 present a new approach for the local grid refinement based on the error estimate and 69 the equidistribution principle.

### 2 Error Estimate and Equidistribution Principle

We shall consider a simple elliptic boundary value problem

$$-\Delta u = f \text{ in } \Omega, \qquad u = 0 \text{ on } \partial \Omega, \tag{2}$$

where, for simplicity, we assume  $\Omega$  is a polygon and is partitioned by a shape regular 73conforming triangulation  $\mathscr{T}_N$  with N number of triangles. Let  $\mathscr{V}_N \subset H^1_0(\Omega)$  be the 74 corresponding continuous piecewise linear finite element space associated with this 75 triangulation  $\mathcal{T}_N$ .

A finite element approximation of the above problem is to find  $u_N \in \mathcal{V}_N$  such that 77

$$a(u_N, v_N) = (f, v_N) \quad \forall v_N \in \mathcal{V}_N, \tag{3}$$

where

$$a(u_N, v_N) = (f, v_N) \quad \forall v_N \in \mathscr{V}_N,$$

$$a(u, v) = \int_{\Omega} \nabla u \cdot \nabla v \, dx, \text{ and } (f, v) = \int_{\Omega} f v \, dx.$$
(3)

For this problem, it is well known that for a fixed finite element space  $\mathcal{V}_N$ 

$$|u - u_N|_{1,\Omega} = \inf_{\nu_N \in \mathscr{V}_N} |u - \nu_N|_{1,\Omega}. \tag{4}$$

We then present a  $H^1$  error estimate for linear triangular element interpolation in 81 two dimensions. We note that in two dimensions, the following two embeddings are 82 both valid:

$$W^{2,1}(\Omega) \subset W^{1,2}(\Omega) \equiv H^1(\Omega) \text{ and } W^{2,1}(\Omega) \subset C(\bar{\Omega}).$$
 (5)

Given  $u \in W^{2,1}(\Omega)$ , let  $u_I$  be the linear nodal value interpolant of u on  $\mathcal{T}_N$ . For any triangle  $\tau \in \mathcal{I}_N$ , thanks to (5) and the assumption that  $\tau$  is shape-regular, we have 85

$$|u - u_I|_{1,\tau} \lesssim |u|_{2,1,\tau}.$$

As a result,

$$|u-u_I|_{1,\Omega}^2 \lesssim \sum_{ au \in \mathscr{T}_N} |u|_{2,1, au}^2.$$

To minimize the error, we can try to minimize the right hand side. By Cauchy-Schwarz inequality, 90

$$|u|_{2,1,\Omega} = \sum_{\tau \in \mathscr{T}_N} |u|_{2,1,\tau} \leq (\sum_{\tau \in \mathscr{T}_N} 1)^{1/2} (\sum_{\tau \in \mathscr{T}_N} |u|_{2,1,\tau}^2)^{1/2} = N^{1/2} (\sum_{\tau \in \mathscr{T}_N} |u|_{2,1,\tau}^2)^{1/2}. \tag{91}$$

Thus, we have the following lower bound:

$$\left(\sum_{\tau \in \mathcal{T}_N} |u|_{2,1,\tau}^2\right)^{1/2} \ge N^{-1/2} |u|_{2,1,\Omega}. \tag{6}$$

The equality holds if and only if

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$$|u|_{2,1,\tau} = \frac{1}{N}|u|_{2,1,\Omega}.\tag{7}$$

The condition (7) is hard to be satisfied in general. But we can considerably relax 94 this condition to ensure the lower bound estimate (6) is still achieved asymptotically. 95 The relaxed condition is as follows:

$$|u|_{2,1,\tau} \le \kappa_{\tau,N} |u|_{2,1,\Omega}$$
 (8)

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and

$$\sum_{\tau \in \mathscr{T}_N} \kappa_{\tau,N}^2 \le c_1 N^{-1}. \tag{9}$$

When the above two inequalities hold, we have

$$|u - u_I|_{1,\Omega} \lesssim N^{-1/2} |u|_{2,1,\Omega}.$$

In summary, we have the following theorem.

**Theorem 1.** If  $\mathcal{T}_N$  is a triangulation with at most N triangles and satisfying (8) and (9), then

$$|u - u_N|_1 \le |u - u_I|_{1,\Omega} \lesssim N^{-1/2} |u|_{2,1,\Omega}.$$
 (10)

In the above analysis, we see how equidistribution principle plays an important 103 role in achieving asymptotically optimal accuracy for adaptive grids. We would like 104 to further elaborate that, in the current setting, equidistribution is indeed a sufficient 105 condition for optimal error, but by no means this has to be a necessary condition. 106 Namely the equidistribution principle can be severely violated but asymptoticly optimal error estimates can still be maintained. For example, the following mild violation 108 of this principle is certainly acceptable:

$$|u|_{2,1,\tau} \le \frac{c}{N} |u|_{2,1,\Omega}.$$
 (11)

In fact, this condition can be more significantly violated on a finitely many elements 110

$$|u|_{2,1,\tau} \le \frac{c}{\sqrt{N}} |u|_{2,1,\Omega}.$$
 (12)

It is easy to see if a bounded number of elements satisfy (12) and the rest satisfy (11), 112 the estimate (9) is satisfied and hence the optimal error estimate (10) is still valid.

As we can see that the condition (12) is a very serious violation of equidistribution principle, nevertheless, as long as such violations do not occur on too many 115 elements, asymptotically optimal error estimates are still valid. This simple obser- 116 vation is important from both theoretical and practical points of view. The marking 117 strategy proposed by Dörfler [11] may also be interpreted in this way in its relation- 118 ship with equidistribution principle. In [5], they propose to use certain penalty in 119 using equidistribution principle. Such a modification certainly has similar spirit.

We shall discuss how to generate a mesh  $\mathcal{T}_N$  to satisfy (8) and (9) in the next two 121 sections. To this end, we shall introduce the local refinement method: newest vertex 122 bisection, in the next section.

#### 3 Newest Vertex Bisection

In this section we shall give a brief introduction of the newest vertex bisection and 125 mainly concern the number of elements added by the completion process. We refer 126 to [14, 19] and [7] for detailed description of the newest vertex bisection refinement 127 procedure.

Given an initial shape regular triangulation  $\mathcal{T}_0$  of  $\Omega$ , it is possible to assign 129 to each  $\tau \in \mathscr{T}_0$  exactly one vertex called *the newest vertex*. The opposite edge of the newest vertex is called *refinement edge*. The rule of the newest vertex bisection 131 includes:

- 1. A triangle is divided to two new children triangles by connecting the newest 133 vertex to the midpoint of the refinement edge;
- 2. The new vertex created at a midpoint of a refinement edge is assigned to be the 135 newest vertex of the children. 136

It is easy to verify that all the descendants of an original triangle fall into four similar- 137 ity classes (see Fig. 1) and hence the angles are bounded away from 0 and  $\pi$  and all 138 triangulations refined from  $\mathcal{T}_0$  using newest vertex bisection forms a shape regular 139 class of triangulations.









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Fig. 1. Four similarity classes of triangles generated by the newest vertex bisection

The triangulation obtained by the newest vertex might have hanging nodes. We 141 have to make additional subdivisions to eliminate the hanging nodes, i.e., complete 142 the new partition. The completion should also follow the bisection rules. We shall 143 consider more combinatory properties of the completion.

Let the triangles of the initial triangulation be assigned generation 0. We refer to 145 the two triangles obtained by splitting a triangle  $\tau$  in two sub-triangles by the newest 146 vertex procedure as being the children of  $\tau$ . For  $i = 1, 2, \dots$ , we define the generation 147 of the children of  $\tau$  to be i if the parent  $\tau$  has the generation i-1. It can be shown 148 that the completion will terminate in finite steps, due to the fact that the completion 149 process will not create new generations of triangles (see [3, 13]).

We ask more than the termination of the completion process. That is we want 151 to control the number of elements refined due to the completion. To this end, we 152 have to carefully assign the newest vertexs for the initial partition  $\mathcal{T}_0$ . A triangle is 153 called compatible divisible if its refinement edge is either the refinement edge of the triangle that shares that edge or an edge on the boundary. A triangulation  $\mathcal{T}$  is called 155 compatible divisible or compatible labled if every triangle is compatible divisible. 156 See Fig. 2 for an example of such compatible initial labeling.

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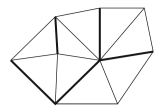


Fig. 2. A conforming divisible labeling of the initial triangulation where edges in bold case are refinement edges

It is obvious that the completion for a compatible triangulation is terminated 158 in one step. Mitchell [13] proves that for any conforming triangulation  $\mathcal{T}$ , there 159 exist a compatible label scheme. Biedl et al. [4] present an O(N) algorithm to find a 160 compatible labeling for a triangulation  $\mathcal{T}$  with N elements.

Let  $\mathscr{T}_0$  be a compatible triangulation and let  $\mathscr{T}_{\underline{1}}$  be a triangulation obtained by the newest vertex bisection by performing  $m_0$  bisections starting from  $\mathcal{T}_0$ . Denote by  $\mathcal{M}_0$  the set of all  $m_0$  marked and split triangles. Note that not all the triangles of  $\mathcal{M}_0$  164 have to be in  $\mathcal{T}_0$ . Let  $\mathcal{T}_1$  be the (minimal) conforming refinement of  $\mathcal{T}_{\frac{1}{2}}$  and denote 165 by  $n_k$  the number of triangles of  $\mathcal{T}_k$ , k = 0, 1 (Fig. 3).

(b)

**Fig. 3.** Marking, splitting, and completing. (a)  $\mathscr{T}_0$ . (b)  $\mathscr{T}_{\frac{1}{2}}$ . (c)  $\mathscr{T}_1$ 

**Theorem 2.** Let  $\mathcal{T}_0$  be a compatible triangulation and  $\mathcal{T}_1$  be obtained as above. 167 Then there exists a constant C only depending on the minimal angle of  $\mathscr{T}_0$  such that

$$n_1 \le n_0 + (C+1) m_0. \tag{13}$$

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*Remark 1.* It is a temptation to repeat the Theorem 2 to conclude: for j = 1, 2, ..., p-1691, we have that  $\mathcal{I}_{j+1}$  is obtained from  $\mathcal{I}_j$ , by  $m_j$  markings and then minimal completion, then

$$n_p \le n_0 + (C+1) (m_0 + m_1 + \dots + m_{p-1}).$$
 (14)

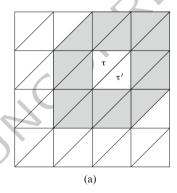
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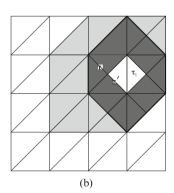
Unfortunately this argument does not work since  $\mathscr{T}_1$  may not be compatible divisible 172 anymore. The inequality (14) still holds but the proof is much involved; See Theorem 173 2.4 in [7]. The bound (13) can be derived from that theorem; See Lemma 2.5 in [7]. However, careful tracing the argument in [7] would give a huge constant in (14) in the 175 magnitude of 10,000. We shall give another more direct and simpler proof based on 176 an improved technique in [1]. The constant in our proof is much smaller and usually 177 below 100. Note that numerically in the average case of the constant is around 4 and 178 in the worst case is around 14; see [1].

Let us introduce notation for uniform bisection by setting  $\overline{\mathcal{T}}_k$  as the triangulation 180 obtained by bisecting each triangle in  $\mathcal{T}_0$  completely up to the k-th generation. The assumption:  $\mathcal{T}_0$  is compatible divisible implies that  $\overline{\mathcal{T}}_k$  is conforming and compatible divisible for all k > 1. Note that this may not hold if the initial labeling is not 183 compatible divisible.

For a triangle  $\tau$ , we define a neighbor of  $\tau$  as another triangle sharing a common 185 edges of  $\tau$ . By the definition, a triangle has at most three neighbors. Among them, for 186  $au \in \overline{\mathcal{T}}_k$ , we define the *refinement neighbor* of au as the triangle  $au' \in \overline{\mathcal{T}}_k$  such that au 187 and  $\tau'$  use the same edge as their refinement edges. We allow  $\tau' = \emptyset$  for  $\tau$  touching 188 the boundary. We define the *barrier* of  $\tau$  as all triangles in  $\overline{\mathcal{F}}_{\varrho(\tau)}$  which intersect 189  $\tau \cup \tau'$  and denoted by  $B(\tau)$ , i.e.,

$$B( au)=\{\hat{ au}\in\overline{\mathscr{T}}_{g( au)},\hat{ au}\cap( au\cup au')
eq\varnothing\}.$$





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Fig. 4. Barrier of a safe triangle. (a) Barrier 1. (b) Barrier 2

**Definition 1.** We say that  $\tau$  is a safe triangle if none of the barrier elements of  $\tau$  is marked in going from  $\mathcal{T}_0$  to  $\mathcal{T}_1$ , namely  $\hat{\tau} \notin \mathcal{M}_0$  for any  $\hat{\tau} \in B(\tau)$ . 193

The following lemma will justify the name of safe triangles. They are triangles 194 that not touched going from  $\mathcal{T}_0$  to  $\mathcal{T}_1$ . 195

**Lemma 1.** Any safe triangle  $\tau$  in  $\mathcal{T}_0$  or born in the marking and completion process of going from  $\mathcal{T}_0$  to  $\mathcal{T}_1$  will never be bisected during the completion process.

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*Proof.* We shall prove it by the induction over the generation of  $\tau$ . Suppose  $g(\tau) = 198$  $\max_{ ilde{ au}\in\mathscr{T}_{\underline{1}}}g( ilde{ au})$  and au is safe. Then au will not be bisected during the completion since the completion will not increase the maximal generation. 200

Assume that our statement holds for all safe triangles of generation p+1. We will 201 show that the statement also holds for a safe triangle with generation p. Note that to 202 trigger the bisection of  $\tau$ , one has to refine one of the two neighbors of  $\tau$  (which 203 do not share the refinement edge with  $\tau$ ) twice or two such neighbors of  $\tau'$  twice 204 (since  $\tau$  and  $\tau'$  share the refinement edge). Without loss of generality, let us say that 205 one of the neighbor  $\tau'$  is bisected once in the completion process. Then it produces 206 a children triangle  $\tau_1$  of generation p+1 which has a common edge with  $\tau'$ . It is 207 important to note that  $B(\tau_1) \subset B(\tau)$  and thus  $\tau_1$  is safe; See Fig. 4 for an illustration. 208 By the inductive hypothesis  $\tau_1$  will never be bisected anymore during the completion 209 process. Consequently,  $\tau$  will never be bisected during the completion process.

Now we are in the position to prove Theorem 2.

*Proof.* (of Theorem 2) We denote by  $\mathcal{M}_{\frac{1}{2}}$  as the set of all triangles  $\tau$  which are split 212 in the completion process of going from  $\mathcal{T}_{\frac{1}{2}}$  to  $\mathcal{T}_1$ . Let us choose a triangle  $\tau \in \mathcal{M}_{\frac{1}{2}}$ . 213 Since  $\tau$  is split in the completion process, by the above Lemma,  $\tau$  is not safe. It 214 implies that there should exist a same-generation triangle  $F(\tau)$  in  $B(\tau)$  such that 215  $F(\tau) \in \mathcal{M}_0$ . In this way, we defined a map from  $F: \mathcal{M}_{\frac{1}{2}} \to \mathcal{M}_0$ .

Note that F is not necessary a one-to-one map, but a triangle  $\tau \in \mathcal{M}_0$  could be 217 in only finite number of barriers, due to the space limitation of the same-generation 218 assumption. Given a triangle  $\tau$ , we define the first ring of  $\tau$  as all triangles intersect 219  $\tau$  and the second ring of  $\tau$  as the union of first rings of triangles in the first ring of  $\tau$ . 220 Then  $\tau$  can be only in the barrier of triangles in its second ring and thus the number 221 is bounded by the maximum number of triangles in the second ring of a triangle, say 222 C, which is usually below 100. Thus any triangle in  $\mathcal{M}_0$  is the image of at most C 223 triangles from  $\mathcal{M}_{\frac{1}{2}}$ . This leads to the fact that the number of splittings needed for 224 completion can be bounded by  $Cm_0$ . Since any splitting in the completion process 225 adds one more triangle towards the completed mesh  $\mathcal{I}_1$ , we have proved (13).

## 4 Local Grid Refinement Algorithm

In this section we shall propose a new approach for the local grid refinement based 228 on the error estimate and the equidistribution principle. We will use newest vertex 229 bisection to refine the grid and use  $|u|_{2,1,\tau}$  as an error indicator. With a little bit higher 230 regularity requirement of u, we are able to prove the effectiveness of our algorithm. 231 Namely, it will end with an optimal asymptotic error estimate similar to (1).

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#### 4.1 Local Refinement Strategy

We will illustrate a way to find a nearly optimal grid for the solution of (2). We will 234 use the newest vertex bisection refinement procedure with the marking strategy given 235 by (11). For the later analysis, we will have to assume that the solution u is in  $W^{2,1}$ and that the Hardy-Littlewood maximal function of  $D^2u$  is in  $L^1(\Omega)$ . Due to a result 237 of [17], this is equivalently  $D^2u \in L\log L(\Omega)$ . Such further assumption holds if for 238 example  $u \in W^{2,p}$  for some p > 1.

The maximal function of an integrable function f on  $\Omega$  is defined by

$$\widetilde{M}f(x) = \sup \frac{1}{|Q|} \int_{Q} |f(y)| \ dy,$$

where the supremum is taken over all square domains contained in  $\Omega$  and containing 241 х. 242

For a triangulation obtained by the newest vertices bisection from  $\mathcal{I}_0$ . The similarity classes are in fact completely represented by the children and grandchildren of 244 all triangles from  $\mathcal{T}_0$ . Let us denote by  $\mathcal{C}_0$  the following family of triangles: 245

> $\mathscr{C}_0 = \{ \tau | \ \tau \text{ is a triangle contained in } \Omega \text{ and is similar with }$ a child or grandchild of a triangle from  $\mathcal{T}_0$

We define another maximal function

$$Mf(x) = \sup \frac{1}{|\tau|} \int_{\tau} |f(y)| \ dy,$$

where the supremum is taken over all triangles  $\tau \in \mathscr{C}_0$  containing x. Then it is easy 247 to show that M and M are equivalent in the sense that 248

$$c_1\widetilde{M}f(x) \le Mf(x) \le c_2\widetilde{M}f(x), \quad \forall x \in \Omega$$

with  $c_1$  and  $c_2$  independent of x. Thus, for theoretical purposes, the two operators M 249 and M are interchangeable. 250

The following result concerns the number of the new triangles added in the re- 251 finement procedure. The main idea of the proof for the 1-D case was showed to the 252 authors by DeVore and can be found in [10]. 253

**Theorem 3.** Let f be an integrable function on  $\Omega$  such that  $Mf \in L^1(\Omega)$ , and let 254  $\varepsilon > 0$  be given. Assume that the newest vertex bisection refinement procedure is 255 applied to an compatible initial triangulation  $\mathcal{T}_0$  with  $n_0$  triangles. Let the marking 256 strategy be given by: a triangle  $\tau$  is marked if 257

$$\int_{\tau} |f(x)| \ dx > \varepsilon.$$

Denote by  $\mathcal{M}_0$  the set of all marked and split triangles. Then, the marking and re- 258 finement procedure will terminate in finite steps and we have 259

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$$n_0 + m_0 < \frac{2}{\varepsilon} \int_{\Omega} Mf(x) \ dx, \tag{15}$$

where  $m_0$  is the number of elements of  $\mathcal{M}_0$ . Assume that  $\mathcal{T}_{\frac{1}{2}}$  is the triangulation 260 obtained from  $\mathcal{T}_0$  after the  $m_0$  bisections. Let  $\mathcal{T}_1$  be the (minimal) conforming refinement of  $\mathcal{T}_{\frac{1}{2}}$  and denote by  $n_1$  the number of triangles of  $\mathcal{T}_1$ . Then,

$$n_1 \le \frac{C_1}{\varepsilon} \int_{\Omega} |Mf(x)| \ dx,\tag{16}$$

with a constant  $C_1$  independent of the function f and the number  $\varepsilon$ . More precisely, 263  $C_1 = 2(C+1)$ , with C the constant of Theorem 2.

*Proof.* Since  $\lim_{|\tau| \to 0} \int_{\tau} |f(x)| dx = 0$  and the areas of new triangles are exponentially decreased, the refinement procedure will terminate in finite steps.

We can assume without loss of generality that each triangle in  $\mathcal{T}_{\frac{1}{2}}$  is not a triangle 267 in  $\mathcal{T}_0$ . Now, let  $\tau \in \mathcal{T}_{\frac{1}{2}}$  and let  $\tilde{\tau}$  be its parent. Then  $\tilde{\tau} \in \mathcal{M}_0$ . (Recall that  $\mathcal{M}_0$  is 268 the collection of marked triangles in the refinement procedure.) By our refinement 269 strategy

 $\int_{\tilde{\tau}} |f(x)| \ dx > \varepsilon,$ 

Thus,

$$Mf(x) > \frac{1}{|\tilde{\tau}|} \int_{\tilde{\tau}} |f(y)| \, dy > \frac{\varepsilon}{|\tilde{\tau}|}, \quad \forall x \in \tau.$$

Integrating the above inequality on  $\tau$  we have,

$$\int_{\tau} Mf(x) \ dx > \frac{\varepsilon}{2}.\tag{17}$$

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Here we use fact  $|\tilde{\tau}| = 2|\tau|$ . If we sum up (17) over all  $n_0 + m_0$  triangles  $\tau \in \mathscr{T}_{\frac{1}{2}}$  we 273 obtain (15).

By using Theorem 2 we have that

$$n_1 < n_0 + m_0 + C m_0 < (C+1) (n_0 + m_0).$$
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The estimate (16) follows now as a direct consequence of (15) and the above inequality.

An application of Theorem 1 and the estimate (16) for  $f = D^2u$  and  $\varepsilon = 1/N$ , 279 leads to the proof of the existence of a nearly optimal grid. Starting from a coarse 280 grid  $\mathscr{T}_0$ , we define the approximation class  $\mathscr{A}_{1/2}$  as

$$\mathscr{A}_{1/2} = \{u \in H^1_0(\Omega): |u|_{\mathscr{A}_{1/2}} := \sup_{N \geq \#\mathscr{T}_0} N^{-1/2} \inf_{\#\mathscr{T} \leq N} \inf_{v_h \in V(\mathscr{T})} |u - v_h|_1 < \infty \}. \tag{282}$$

**Corollary 1.** If 
$$u \in W^{2,L\log L}(\Omega)$$
, then  $u \in \mathcal{A}_{1/2}$ .

Remark 2. The  $(L\log L)$  norm is needed only for proving the success of the algorithm 284 but is not effectively needed for the implementation of the algorithm. If we can find 285 good approximations or upper bound for  $\int_{\tau} D^2 u dx$  on triangles using e.g., gradient 286 and Hessian recovery methods (from the discrete Galerkin approximation of u) or 287 using regularity result in [12], then the ideas presented in this paper can lead to new 288 and optimal adaptive methods. 289

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# Some Recent Tools and a BDDC Algorithm for 3D **Problems in H(curl)**

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Summary. We present some recent domain decomposition tools and a BDDC algorithm for 13 3D problems in the space  $H(\text{curl}; \Omega)$ . Of primary interest is a face decomposition lemma 14 which allows us to obtain improved estimates for a BDDC algorithm under less restrictive assumptions than have appeared previously in the literature. Numerical results are also presented 16 to confirm the theory and to provide additional insights.

1 Introduction 18

We investigate a BDDC algorithm for three-dimensional (3D) problems in the space 19  $H_0(\text{curl};\Omega)$ . The subject problem is to obtain edge finite element approximations of 20 the variational problem: Find  $\mathbf{u} \in H_0(\text{curl}; \Omega)$  such that

$$a_{\Omega}(\boldsymbol{u},\boldsymbol{v}) = (\boldsymbol{f},\boldsymbol{v})_{\Omega} \quad \forall \boldsymbol{v} \in H_0(\operatorname{curl};\Omega),$$

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where 23

$$a_{\Omega}(\boldsymbol{u},\boldsymbol{v}) := \int_{\Omega} \left[ (\boldsymbol{\alpha} \nabla \times \boldsymbol{u} \cdot \nabla \times \boldsymbol{v}) + (\boldsymbol{\beta} \boldsymbol{u} \cdot \boldsymbol{v}) \right] dx, \quad (\boldsymbol{f},\boldsymbol{v})_{\Omega} = \int_{\Omega} \boldsymbol{f} \cdot \boldsymbol{v} dx.$$
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The norm of  $\mathbf{u} \in H(\text{curl}; \Omega)$ , for a domain with diameter 1, is given by  $a_{\Omega}(\mathbf{u}, \mathbf{u})^{1/2}$ with  $\alpha = 1$  and  $\beta = 1$ ; the elements of  $H_0(\text{curl})$  have vanishing tangential compo- 26 nents on  $\partial \Omega$ . We could equally well consider cases where this boundary condition 27 is imposed only on one or several subdomain faces which form part of  $\partial \Omega$ . We will 28 assume that  $\alpha \geq 0$  and  $\beta > 0$  are constant in each of the subdomains  $\Omega_1, \ldots, \Omega_N$ . 29 Our results could be presented in a form which accommodates properties which are 30 not constant or isotropic in each subdomain, but we avoid this generalization for 31 purposes of clarity. 32

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In the pioneering work of [12], two different cases were analyzed for FETI-DP 33 algorithms:

$$\alpha_i = \alpha$$
 for  $i = 1, \dots, N$ 

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The condition number bound reported for the preconditioned operator is

$$\kappa \le C \max_{i} (1 + H_i^2 \beta_i / \alpha) (1 + \log(H/h))^4, \tag{1}$$

where 
$$H/h := \max_i H_i/h_i$$
.

Case 2: 
$$\beta_i = \beta \quad \text{for} \quad i = 1....N$$

for which the reported condition number bound is

$$\kappa \le C \max_{i} (1 + H_i^2 \beta / \alpha_i) (1 + \log(H/h))^4. \tag{2}$$

We address the following basic questions regarding [12] in this study.

- 1. Is is possible to remove the assumption of  $\alpha_i = \alpha$  or  $\beta_i = \beta$  for all *i*?
- 2. Is it possible to remove the factor of  $H_i^2 \beta_i / \alpha_i$  from the estimates?
- 3. Is is possible to reduce the logarithmic factor from four powers to two powers as 45 is typical of other iterative substructuring algorithms? 46
- 4. Do FETI-DP or BDDC algorithms for 3D H(curl) problems have certain complications not present for problems with just a single parameter? 48

We find in the following sections that the answers are yes to all four questions. However, due to page limitations, we only consider here the relatively rich coarse space 50 of Algorithm C of [12]. We remark that the analysis of 3D H(curl) problems with 51 material property jumps between subdomains is quite limited in the literature. A 52 comprehensive treatment of problems in 2D can be found in [3]. A different iterative 53 substructuring algorithm for 3D problems is given in [6], but the authors were un- 54 able to conclude whether their condition number bound was independent of material 55 property jumps. A related study on substructuring preconditioners can also be found 56 in [7].

2 Tools 58

We assume that  $\Omega$  is decomposed into N non-overlapping subdomains,  $\Omega_1, \ldots, \Omega_N$ , 59 each the union of elements of the triangulation of  $\Omega$ . We denote by  $H_i$  the diameter 60 of  $\Omega_i$ . The interface of the domain decomposition is given by 61

$$\Gamma := \left(igcup_{i=1}^N \partial \Omega_i
ight) ackslash \partial \Omega,$$
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and the contribution to  $\Gamma$  from  $\partial \Omega_i$  by  $\Gamma_i := \partial \Omega_i \setminus \partial \Omega$ . These sets are unions of 63 subdomain faces, edges, and vertices. For simplicity, we assume that each subdomain 64 is a shape-regular and convex tetrahedron or hexahedron with planar faces. 65

We assume a shape-regular triangulation  $\mathcal{T}_{h_i}$  of each  $\Omega_i$  with nodes matching 66 across the interfaces. The smallest element diameter of  $\mathcal{T}_{h_i}$  is denoted by  $h_i$ . Associated with the triangulation  $\mathcal{T}_{h_i}$  are the two finite element spaces  $W_{\mathrm{grad}}^{h_i} \subset H(\mathrm{grad},\Omega_i)$  68 and  $W_{\mathrm{curl}}^{h_i} \subset H(\mathrm{curl},\Omega_i)$  based on continuous, piecewise linear, tetrahedral nodal elements and linear, tetrahedral edge (Nédeléc) elements, respectively. We could equally 70 well develop our algorithms and theory for low order hexahedral elements.

The energy of a vector function  $\mathbf{u} \in W_{\mathrm{curl}}^{h_i}$  for subdomain  $\Omega_i$  is defined as

$$E_i(\boldsymbol{u}) := \alpha_i (\nabla \times \boldsymbol{u}, \nabla \times \boldsymbol{u})_{\Omega_i} + \beta_i (\boldsymbol{u}, \boldsymbol{u})_{\Omega_i}, \tag{3}$$

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where  $\alpha_i$  and  $\beta_i$  are assumed constant in  $\Omega_i$ .

Let  $N_e \in W_{\text{curl}}^{h_i}$  and  $t_e$  denote the finite element shape function and unit tangent 74 vector, respectively, for an edge e of  $\mathcal{T}_{h_i}$ . We assume that  $N_e$  is scaled such that 75  $N_e \cdot t_e = 1$  along e. The edge finite element interpolant of a sufficiently smooth vector 76 function  $u \in H(\text{curl}, \Omega_i)$  is then defined as

$$\Pi^{h_i}(\boldsymbol{u}) := \sum_{e \in \mathcal{M}_{\tilde{\Omega}_i}} u_e \boldsymbol{N}_e, \quad u_e := (1/|e|) \int_e \boldsymbol{u} \cdot \boldsymbol{t}_e \, ds, \tag{4}$$

where  $\mathcal{M}_{\bar{\Omega}_i}$  is the set of edges of  $\mathcal{T}_{h_i}$ , and |e| is the length of e. We will also make use 78 of other sets of edges of  $\mathcal{T}_{h_i}$ . Namely,  $\mathcal{M}_{\partial\Omega_i}$ ,  $\mathcal{M}_{\mathcal{E}}$ ,  $\mathcal{M}_{\mathcal{F}}$ , and  $\mathcal{M}_{\partial\mathcal{F}}$  contain the edges 79 of  $\partial\Omega_i$ , subdomain edge  $\mathcal{E}$ , subdomain face  $\mathcal{F}$ , and  $\partial\mathcal{F}$ , respectively. We denote 80 by  $\mathcal{G}_{i\mathcal{F}}$ ,  $\mathcal{G}_{i\mathcal{E}}$ , and  $\mathcal{G}_{i\mathcal{V}}$  sets of subdomain faces, subdomain edges, and subdomain 81 vertices for  $\Omega_i$ . The wire basket  $\mathcal{W}_i$  is the union of all subdomain edges and vertices 82 for  $\Omega_i$ . We will also make use of the symbol  $\omega_i := 1 + \log(H_i/h_i)$ , and bold faced 83 symbols refer to vector functions. We denote by  $\bar{p}_i$  the mean of  $p_i$  over  $\Omega_i$ .

The estimate in the next lemma can be found in several references, see e.g., 85 Lemma 4.16 of [13].

**Lemma 1.** For any  $p_i \in W_{grad}^{h_i}$  and subdomain edge  $\mathscr{E}$  of  $\Omega_i$ ,

$$||p_i||_{L^2(\mathscr{E})}^2 \le C\omega_i ||p_i||_{H^1(\Omega_i)}^2.$$
 (5)

**Lemma 2.** For any  $p_i \in W_{grad}^{h_i}$ , there exist  $p_{i\mathscr{V}}, p_{i\mathscr{E}}, p_{i\mathscr{F}} \in W_{grad}^{h_i}$  such that

$$p_i|_{\partial\Omega_i} = \sum_{\mathcal{V} \in \mathcal{G}_{i\mathcal{V}}} p_{i\mathcal{V}}|_{\partial\Omega_i} + \sum_{\mathcal{E} \in \mathcal{G}_{i\mathcal{E}}} p_{i\mathcal{E}}|_{\partial\Omega_i} + \sum_{\mathcal{F} \in \mathcal{G}_{i\mathcal{F}}} p_{i\mathcal{F}}|_{\partial\Omega_i}, \tag{6}$$

where the nodal values of  $p_{i\mathscr{V}}$ ,  $p_{i\mathscr{E}}$ , and  $p_{i\mathscr{F}}$  on  $\partial\Omega_i$  may be nonzero only at the 90 nodes of  $\mathscr{V}$ ,  $\mathscr{E}$ , and  $\mathscr{F}$ , respectively. Further,

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$$|p_{i\mathscr{V}}|_{H^1(\Omega_i)}^2 \le C||p_i||_{H^1(\Omega_i)}^2,$$
 (7)

$$|p_{i\mathscr{E}}|_{H^{1}(\Omega_{i})}^{2} \le C\omega_{i}||p_{i}||_{H^{1}(\Omega_{i})}^{2},$$
 (8)

$$|p_{i\mathscr{F}}|_{H^{1}(\Omega_{i})}^{2} \le C\omega_{i}^{2} ||p_{i}||_{H^{1}(\Omega_{i})}^{2}.$$
 (9)

*Proof.* The estimates in (7)–(9) are standard, and follow from Corollary 4.20 and 93 Lemma 4.24 of [13] and elementary estimates.

We note that a Poincaré inequality allows us to replace the  $H^1$ -norm of  $p_i$  by its 95  $H^1$ -seminorm in Lemmas 1 and 2 if  $\bar{p}_i = 0$ .

The next lemma is stated without proof due to page restrictions.

**Lemma 3.** Let  $f_i \in W_{grad}^{h_i}$  have vanishing nodal values everywhere on  $\partial \Omega_i$  except on 98 the wire basket  $W_i$  of  $\Omega_i$ . For each subdomain face  $\mathscr{F}$  of  $\Omega_i$  and  $Ch_i \leq d \leq H_i/C$ , C>1, there exists a  $\mathbf{v}_i \in W_{curl}^{h_i}$  such that  $v_{ie} = \nabla f_{ie}$  for all  $e \in \mathcal{M}_{\mathscr{F}}$ ,  $v_{ie} = 0$  for all 100 other edges of  $\partial \Omega_i$ , and 101

$$\|\mathbf{v}_i\|_{L^2(\Omega_i)}^2 \le C(\boldsymbol{\omega}_i \|f_i\|_{L^2(\partial \mathscr{F})}^2 + d^2 \|\nabla f_i \cdot \mathbf{t}_{\partial \mathscr{F}}\|_{L^2(\partial \mathscr{F})}^2), \tag{10}$$

$$\|\nabla \times \mathbf{v}_i\|_{L^2(\Omega_i)}^2 \le C(\tau(d)\|f_i\|_{L^2(\partial\mathscr{F})}^2 + \|\nabla f_i \cdot \mathbf{t}_{\partial\mathscr{F}}\|_{L^2(\partial\mathscr{F})}^2),\tag{11}$$

where  $t_{\partial \mathcal{F}}$  is a unit tangent along  $\partial \mathcal{F}$ , and

$$au(d) = \left\{ egin{array}{ll} 0 & \mbox{if } d > H_i/C \ d^{-2} & \mbox{otherwise}. \end{array} 
ight.$$

The Helmholtz-type decomposition and estimates in the next lemma will allow 105 us to make use of and build on existing tools for scalar functions in  $H^1(\Omega_i)$ . We refer 106 the reader to Lemma 5.2 of [4] for the case of convex polyhedral subdomains; this 107 important paper was preceded by Hiptmair et al. [5], which concerns other applications of the same decomposition.

**Lemma 4.** For a convex and polyhedral subdomain  $\Omega_i$  and any  $m{u}_i \in W^{h_i}_{curb}$ , there is a 110  $\mathbf{q}_i \in W_{curl}^{h_i}, \ \mathbf{\Psi}_i \in (W_{grad}^{h_i})^3$ , and  $p_i \in W_{grad}^{h_i}$  such that 111

$$\boldsymbol{u}_i = \boldsymbol{q}_i + \Pi^{h_i}(\boldsymbol{\Psi}_i) + \nabla p_i, \tag{12}$$

$$\|\nabla p_i\|_{L^2(\Omega_i)} \le C \|\mathbf{u}_i\|_{L^2(\Omega_i)},$$
 (13)

$$\|\mathbf{\Psi}_i\|_{L^2(\Omega_i)} \le C \|\mathbf{u}_i\|_{L^2(\Omega_i)},$$
 (14)

$$\|h_i^{-1} \boldsymbol{q}_i\|_{L^2(\Omega_i)}^2 + \|\boldsymbol{\Psi}_i\|_{H^1(\Omega_i)}^2 \le C \|\nabla \times \boldsymbol{u}_i\|_{L^2(\Omega_i)}^2. \tag{15}$$

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**Lemma 5.** For any  $\mathbf{u}_i \in W_{curl}^{h_i}$  with  $u_{ie} = 0$  for all  $e \in \mathcal{M}_{\partial \mathcal{F}}$ , there exists a  $\mathbf{v}_{i\mathcal{F}} \in W_{curl}^{h_i}$ such that  $v_{i\mathscr{F}_e} = u_{ie}$  for all  $e \in \mathscr{M}_{\mathscr{F}}$ ,  $v_{i\mathscr{F}_e} = 0$  for all  $e \in \mathscr{M}_{\partial\Omega} \setminus \mathscr{M}_{\mathscr{F}}$ , and

$$E_i(\mathbf{v}_{i\mathscr{F}}) \le C\omega_i^2 E_i(\mathbf{u}_i),\tag{16}$$

where the energy  $E_i$  is defined in (3).

*Proof.* Let  $p_i$  in (12) be chosen so  $\bar{p}_i = 0$ . This is possible since a constant can be 116 added to  $p_i$  without changing its gradient. Because  $u_{ie} = 0$  for all  $e \in \mathcal{M}_{\partial \mathcal{R}}$ , it follows from Lemmas 1 and 4 and elementary estimates that 118

$$\|\nabla p_{i} \cdot \boldsymbol{t}_{\mathscr{E}}\|_{L^{2}(\partial \mathscr{F})}^{2} = \|(\Pi^{h_{i}}(\boldsymbol{\Psi}_{i}) + \boldsymbol{q}_{i}) \cdot \boldsymbol{t}_{\mathscr{E}}\|_{L^{2}(\partial \mathscr{F})}^{2}$$

$$\leq C\omega_{i} \|\nabla \times \boldsymbol{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}.$$
(17)

We then find from Lemmas 2 and 4 that

$$\|\nabla p_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq C\omega_{i}^{2}\|\mathbf{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}.$$
(18)

Define 120

$$p_{i\mathscr{W}} := \sum_{\mathscr{V} \in \mathscr{G}_{i\mathscr{V}}} p_{i\mathscr{V}} + \sum_{\mathscr{E} \in \mathscr{G}_{i\mathscr{E}}} p_{i\mathscr{E}}, \quad d := \begin{cases} H_i & \text{if } d_i \geq H_i \\ \max(d_i, Ch_i) & \text{otherwise,} \end{cases}$$

where  $d_i := \sqrt{\alpha_i/\beta_i}$ . Further, let  $p_{i\mathscr{W}}$  and  $p_{i\mathscr{F}}$  denote the functions  $f_i$  and  $v_i$ , respectively, of Lemma 3. We then find from Lemmas 1 and 3 and (17) that 123

$$E_i(\boldsymbol{p}_{i\mathscr{F}}) \le C\omega_i^2 E_i(\boldsymbol{u}_i), \tag{19}$$

where  $p_{i\mathscr{F}e}=\nabla p_{i\mathscr{W}e}\ \forall e\in\mathscr{M}_{\mathscr{F}}$  and  $p_{i\mathscr{F}e}=0\ \forall e\in\mathscr{M}_{\partial\Omega_i}\setminus\mathscr{M}_{\mathscr{F}}.$  With reference to (12) and (4), we define 125

$$\boldsymbol{q}_{i\mathscr{F}} := \sum_{e \in \mathscr{M}_{\mathscr{F}}} q_{ie} \boldsymbol{N}_e, \tag{20}$$

and from elementary finite element estimates and Lemma 4 find

$$\|\boldsymbol{q}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq Ch_{i}^{3} \sum_{e \in \mathscr{M}_{\mathscr{F}}} q_{ie}^{2} \leq C\|\boldsymbol{q}_{i}\|_{L^{2}(\Omega_{i})}^{2} \leq C\|\boldsymbol{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}, \tag{21}$$

$$\|\nabla \times \boldsymbol{q}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq Ch_{i} \sum_{e \in \mathscr{M}_{\mathscr{F}}} q_{ie}^{2} \leq C\|\nabla \times \boldsymbol{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}. \tag{22}$$

It follows from Lemmas 2 and 4 that there exists a  $\Psi_{i\mathscr{F}} \in (W^{h_i}_{\mathrm{grad}})^3$  such that  $\Psi_{i\mathscr{F}} = \Psi_i$  at all nodes of  $\mathscr{F}$ , that vanishes at all other nodes of  $\partial \Omega_i$ , and 128

$$\|\mathbf{\Psi}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq C\|\mathbf{\Psi}_{i}\|_{L^{2}(\Omega_{i})}^{2} \leq C\|\mathbf{u}_{i}\|_{L^{2}(\Omega_{i})}^{2},\tag{23}$$

$$\|\nabla \times \boldsymbol{\Psi}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq C\omega_{i}^{2}\|\boldsymbol{\Psi}_{i}\|_{H^{1}(\Omega_{i})}^{2} \leq C\omega_{i}^{2}\|\nabla \times \boldsymbol{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}. \tag{24}$$

From Lemmas 1 and 4, we obtain

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$$\|\boldsymbol{\Psi}_i\|_{L^2(\partial\mathscr{F})}^2 \leq C\omega_i\|\boldsymbol{\Psi}_i\|_{H^1(\Omega_i)}^2 \leq C\omega_i\|\nabla \times \boldsymbol{u}_i\|_{L^2(\Omega_i)}^2. \tag{25}$$

Let  $\Psi_{i\partial\mathscr{F}} \in (W_{\mathrm{grad}}^{h_i})^3$  be identical to  $\Psi_i$  at all nodes of  $\partial\mathscr{F}$  and vanish at all other nodes of  $\Omega_i$ . For  $\mathbf{g} := \Pi^{h_i}(\Psi_{i\partial\mathscr{F}})$ , we define

$$\mathbf{g}_{i\mathscr{F}} := \sum_{e \in \mathscr{M}_{\mathscr{F}}} g_e^{h_i} \mathbf{N}_e. \tag{26}$$

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From elementary estimates and (25,) we then obtain

$$\|\boldsymbol{g}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq Ch_{i}^{2}|\boldsymbol{\Psi}_{i}\|_{L^{2}(\partial\mathscr{F})}^{2} \leq C\omega_{i}h_{i}^{2}\|\nabla \times \boldsymbol{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}, \tag{27}$$

$$\|\nabla \times \mathbf{g}_{i\mathscr{F}}\|_{L^{2}(\Omega_{i})}^{2} \leq C\omega_{i}\|\nabla \times \mathbf{u}_{i}\|_{L^{2}(\Omega_{i})}^{2}.$$
(28)

Defining

$$\mathbf{v}_{i\mathscr{F}} := \nabla p_{i\mathscr{F}} + \mathbf{p}_{i\mathscr{F}} + \mathbf{q}_{i\mathscr{F}} + \Pi^{h_i}(\mathbf{\Psi}_{i\mathscr{F}}) + \mathbf{g}_{i\mathscr{F}}, \tag{29}$$

we find that  $v_{i\mathscr{F}e} = u_{ie} \ \forall e \in \mathscr{M}_{\mathscr{F}}$  and  $v_{i\mathscr{F}e} = 0 \ \forall e \in \mathscr{M}_{\partial\Omega_i} \setminus \mathscr{M}_{\mathscr{F}}$ . The estimate in 134 (16) then follows from the bounds for each of the terms on the right-hand-side of 135 (29) along with elementary estimates for  $\Pi^{h_i}(\mathbf{\Psi}_{i\mathscr{F}})$ .  $\square$ 

**3 BDDC** 

Background information and related theory for BDDC can be found in several references including [1, 2, 9–11]. Let  $u_i$  and u denote vectors of finite element coefficients associated with  $\Gamma_i$  and  $\Gamma$ . In general, entries in  $u_i$  and  $u_j$  are allowed to differ for  $j \neq i$  even though they refer to the same finite element edge. Entries in the vector  $\tilde{u}_i$  are partially continuous in the sense that specific edge values or edge averages over certain subsets of  $\Gamma$  are required to match for adjacent subdomains. In order to obtain consistent entries, we define the weighted average

$$\hat{u}_i = R_i \sum_{j=1}^N R_j^T D_j \tilde{u}_j, \tag{30}$$

where  $R_j$  is a 0–1 (Boolean) matrix that selects the rows of  $u_j$  from u and  $D_j$  is a 145 weight matrix. The weight matrices form a partition of unity in the sense that

$$\sum_{i=1}^{N} R_i^T D_i R_i = I, \tag{31}$$

where I is the identity matrix. To summarize,  $\hat{u}_i$  is fully continuous while  $\tilde{u}_i$  is only partially continuous. The number of continuity constraints that must be satisfied by all the  $\tilde{u}_i$  determines the dimension of the coarse space.

The energy of  $\boldsymbol{u}$  for  $\Omega_i$  can be expressed as

$$E_i(\mathbf{u}) = E_i(u_i) = u_i^T S_i u_i, \tag{32}$$

where  $S_i$  is the Schur complement matrix associated with  $\Omega_i$  and  $\Gamma_i$ . The system operator for BDDC is the assembled Schur complement 152

$$S = \sum_{i=1}^{N} R_i^T S_i R_i. \tag{33}$$

From Theorem 25 of [11], the condition number of the BDDC preconditioned operator is bounded above by 154

$$\kappa(M^{-1}S) \le \sup_{\tilde{u}_i} \frac{\sum_{i=1}^{N} \hat{u}_i^T S_i \hat{u}_i}{\sum_{i=1}^{N} \tilde{u}_i^T S_i \tilde{u}_i}.$$
 (34)

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This remarkably simple expression shows that the continuity constraints for  $\tilde{u}_i$  should 155 be chosen so that large increases in energy do not result from the averaging operation 156 in (30).

Let  $R_{i\partial\mathscr{F}_{ii}}$  select the rows of  $u_i$  corresponding to the edge coefficients on the 158 boundary of the face  $\mathscr{F}_{ij}$ , the closure of which is  $\partial \Omega_i \cap \partial \Omega_j$ . Similarly, let  $R_{i\mathscr{F}_{ij}}$ select the rows of  $u_i$  corresponding to the interior of the face  $\mathscr{F}_{ij}$ . We define the vector of face edge coefficients by  $u_{iF} := R_{i\mathscr{F}_{ii}}u_{i}$  and the face Schur complement matrix by  $S_{iFF} := R_{i\mathscr{F}_{ii}} S_i R_{i\mathscr{F}_{ii}}^T$ .

Because of page restrictions, we only consider a very rich coarse space which 163 includes every edge variable of each subdomain edge. This coarse space corresponds 164 to Algorithm C of [12]. For this case, we choose the weighted average of  $u_{iF}$  and  $u_{jF}$ as

$$\hat{u}_F = (S_{iFF} + S_{jFF})^{-1} (S_{iFF} u_{iF} + S_{jFF} u_{jF}). \tag{35}$$

Thus.

$$\hat{u}_F = (S_{iFF} + S_{jFF})^{-1} (S_{iFF} u_{iF} + S_{jFF} u_{jF}).$$

$$u_{iF} - \hat{u}_F = (S_{iFF} + S_{jFF})^{-1} S_{jFF} (u_{iF} - u_{jF}).$$
(35)

Using the eigenvectors of the generalized eigenvalue problem  $S_{iFF}x = \lambda S_{iFF}x$  as a 168 convenient basis, we find 169

$$u_{kF}^T \bar{S}_{iFF} u_{kF} \le u_{kF}^T S_{kFF} u_{kF}, \quad \forall u_{kF} \quad k \in \{i, j\}, \tag{37}$$

where

$$\bar{S}_{iFF} := S_{jFF} (S_{iFF} + S_{jFF})^{-1} S_{iFF} (S_{iFF} + S_{jFF})^{-1} S_{jFF}$$
(38)

Let us assume for the moment that there are vectors  $u_{ij}$ ,  $u_{ji}$ , and a scalar  $\hat{C} > 0$  such that 172

$$R_{i\partial\mathscr{F}_{ii}}u_{ij} = R_{j\partial\mathscr{F}_{ii}}u_{ji} = u_{\partial F}, \tag{39}$$

$$R_{i\mathscr{F}_{ij}}u_{ij} = R_{j\mathscr{F}_{ij}}u_{ji},\tag{40}$$

$$u_{ij}^{T} S_{i} u_{ij} + u_{ji}^{T} S_{j} u_{ji} \le \hat{C}(u_{i}^{T} S_{i} u_{i} + u_{j}^{T} S_{j} u_{j}). \tag{41}$$

In other words,  $u_{ij}$ ,  $u_{ji}$ ,  $u_i$  and  $u_j$  are all identical along the boundary of  $\mathscr{F}_{ij}$ . Further, 173  $u_{ij}$  and  $u_{ji}$  are identical in the interior of  $\mathcal{F}_{ij}$ , and the sum of their energies is bounded 174 uniformly by the sum of the energies of  $u_i$  and  $u_i$ . 175

In order to establish a condition number bound for Algorithm C, we need an estimate for  $E_i(R_{i\mathscr{F}_{ii}}^T(u_{iF} - \hat{u}_F))$ ; see (34). By construction, we have  $R_{i\partial\mathscr{F}_{ij}}(u_i - u_{ij}) = 0$ and  $R_{j\partial\mathscr{F}_{ii}}(u_j-u_{ji})=0$ . Since  $u_{iF}-u_{jF}=(u_{iF}-u_{ijF})-(u_{jF}-u_{jiF})$ , it then follows 178 from (36), (37), (41), and Lemma 5 that 179

$$E_{i}(R_{iF_{ij}}^{T}(u_{iF} - \hat{u}_{F})) = E_{i}(R_{iF_{ij}}^{T}(S_{iFF} + S_{jFF})^{-1}S_{jFF}(u_{iF} - u_{jF}))$$

$$\leq 2(u_{iF} - u_{ijF})^{T}S_{iFF}(u_{iF} - u_{ijF}) +$$

$$2(u_{jF} - u_{jiF})^{T}S_{jFF}(u_{jF} - u_{jiF})$$

$$\leq \hat{C}C\omega_{i}^{2}(E_{i}(u_{i}) + E_{j}(u_{j})). \tag{42}$$

We are able to show there exist  $u_{ij}$  and  $u_{ji}$  which satisfy the conditions in (39)–(41) 180 with  $\hat{C}$  independent of mesh parameters and the material properties  $\alpha_i$ ,  $\beta_i$ ,  $\alpha_j$ , and  $\beta_j$  181 under the assumption

$$\alpha_m \le C\alpha_n$$
 and  $\beta_m \le C\beta_n$  for  $\{m, n\} = \{i, j\}$  or  $\{m, n\} = \{j, i\}$ . (43)

This can be done using Lemma 4 together with an extension theorem for  $H^1$  functions on Lipschitz domains. We note that numerical experiments suggest that no 184 assumptions on subdomain material properties are needed, other than them being constant in each subdomain, for  $\hat{C}$  in (41) to be uniformly bounded.

Our main result follows from the estimate in (42).

**Theorem 1** (Condition Number Estimate). *Under the assumption in (43), the con-*188 dition number of the BDDC preconditioned operator for this study is bounded by 189

$$\kappa < C\omega^2$$
, (44)

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where

$$\omega = \max_{i} (1 + \log(H_i/h_i)). \tag{45}$$

In summary, we have obtained a favorable condition number estimate with less re- 191 strictive assumptions on the material properties of the subdomains than in previous 192 studies. Comparing the condition number estimate of Theorem 1 with those in (1) 193 and (2), we see that the factor of  $H_i^2 \beta_i / \alpha_i$  can be removed provided the assumption 194 in (43) holds. In addition, the logarithmic factor has been reduced from four pow- 195 ers to two. We note that the estimate in Theorem 1 also holds for FETI-DP due its 196 spectral equivalence with BDDC.

We note that the algorithm involves a non-standard averaging given by (35). This 198 averaging requires the solution of Dirichlet problems over the union of each pair of 199 subdomains sharing a face. The importance of this method of averaging for some 200 problems is shown in the next section.

#### 4 Numerical Results

In this section, we present some numerical results to verify the theory and also to 203 provide some additional insights. The domain is a unit cube discretized into smaller 204 cubic elements. All the examples are solved to a relative residual tolerance of  $10^{-8}$  205 for random right-hand-sides using the conjugate gradient algorithm with BDDC as 206 the preconditioner. The number of iterations and condition number estimates from 207 conjugate gradients are under the headings of iter and cond in the tables. We con- 208 sider three different types of weights for the averaging operator. The first one, des- 209 ignated SC, is the one based on (35). Unless otherwise specified in the tables, this 210 is the weighting used. The second type, stiff, is based on a conventional approach 211 in which the weights are proportional to the entries on the diagonals of subdomain 212 matrices. The third, card, uses the inverse of the cardinality of an edge, i.e. the recip-213 rocal of the number of subdomains sharing the edge, for the weight.

The results in Table 1 are consistent with theory, suggesting condition numbers 215 that are bounded independently of the number of subdomains, while the results in 216 Table 2 are consistent with the  $log(H/h)^2$  estimate of Theorem 1.

We also consider a checkerboard distribution of material properties in which 218  $(\alpha, \beta)$  for a subdomain is either  $(\alpha_1, \beta_1)$  or  $(\alpha_2, \beta_2)$ , and note that subdomains with 219 the same properties only share a subdomain vertex and no degrees of freedom. Re- 220 sults for 64 cubic subdomains each with H/h = 4 are shown in Table 3. Notice that 221 for only one choice of material properties in the table do all three types of weighting 222 lead to small condition numbers, and only the SC approach always gives condition 223 numbers which are independent of the material properties. We have also investigated 224 another type of weighting similar to card, but with weights  $\gamma$ ,  $0 < \gamma < 1$  for faces of 225 subdomains with properties  $\alpha_1$ ,  $\beta_1$  and  $1-\gamma$  for faces of subdomains with properties  $\alpha_2$ ,  $\beta_2$ . Regardless of the choice of  $\gamma$ , large condition numbers were observed for 227 the coefficients of the final row of Table 3. We note also that the choice of material 228 properties in the final row is not covered by the theory of [12].

In the final example, we consider a cubic mesh of 20<sup>3</sup> elements that is partitioned 230 into different numbers of subdomains using the graph partitioner Metis [8]. Although 231 this example is not covered by our theory because the subdomains have irregular 232 shapes, the results in Table 4 indicate that the algorithm of this study continues to 233 perform well. The results in Tables 3 and 4 suggest that the SC weighting of this 234 study may be necessary in order to effectively solve problems with material property 235 jumps or with subdomains of irregular shape.

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**Table 1.** Results for N cubic subdomains, each with  $\beta = 1$  and H/h = 4.

N	$\alpha = 10^2$ iter (cond)	$\alpha = 1$ iter (cond)	$\alpha = 10^{-2}$ iter (cond)
43	15 (2.70)	14 (2.63)	10 (1.77)
$6^3$	16 (2.88)	15 (2.81)	11 (2.05)
$8^3$	16 (2.95)	15 (2.87)	12 (2.23)
$10^{3}$	17 (2.98)	16 (2.91)	13 (2.33)

**Table 2.** Results for 64 cubic subdomains, each with  $\beta = 1$ .

H/k	$\alpha = 10^2$	$\alpha = 1$	$\alpha = 10^{-2}$
	iter (cond)	iter (cona)	iter (cond)
4	15 (2.70)	14 (2.63)	10 (1.77)
6	17 (3.30)	16 (3.21)	11 (2.14)
8	18 (3.77)	16 (3.66)	13 (2.46)
10	19 (4.16)	18 (4.03)	13 (2.72)

**Table 3.** Checkerboard material property results for 64 cubic subdomains with H/h = 4.

$\alpha_1$	$\beta_1$	$\alpha_2$	$\beta_2$		stiff iter (cond)	card iter (cond)
1	1	$10^{3}$	1	10 (1.59)	19 (4.57)	196 (1.64e3)
1	1	1	$10^{3}$	11 (1.96)	84 (2.69e2)	109 (4.72e2)
1	1	1	1.01	14 (2.63)	14 (2.63)	14 (2.63)
$10^{2}$	$10^{-2}$	1	1	6 (1.07)	65 (3.17e2)	74 (1.65e2)

**Table 4.** Results for  $20^3$  elements partitioned into N subdomains using a graph partitioner. Material properties are constant with  $\alpha = 1$  and  $\beta = 1$ .

N	SC iter (cond)	stiff iter (cond)	card iter (cond)
60	19 (4.30)	189 (6.31e2)	24 (9.06)
65	19 (4.40)	184 (6.34e2)	29 (1.55e3)
70	18 (3.89)	188 (6.47e2)	23 (7.48)
75	19 (4.16)	176 (6.12e2)	23 (6.49)

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# **Symbolic Techniques for Domain Decomposition Methods**

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1 Introduction 14

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Some algorithmic aspects of systems of PDEs based simulations can be better clar- 15 ified by means of symbolic computation techniques. This is very important since 16 numerical simulations heavily rely on solving systems of PDEs. For the large-scale 17 problems we deal with in today's standard applications, it is necessary to rely on 18 iterative Krylov methods that are scalable (i.e., weakly dependent on the number 19 of degrees on freedom and number of subdomains) and have limited memory re- 20 quirements. They are preconditioned by domain decomposition methods, incomplete 21 factorizations and multigrid preconditioners. These techniques are well understood 22 and efficient for scalar symmetric equations (e.g., Laplacian, biLaplacian) and to 23 some extent for non-symmetric equations (e.g., convection-diffusion). But they have 24 poor performances and lack robustness when used for symmetric systems of PDEs, 25 and even more so for non-symmetric complex systems (fluid mechanics, porous me- 26 dia...). As a general rule, the study of iterative solvers for systems of PDEs as op- 27 posed to scalar PDEs is an underdeveloped subject.

We aim at building new robust and efficient solvers, such as domain decomposi- 29 tion methods and preconditioners for some linear and well-known systems of PDEs. 30 In particular, we shall concentrate on Neumann-Neumann and FETI type algorithms 31 which are very popular for scalar symmetric positive definite second order problems 32 (see, for instance, [9, 11]), and to some extent to different other problems, like the 33 advection-diffusion equations [1], plate and shell problems [16] or the Stokes equa- 34 tions [13]. This work is motivated by the fact that, in some sense, these methods 35 applied to systems of PDEs (such as Stokes, Oseen, linear elasticity) are less op- 36 timal than the domain decomposition methods for scalar problems. Indeed, in the 37

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case of two subdomains consisting of the two half planes, it is well-known that the 38 Neumann-Neumann preconditioner is an exact preconditioner (the preconditioned 39 operator is the identity operator) for the Schur complement equation for scalar equations like the Laplace problem. Unfortunately, this does not hold in the vector case.

In order to achieve this goal, we use algebraic methods developed in construc- 42 tive algebra, D-modules (differential modules) and symbolic computation such as the 43 so-called Smith or Jacobson normal forms and Gröbner basis techniques for trans- 44 forming a linear system of PDEs into a set of independent PDEs. These algebraic and 45 symbolic methods provide important intrinsic information (e.g., invariants) about the 46 linear system of PDEs to solve. These build-in properties need to be taken into ac- 47 count in the design of new numerical methods, which can supersede the usual ones 48 based on a direct extension of the classical scalar methods to linear systems of PDEs. 49

By means of these techniques, it is also possible to transform the linear system of 50 PDEs into a set of decoupled PDEs under certain types of invertible transformations. 51 One of these techniques is the so-called Smith normal form of the matrix of OD 52 operators associated with the linear system. This normal form was introduced by H. 53 J. S. Smith (1826–1883) for matrices with integer entries (see, e.g., [17], Theorem 54 1.4). The Smith normal form has already been successfully applied to open problems 55 in the design of Perfectly Matched Layers (PML). The theory of PML for scalar 56 equations was well-developed and the usage of the Smith normal form allowed to 57 extend these works to systems of PDEs. In [12], a general approach is proposed and 58 applied to the particular case of the compressible Euler equations that model aero- 59 acoustic phenomena and in [2] for shallow-water equations.

For domain decomposition methods, several results have been obtained on com- 61 pressible Euler equations [7], Stokes and Oseen systems [8] or in [10] where a new 62 method in the "Smith" spirit has been derived. Previously the computations were 63 performed heuristically, whereas in this work, we aim at finding a systematic way to 64 build optimal algorithms for given PDE systems.

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**Notations.** If R is a ring, then  $R^{p\times q}$  is the set of  $p\times q$  matrices with entries in 66 R and  $\operatorname{GL}_n(R)$  is the group of invertible matrices of  $R^{p \times p}$ , namely  $\operatorname{GL}_n(R) = \{E \in {}^{67}$  $R^{p \times p} \mid \exists F \in R^{p \times p} : EF = FE = I_p$ . An element of  $GL_p(R)$  is called an *unimodular* 68 matrix. A diagonal matrix with elements  $d_i$ 's will be denoted by diag $(d_1, \dots, d_p)$ . If k 69 is a field (e.g.,  $k = \mathbb{Q}, \mathbb{R}, \mathbb{C}$ ), then  $k[x_1, \dots, x_n]$  is the commutative ring of polynomials 70 in  $x_1, \ldots, x_n$  with coefficients in k. In what follows,  $k(x_1, \ldots, x_n)$  will denote the field 71 of rational functions in  $x_1, \dots, x_n$  with coefficients in k. Finally, if  $r, r' \in R$ , then  $r' \mid r$ means that r' divides r, i.e., there exists  $r'' \in R$  such that r = r'' r'.

# 2 Smith Normal Form of Linear Systems of PDEs

We first introduce the concept of Smith normal form of a matrix with polynomial 75 entries (see, e.g., [17], Theorem 1.4). The Smith normal form is a mathematical 76 technique which is classically used in module theory, linear algebra, symbolic computation, ordinary differential systems, and control theory. It was first developed to 78 study matrices with integer entries. But, it was proved to exist for any principal ideal 79 domain (namely, a commutative ring R whose ideals can be generated by an element 80 of R) [15]. Since R = k[s] is a principal ideal domain when k is a field, we have the following theorem only stated for square matrices.

**Theorem 1.** Let k be a field, R = k[s], p a positive integer and  $A \in \mathbb{R}^{p \times p}$ . Then, there exist two matrices  $E \in GL_p(R)$  and  $F \in GL_p(R)$  such that

$$A = ESF,$$
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where  $S = \text{diag}(d_1, \dots, d_p)$  and the  $d_i \in R$  satisfying  $d_1 | d_2 | \dots | d_p$ . In particular, 86 we can take  $d_i = m_i/m_{i-1}$ , where  $m_i$  is the greatest common divisor of all the  $i \times i$ -87 minors of A (i.e., the determinants of all  $i \times i$ -submatrices of A), with the convention 88 that  $m_0 = 1$ . The matrix  $S = \text{diag}(d_1, \dots, d_p) \in \mathbb{R}^{p \times p}$  is called a Smith normal form 89

We note that  $E \in GL_p(R)$  is equivalent to det(E) is an invertible polynomial, i.e., 91  $det(E) \in k \setminus \{0\}$ . Also, in what follows, we shall assume that the  $d_i$ 's are monic polynomials, i.e., their leading coefficients are 1, which will allow us to call the matrix 93  $S = \operatorname{diag}(d_1, \dots, d_p)$  the Smith normal form of A. But, the unimodular matrices E and 94 F are not uniquely defined by A. The proof of Theorem 1 is constructive and gives 95 an algorithm for computing matrices E, S and F. The computation of Smith normal 96 forms is available in many computer algebra systems such as Maple, Mathematica, 97 Magma...

Consider now the following model problem in  $\mathbb{R}^d$  with d = 2, 3:

$$\mathcal{L}_d(\mathbf{w}) = \mathbf{g} \quad \text{in } \mathbb{R}^d, \quad |\mathbf{w}(\mathbf{x})| \to 0 \quad \text{for } |\mathbf{x}| \to \infty.$$
 (1)

For instance,  $\mathcal{L}_d(\mathbf{w})$  can represent the Stokes/Oseen/linear elasticity operators in 100 dimension d. Moreover, if we suppose that the inhomogeneous linear system of PDEs 101 (1) has constant coefficients, then it can be rewritten as

$$A_d \mathbf{w} = \mathbf{g},\tag{2}$$

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where  $A_d \in R^{p \times p}$ ,  $R = k[\partial_x, \partial_y]$  (resp.,  $R = k[\partial_x, \partial_y, \partial_z]$ ) for d = 2 (resp., d = 3) and k is a field.

In what follows, we shall study the domain decomposition problem in which  $\mathbb{R}^d$ is divided into subdomains. We assume that the direction normal to the interface 106 of the subdomains is particularized and denoted by  $\partial_x$ . If  $R_x = k(\partial_y)[\partial_x]$  for d =2 or  $R_x = k(\partial_y, \partial_z)[\partial_x]$  for d = 3, then, computing the Smith normal form of the matrix  $A_d \in R_x^{p \times p}$ , we obtain  $A_d = ESF$ , where  $S \in R_x^{p \times p}$  is a diagonal matrix,  $E \in {}^{109}$  $\operatorname{GL}_n(R_x)$  and  $F \in \operatorname{GL}_n(R_x)$ . The entries of the matrices E, S, F are polynomials in  $\partial_x$ , and E and F are unimodular matrices, i.e.,  $\det(E)$ ,  $\det(F) \in k(\partial_y) \setminus \{0\}$  if d=2, or  $\det(E)$ ,  $\det(F) \in k(\partial_{v}, \partial_{z}) \setminus \{0\}$  if d = 3. We recall that the matrices E and F are 112 not unique contrary to S. Using the Smith normal form of  $A_d$ , we get:

$$A_d \mathbf{w} = \mathbf{g} \quad \Leftrightarrow \quad \{ \mathbf{w_s} := F \mathbf{w}, S \mathbf{w_s} = E^{-1} \mathbf{g} \}.$$
 (3)

In other words, (3) is equivalent to the uncoupled linear system:

$$S\mathbf{w_s} = E^{-1}\mathbf{g}. (4)$$

Since  $E \in GL_p(R_x)$  and  $F \in GL_p(R_x)$ , the entries of their inverses are still polynomial in  $\partial_x$ . Thus, applying  $E^{-1}$  to the right-hand side  $\mathbf{g}$  of  $A_d$   $\mathbf{w} = \mathbf{g}$  amounts 116 to taking k-linear combinations of derivatives of **g** with respect to x. If  $\mathbb{R}^d$  is split 117 into two subdomains  $\mathbb{R}^- \times \mathbb{R}^{d-1}$  and  $\mathbb{R}^+ \times \mathbb{R}^{d-1}$ , where  $\mathbb{R}^- = \{x \in \mathbb{R} \mid x < 0\}$  and 118  $\mathbb{R}^+ = \{x \in \mathbb{R} \mid x > 0\}$ , then the application of  $E^{-1}$  and  $F^{-1}$  to a vector can be done for each subdomain independently. No communication between the subdomains is 120 necessary.

In conclusion, it is enough to find a domain decomposition algorithm for the 122 uncoupled system (4) and then transform it back to the original one (2) by means of 123 the invertible matrix F over  $R_x$ . This technique can be applied to any linear system 124 of PDEs once it is rewritten in a polynomial form. The uncoupled system acts on the 125 new dependent variables ws, which we shall further call Smith variables since they 126 are issued from the Smith normal form.

Remark 1. Since the matrix F is used to transform (4) to (2) (see the first equation of 128 the right-hand side of (3)) and F is not unique, we need to find a matrix F as simple 129 as possible (e.g., F has minimal degree in  $\partial_x$ ) so that to obtain a final algorithm 130 whose form can be used for practical computations. 131

**Example 1** Consider the two dimensional elasticity operator defined by  $\mathscr{E}_2(\mathbf{u}) :=$  $-\mu \Delta \mathbf{u} - (\lambda + \mu) \nabla \text{div } \mathbf{u}$ . If we consider the commutative polynomial rings R = $\mathbb{Q}(\lambda,\mu)[\partial_x,\partial_y], R_x = \mathbb{Q}(\lambda,\mu)(\partial_y)[\partial_x] = \mathbb{Q}(\lambda,\mu,\partial_y)[\partial_x]$  and

$$A_2 = \begin{pmatrix} (\lambda + 2\mu) \, \partial_x^2 + \mu \, \partial_y^2 & (\lambda + \mu) \, \partial_x \, \partial_y \\ (\lambda + \mu) \, \partial_x \, \partial_y & \mu \, \partial_x^2 + (\lambda + 2\, \mu) \, \partial_y^2 \end{pmatrix} \in R^{2\times 2}$$
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the matrix of PD operators associated with  $\mathcal{E}_2$ , i.e.,  $\mathcal{E}_2(\mathbf{u}) = A_2 \mathbf{u}$ , then the Smith 136 normal form of  $A_2 \in R_x^{2 \times 2}$  is defined by: 137

$$S_{A_2} = \begin{pmatrix} 1 & 0 \\ 0 & \Delta^2 \end{pmatrix}. \tag{5}$$

The particular form of  $S_{A_2}$  shows that, over  $R_x$ , the system of PDEs for the linear 138 elasticity in  $\mathbb{R}^2$  is algebraically equivalent to a biharmonic equation. 139

**Example 2** Consider the two dimensional Oseen operator  $\mathscr{O}_2(\mathbf{w}) = \mathscr{O}_2(\mathbf{v},q) := 140$  $(c\mathbf{v} - v\Delta\mathbf{v} + \mathbf{b} \cdot \nabla\mathbf{v} + \nabla q, \nabla \cdot \mathbf{v})$ , where **b** is the convection velocity. If **b** = 0, then 141 we obtain the Stokes operator  $\mathscr{S}_2(\mathbf{w}) = \mathscr{S}_2(\mathbf{v},q) := (c\mathbf{v} - v\Delta\mathbf{v} + \nabla q, \nabla \cdot \mathbf{v})$ . If 142  $R = \mathbb{Q}(b_1, b_2, c, v)[\partial_x, \partial_y], R_x = \mathbb{Q}(b_1, b_2, c, v)(\partial_y)[\partial_x] = \mathbb{Q}(b_1, b_2, c, v, \partial_y)[\partial_x]$  and

$$O_{2} = \begin{pmatrix} -v\left(\partial_{x}^{2} + \partial_{y}^{2}\right) + b_{1}\partial_{x} + b_{2}\partial_{y} + c & 0 & \partial_{x} \\ 0 & -v\left(\partial_{x}^{2} + \partial_{y}^{2}\right) + b_{1}\partial_{x} + b_{2}\partial_{y} + c & \partial_{y} \\ \partial_{x} & \partial_{y} & 0 \end{pmatrix}$$
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the matrix of PD operators associated with  $\mathcal{O}_2$ , i.e.,  $\mathcal{O}_2(\mathbf{w}) = O_2 \mathbf{w}$ , then the Smith normal form of  $O_2 \in R_x^{3 \times 3}$  is defined by:

$$S_{O_2} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & \Delta L_2 \end{pmatrix}, \quad L_2 = c - v \Delta + \mathbf{b} \cdot \nabla. \tag{6}$$

From the form of  $S_{O_2}$  we can deduce that the two-dimensional Oseen equations can be mainly characterized by the scalar fourth order PD operator  $\Delta L_2$ . This is not surprising since the stream function formulation of the Oseen equations for d=2 150 gives the same PDE for the stream function.

Remark 2. The above applications of Smith normal forms suggest that one should design an optimal domain decomposition method for the biharmonic operator  $\Delta^2$  153 (resp.,  $L_2\Delta$ ) in the case of linear elasticity (resp., the Oseen/Stokes equations) for 154 the two-dimensional problems, and then transform it back to the original system.

### 3 An Optimal Algorithm for the Biharmonic Operator

We give here an example of Neumann-Neumann methods in its iterative version 157 for Laplace and biLaplace equations. For simplicity, consider a decomposition of 158 the domain  $\Omega = \mathbb{R}^2$  into two half planes  $\Omega_1 = \mathbb{R}^- \times \mathbb{R}$  and  $\Omega_2 = \mathbb{R}^+ \times \mathbb{R}$ . Let the 159 interface  $\{0\} \times \mathbb{R}$  be denoted by  $\Gamma$  and  $(\mathbf{n}_i)_{i=1,2}$  be the outward normal of  $(\Omega_i)_{i=1,2}$ . 160 We consider the following problem:

$$-\Delta u = f \text{ in } \mathbb{R}^2, \quad |u(\mathbf{x})| \to 0 \text{ for } |\mathbf{x}| \to \infty.$$
 (7)

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and the following **Neumann-Neumann algorithm** applied to problem (7): 162 Let  $u_{\Gamma}^n$  be the interface solution at iteration n. We obtain  $u_{\Gamma}^{n+1}$  from  $u_{\Gamma}^n$  by the following iterative procedure

$$\begin{cases}
-\Delta u^{i,n} = f, & \text{in } \Omega_i, \\
u^{i,n} = u_{\Gamma}^n, & \text{on } \Gamma,
\end{cases}
\begin{cases}
-\Delta \tilde{u}^{i,n} = 0, & \text{in } \Omega_i, \\
\frac{\partial \tilde{u}^{i,n}}{\partial \mathbf{n}_i} = -\frac{1}{2} \left( \frac{\partial u^{1,n}}{\partial \mathbf{n}_1} + \frac{\partial u^{2,n}}{\partial \mathbf{n}_2} \right), & \text{on } \Gamma,
\end{cases} (8)$$

and then  $u_{\Gamma}^{n+1} = u_{\Gamma}^{n} + \frac{1}{2} (\tilde{u}^{1,n} + \tilde{u}^{2,n}).$ 

This algorithm is *optimal* in the sense that it converges in two iterations.

$$\Delta^2 \phi = g \text{ in } \mathbb{R}^2, \quad |\phi(\mathbf{x})| \to 0 \text{ for } |\mathbf{x}| \to \infty.$$
 (9)

and the following "Neumann-Neumann" type algorithm applied to (9): 171 Let  $(\phi_{\Gamma}^n, D\phi_{\Gamma}^n)$  be the interface solution at iteration n (suppose also that  $\phi_{\Gamma}^0 = 172$ 

 $\phi^0|_{\Gamma}$ ,  $D\phi^0_{\Gamma}=(\Delta\phi^0)_{\Gamma}$ ). We obtain  $(\phi^{n+1}_{\Gamma},D\phi^n_{\Gamma})$  from  $(\phi^n_{\Gamma},D\phi^n_{\Gamma})$  by the following iterative procedure 174

$$\begin{cases} -\Delta^{2}\phi^{i,n} = f, & \text{in } \Omega_{i}, \\ \phi^{i,n} = \phi_{\Gamma}^{n}, & \text{on } \Gamma, \\ \Delta\phi^{i,n} = D\phi_{\Gamma}^{n}, & \text{on } \Gamma, \end{cases} \begin{cases} \frac{\partial\tilde{\phi}^{i,n}}{\partial\mathbf{n}_{i}} = -\frac{1}{2}\left(\frac{\partial\phi^{1,n}}{\partial\mathbf{n}_{1}} + \frac{\partial\phi^{2,n}}{\partial\mathbf{n}_{2}}\right), & \text{on } \Gamma, \\ \frac{\partial\Delta\tilde{\phi}^{i,n}}{\partial\mathbf{n}_{i}} = -\frac{1}{2}\left(\frac{\partial\Delta\phi^{1,n}}{\partial\mathbf{n}_{1}} + \frac{\partial\Delta\phi^{2,n}}{\partial\mathbf{n}_{2}}\right), & \text{on } \Gamma, \end{cases}$$
and then  $\phi_{\Gamma}^{n+1} = \phi_{\Gamma}^{n} + \frac{1}{2}\left(\tilde{\phi}^{1,n} + \tilde{\phi}^{2,n}\right), D\phi_{\Gamma}^{n+1} = D\phi_{\Gamma}^{n} + \frac{1}{2}\left(\tilde{\Delta}\phi^{1,n} + \tilde{\Delta}\phi^{2,n}\right).$ (10)

This is a generalization of the Neumann-Neumann algorithm for the  $\Delta$  operator 177 and is also *optimal* (the proof can be found in [8]).

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Now, in the case of the two dimensional linear elasticity,  $\phi$  represents the second component of the vector of Smith variables, that is,  $\phi = (\mathbf{w}_s)_2 = (F\mathbf{u})_2$ , where  $\mathbf{u} = (u, v)$  is the displacement field. Hence, we need to replace  $\phi$  with  $(F\mathbf{u})_2$  into the algorithm for the biLaplacian, and then simplify it using algebraically admissible op- 182 erations. Thus, one can obtain an optimal algorithm for the Stokes equations or linear 183 elasticity depending on the form of F. From here comes the necessity of choosing 184 in a proper way the matrix F (which is not unique), used to define the Smith normal 185 form, in order to obtain a "good" algorithm for the systems of PDEs from the optimal 186 one applied to the biharmonic operator. In [7] and [8], the computation of the Smith 187 normal forms for the Euler equations and the Stokes equations was done by hand or 188 using the Maple command Smith. Surprisingly, the corresponding matrices F have 189 provided good algorithms for the Euler equations and the Stokes equations even if 190 the approach was entirely heuristic.

## 4 Relevant Smith Variables: A Completion Problem

The efficiency of our algorithms heavily relies on the simplicity of the Smith vari- 193 ables, that is on the entries of the unimodular matrix F used to compute the Smith 194 normal form of the matrix A. In this section, within a constructive algebraic analysis 195 approach, we develop a method for constructing many possible Smith variables. Tak- 196 ing into account physical aspects, the user can then choose the simplest one among 197 them. We are going to show that the problem of finding Smith variables can be re- 198 duced to a completion problem. First of all, we very briefly introduce some notions 199 of module theory [15].

Given a ring R (e.g.,  $R = k[\partial_1, \dots, \partial_d]$ , where k is a field (e.g.,  $\mathbb{Q}$ ,  $\mathbb{R}$ ,  $\mathbb{C}$ )), the 201 definition of a R-module M is similar to the one of a vector space but where the 202 scalars are taken in the ring R and not in a field as for vector spaces. If  $A \in \mathbb{R}^{p \times p}$ , 203 then the kernel of the *R*-linear map (*R*-homomorphism)  $A: R^{1 \times p} \longrightarrow R^{1 \times p}$ , defined 204 by  $(A)(\mathbf{r}) = \mathbf{r}A$ , is the *R*-module defined by:

$$\ker_{R}(A) = \{ \mathbf{r} \in R^{1 \times p} \mid \mathbf{r}A = 0 \}.$$
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The image  $\operatorname{im}_R(A)$  of A, simply denoted by  $R^{1\times p}A$ , is the R-module defined by 207 all the R-linear combinations of the rows of A. The cokernel  $\operatorname{coker}_R(A)$  of A is 208 the *factor* R-module defined by  $\operatorname{coker}_R(A) = R^{1\times p}/(R^{1\times p}A)$ . To simplify the no-209 tation, we shall denote this module by M. M is nothing more than the R-module 210 of the row vectors of  $R^{1\times p}$  modulo the R-linear combinations of rows of A. Let 211  $R_1 = k(\partial_2, \ldots, \partial_d)[\partial_1], \ R_i = k(\partial_1, \ldots, \partial_{i-1}, \partial_{i+1}, \ldots, \partial_d)[\partial_i], \ i = 2, \ldots, d-1$ , and 212  $R_d = k(\partial_1, \ldots, \partial_{d-1})[\partial_d]$  be the polynomial rings in  $\partial_i$  with coefficients in the field 213 of rational functions in all other PD operators.

Since the R-module  $M = R^{1 \times p}/(R^{1 \times p}A)$  plays a fundamental role in what follows, let us describe it in terms of generators and relations. Let  $\{\mathbf{f_j}\}_{j=1,\dots,p}$  be the 216 standard basis of  $R^{1 \times p}$ , namely  $\mathbf{f_j}$  is the row vector of  $R^{1 \times p}$  defined by 1 at the jth 217 position and 0 elsewhere, and  $m_j$  the residue class of  $\mathbf{f_j}$  in M. Then,  $\{m_j\}_{j=1,\dots,p}$  218 is a family of generators of the R-module M, i.e., for any  $m \in M$ , then there explicts  $\mathbf{r} = (r_1, \dots, r_p) \in R^{1 \times p}$  such that  $m = \sum_{j=1}^p r_j m_j$  [3]. The family of generators 220  $\{m_j\}_{j=1,\dots,p}$  of M satisfies the relations  $\sum_{j=1}^p A_{ij} m_j = 0$  for all  $i = 1,\dots,p$  [3]. For 221 more details, see [3, 15].

Let  $E, F \in \operatorname{GL}_p(R_i)$  be two unimodular matrices such that A = ESF, where 223  $S = \operatorname{diag}(1, \dots, 1, d_{r+1}, \dots, d_q)$  is the Smith normal form of A. Moreover, let us split 224  $F \in \operatorname{GL}_p(R_i)$  into two parts row-wise, i.e.,  $F = (F_1^T \quad F_2^T)^T$ , where  $F_1 \in R_i^{r \times p}$ ,  $F_2 \in P_i^{r \times p}$ , and  $F_i \in R_i^{r \times p}$ , and  $F_i \in R_$ 

$$A = ESF \quad \Leftrightarrow \quad \begin{pmatrix} F_1 \\ S_2 F_2 \end{pmatrix} = E^{-1}A, \quad S_2 = \operatorname{diag}(d_{r+1}, \dots, d_p). \tag{11}$$

Cleaning the denominators of the entries of  $S_2$  (resp.,  $F_2$ ), we can assume without 227 loss of generality that the  $d_j$ 's (resp., the entries of  $F_2$ ) belong to R. Then, (11) shows 228 that the jth row of  $F_2$  must be an element of the  $R_i$ -module  $M_i = R_i^{1 \times p}/(R_i^{1 \times p}A)$  annihilated by  $d_j$ . Consequently, the possible  $F_2$ 's can be found by computing a family 230 of generators of the  $R_i$ -modules ann $M_i(d_j) = \{m \in M_i \mid d_j m = 0\}$  for  $j = r+1, \ldots, p$ . 231 These  $R_i$ -modules can be computed by means of *Gröbner basis techniques* (see, e.g., 232 [6]). Hence, we get  $S_2F_2 = G_2A$  for some  $G_2 \in R_i^{(p-r) \times p}$ . Then, for each choice for 233  $F_2$ , we are reduced to the following *completion problem*:

Find 
$$F_1 \in R_i^{r \times p}$$
 such that  $F = (F_1^T \quad F_2^T)^T \in GL_p(R_i)$  and  $F_1 = G_1A$  for some  $G_1 \in R_i^{r \times p}$ . (12)

**Example 3** Let  $R = \mathbb{Q}(\lambda, \mu)[\partial_x, \partial_y, \partial_z]$  be the commutative polynomial ring of PD 235 operators in  $\partial_x$ ,  $\partial_y$  and  $\partial_z$  with coefficients in the field  $\mathbb{Q}(\lambda, \mu)$ , 236

$$A = \begin{pmatrix} -(\lambda + \mu) \, \partial_x^2 - \mu \, \Delta & -(\lambda + \mu) \, \partial_x \, \partial_y & -(\lambda + \mu) \, \partial_x \, \partial_z \\ -(\lambda + \mu) \, \partial_x \, \partial_y & -(\lambda + \mu) \, \partial_y^2 - \mu \, \Delta & -(\lambda + \mu) \, \partial_y \, \partial_z \\ -(\lambda + \mu) \, \partial_x \, \partial_z & -(\lambda + \mu) \, \partial_y \, \partial_z & -(\lambda + \mu) \, \partial_z^2 - \mu \, \Delta \end{pmatrix} \in R^{3 \times 3}$$
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the matrix of PD operators defining the elastostatic equations in  $\mathbb{R}^3$ , where  $\Delta = \partial_x^2 + 238$   $\partial_y^2 + \partial_z^2$ , and the associated *R*-module  $M = R^{1\times3}/(R^{1\times3}A)$ . The Smith normal form 239

of A with respect to x is given by  $S = \operatorname{diag}(1, \Delta, \Delta^2)$ . With the above notations, we get r = 1 and  $S_2 = \operatorname{diag}(\Delta, \Delta^2) \in R^{2 \times 2}$ . Let  $R_x = \mathbb{Q}(\lambda, \mu)(\partial_y, \partial_z)[\partial_x]$ ,  $F_1 \in R_x^{1 \times 3}$  and  $F_2 \in \mathbb{Z}^2$  and  $F_2 \in \mathbb{Z}^2$ . Then, the first (resp. second) row of  $F_2$  must be an element of the  $R_x$ -module  $R_x = R_x^{1 \times 3}/(R_x^{1 \times 3}A)$  annihilated by  $A \in R$  (resp.  $A^2 \in R$ ). Using the Oremodules package [4], we find that families of generators of  $\operatorname{ann}_{M_x}(\Delta)$  and  $\operatorname{ann}_{M_x}(\Delta^2)$  are  $R_x = R_x = R_$ 

$$A_{\Delta} = \begin{pmatrix} 0 & -\partial_{z} & \partial_{y} \\ \partial_{z} & 0 & -\partial_{x} \\ -\partial_{y} & \partial_{x} & 0 \\ \partial_{x} & \partial_{y} & \partial_{z} \end{pmatrix}, \quad A_{\Delta^{2}} = I_{3}.$$

That simply means that a family of generators of  $\operatorname{ann}_{M_x}(\Delta)$  is given by the divergence 248 and the curl of the displacement field and for  $\operatorname{ann}_{M_x}(\Delta^2)$  by the components of the 249 displacement fields. Now, the first (resp., second) row of  $F_2$  must be a  $R_x$ -linear 250 combination of the rows of  $A_\Delta$  (resp.,  $A_{\Delta^2}$ ). We thus have several choices and for 251 each of them, we are reduced to a completion problem (12). For instance, choosing 252 the first row of  $A_\Delta$  (resp., the third row of  $A_{\Delta^2}$ ) as first (resp., second) row of  $F_2$ , 253 namely

$$F_2 = \begin{pmatrix} 0 & -\partial_z & \partial_y \\ 0 & 0 & 1 \end{pmatrix}, \tag{255}$$

we then have to find a row vector  $F_1 \in R_x^{1 \times 3}$  such that  $F_1 = G_1 A$  for some  $G_1 \in R_x^{1 \times 3}$  256 and  $F = (F_1^T \quad F_2^T)^T \in GL_3(R_x)$ . If such a row vector  $F_1$  exists, then the matrix 257  $F = (F_1^T \quad F_2^T)^T$  provides a good choice of Smith variables.

We first give two necessary conditions for a choice of  $F_2$  to provide a solution of the completion problem (straightforward from the relation A = ESF):

**Lemma 1.** With the above notations, given  $F_2 \in R^{(p-r)\times p}$ , necessary conditions for the solvability of the completion problem (12) are:

1. 
$$F_2$$
 admits a right inverse over  $R_i$ , i.e.  $\exists S_2 \in R_i^{p \times (p-r)}$ :  $F_2 S_2 = I_{p-r}$ .

2. There exists a matrix  $G_2 \in R_i^{(p-r) \times p}$  such that  $S_2 F_2 = G_2 A$ .

Since  $R_i$  is a *principal ideal domain* (namely, every ideal of  $R_i$  can be generated 265 by an element of  $R_i$ ), Condition 1 of Lemma 1 is equivalent to the condition that 266 the  $R_i$ -module  $\operatorname{coker}_{R_i}(.F_2) = R_i^{1 \times p}/(R_i^{1 \times (p-r)}F_2)$  is free of rank r, i.e.  $\operatorname{coker}_{R_i}(.F_2)$  267 admits a basis of cardinality r [3, 15]. It is equivalent to the existence of two matrices 268  $Q_2 \in R_i^{p \times r}$  and  $T_2 \in R_i^{r \times p}$  such that  $\ker_{R_i}(.Q_2) = R_i^{1 \times (p-r)}F_2$  and  $T_2 = I_r$  [3]. Such 269 a matrix  $Q_2$  is called an *injective parametrization* of  $\operatorname{coker}_{R_i}(.F_2)$ . Matrices  $Q_2$  and 270  $T_2$  can be computed by Gröbner basis techniques [3]. The corresponding algorithms 271 are implemented in the OREMODULES package [4]. The next theorem characterizes 272 the solvability of the completion problem (12).

**Theorem 2.** Let  $F_2 \in R^{(p-r)\times p}$  admit a right inverse over  $R_i$  and satisfy  $S_2F_2 = G_2A$  274 for some  $G_2 \in R_i^{(p-r)\times p}$ . If  $Q_2$  is an injective parametrization of the free  $R_i$ -module 275  $\operatorname{coker}_{R_i}(.F_2)$  of rank r, and  $T_2 \in R_i^{r\times p}$  a left inverse of  $Q_2$ , then a necessary and 276 sufficient condition for the existence of a solution of the completion problem (12) is 277 the existence of two matrices  $H \in R_i^{r\times (p-r)}$  and  $G_1 \in R_i^{r\times p}$  such that  $T_2 = G_1A - HF_2$ . 278 Then,  $F_1 = T_2 + HF_2 = G_1A$  is a solution of the completion problem (12), i.e., F = 279  $((T_2 + HF_2)^T F_2^T)^T \in \operatorname{GL}_p(R_i)$  is such that A = ESF for some  $E \in \operatorname{GL}_p(R_i)$ , where 280 S is the Smith normal form of A.

From the explanations above, we deduce the following algorithm that, given 282 A,  $S_2 = \text{diag}(d_{r+1}, \ldots, d_p)$ , and a choice for  $F_2$  computed from the calculations of 283  $\text{ann}_{M_i}(d_j)$  for  $d_j \in R$ , find (if it exists) a completion of  $F_2$ . The following algorithm

Input:  $A \in R^{p \times p}$ ,  $S_2 \in R^{(p-r) \times (p-r)}$  and  $F_2 \in R^{(p-r) \times p}$ .

Output: A completion  $F = (F_1^T \quad F_2^T)^T$  of  $F_2$  or "No completion exists".

- 1. Compute a right inverse of  $F_2$  over  $R_i$ ;
- 2. If no right inverse exists, then RETURN "No completion exists", Else
  - (a) Factorize  $S_2 F_2$  with respect to A over  $R_i$ ;
  - (b) If no factorization exists, then RETURN "No completion exists", Else
    - i. Compute an injective parametrization  $Q_2$  of  $\operatorname{coker}_{R_i}(.F_2)$ ;
    - ii. Compute a left inverse  $T_2$  of  $Q_2$  over  $R_i$ ;
    - iii. Factorize  $T_2$  with respect to  $(F_2^T A^T)^T$  over  $R_i$ ;
    - iv. If no factorization exists, then RETURN "No completion exists", Else note  $T_2 = (-H G_1) \begin{pmatrix} F_2 \\ A \end{pmatrix}$  and RETURN  $F = \begin{pmatrix} T_2 + H F_2 \\ F_2 \end{pmatrix}$ .

was implemented in Maple based on the OREMODULES package.

**Example 4** Consider again the elastostatic equations introduced in Example 3. For the choice of  $F_2$  given at the end of Example 3, our implementation succeeds in finding a completion and we get the following completion of  $F_2$ : 288

$$F = \begin{pmatrix} 1 - \frac{\partial_{x} \partial_{y}}{\partial_{y}^{2} + \partial_{z}^{2}} - \frac{\partial_{x} ((\lambda + 2\mu)(\partial_{x}^{2} + \partial_{y}^{2}) + (2\lambda + 3\mu)\partial_{z}^{2})}{(\lambda + \mu)\partial_{z}(\partial_{y}^{2} + \partial_{z}^{2})} \\ 0 - \partial_{z} & \partial_{y} \\ 0 & 0 & 1 \end{pmatrix} \in GL_{3}(R_{x}).$$
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For more details and explicit computations, we refer the reader to [5].

#### 5 Reduction of the Interface Conditions

In the algorithms presented in the previous sections, we have equations in the domains  $\Omega_i$  and interface conditions on  $\Gamma$  obtained heuristically. We need to find an 293

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automatic way to reduce the interface conditions with respect to the equations in the 294 domains. In this section, we show how symbolic computations can be used to per- 295 form such reductions. The naïve idea consists in gathering all equations and compute 296 a Gröbner basis [6]. However, one has to keep in mind that the independent variables 297 do not play the same role. More precisely, the interface conditions cannot be dif- 298 ferentiated with respect to x since the border of the interface is defined by  $x \neq 0$ . 299 Consequently, we have developed and implemented an alternative method in Maple 300 using the OREMODULES package, which can be sketched as follows:

1. Compute a Gröbner basis of the polynomial equations inside the domain for a 302 relevant monomial order:

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- 2. Compute the normal forms of the interface conditions with respect to the latter 304 Gröbner basis;
- 3. Write these normal forms in the jet notations with respect to the independent 306 variable x, i.e., rewrite the derivatives  $\partial_x^i y_k$  of the dependent variables  $y_k$  as new 307 indeterminates  $y_{k,i}$ ;
- 4. Perform linear algebra manipulations to simplify the normal forms.

For more details and explicit computations, we refer the reader to [5].

#### 6 Some Optimal Algorithms

After performing the completion and the reduction of the interface conditions, we 312 can give examples of optimal algorithms (elasticity and Stokes equations). 313

**Example 5** Consider the elasticity operator:

$$\mathcal{E}_d \mathbf{u} = -\operatorname{div} \sigma(\mathbf{u}), \quad \sigma(\mathbf{u}) = \mu \left( \nabla \mathbf{u} + (\nabla \mathbf{u})^T \right) + \lambda \operatorname{div} \mathbf{u} I_d.$$
 315

If d = 2, then the completion algorithm gives two possible choices for F:

$$F = \begin{pmatrix} -\frac{\partial_x (\mu \, \partial_x^2 - \lambda \, \partial_y^2)}{(\lambda + \mu) \, \partial_y^3} & 1\\ 1 & 0 \end{pmatrix}, \quad F = \begin{pmatrix} 1 - \frac{(\lambda + \mu)\partial_x ((3\,\mu + 2\,\lambda) \, \partial_y^2 + (2\,\mu + \lambda) \, \partial_x^2)}{\partial_y^3}\\ 0 & 1 \end{pmatrix}. \quad (13)$$

By replacing  $\phi$  into the Neumann-Neumann algorithm for the biLaplacian by  $(F\mathbf{u})_2$  317 and re-writing the interface conditions, using the equations inside the domain like in 318 [8], we get two different algorithms for the elasticity system. Note that, in the first 319 case of (13),  $\phi = u$ , and, in the second one,  $\phi = v$  (where  $\mathbf{u} = (u, v)$ ). Below, we shall 320 write in detail the algorithm in the second case. To simplify the writing, we denote 321 by  $u_{\tau} = \mathbf{u} \cdot \mathbf{\tau}$ ,  $u_{\mathbf{n}} = \mathbf{u} \cdot \mathbf{n}$ ,  $\sigma_{\mathbf{nn}}(\mathbf{u}) = (\sigma(\mathbf{u}) \cdot \mathbf{n}) \cdot \mathbf{n}$ ,  $\sigma_{\mathbf{n\tau}}(\mathbf{u}) = (\sigma(\mathbf{u}) \cdot \mathbf{n}) \cdot \mathbf{\tau}$ . 322

Let  $(u_{\Gamma}^n, \sigma_{\Gamma}^n)$  be the interface solution at iteration n (suppose also that  $u_{\Gamma}^0 = (u_{\tau}^0)|_{\Gamma}$ , 323  $\sigma_{\Gamma}^0 = (\sigma_{\rm snn}(u^0))|_{\Gamma}$ ). We obtain  $(u_{\Gamma}^{n+1}, \sigma_{\Gamma}^n)$  from  $(u_{\Gamma}^n, \sigma_{\Gamma}^n)$  by the following iterative 324 procedure 325

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$$\begin{cases} \mathscr{E}_{2}(\mathbf{u}^{i,n}) &= f, & \text{in } \Omega_{i}, \\ u_{\tau_{i}}^{1,n} &= u_{\Gamma}^{n}, & \text{on } \Gamma, \\ \sigma_{\mathbf{n}_{i}\mathbf{n}_{i}}(\mathbf{u}^{i,n}) &= \sigma_{\Gamma}^{n}, & \text{on } \Gamma, \end{cases} \begin{cases} \mathscr{E}_{2}(\tilde{\mathbf{u}}^{i,n}) &= 0, & \text{in } \Omega_{i}, \\ \tilde{\mathbf{u}}_{\tau_{i}}^{i,n} &= -\frac{1}{2}\left(\mathbf{u}_{\mathbf{n}_{1}}^{1,n} + \mathbf{u}_{\mathbf{n}_{2}}^{2,n}\right), & \text{on } \Gamma, \\ \sigma_{\mathbf{n}_{i}\tau_{i}}(\tilde{\mathbf{u}}^{i,n}) &= -\frac{1}{2}\left(\sigma_{\mathbf{n}_{1}\tau_{1}}(\mathbf{u}^{1,n}) + \sigma_{\mathbf{n}_{2}\tau_{2}}(\mathbf{u}^{2,n})\right), & \text{on } \Gamma, \end{cases}$$

$$(14)$$

and 
$$u_{\Gamma}^{n+1} = u_{\Gamma}^{n} + \frac{1}{2} \left( \tilde{u}_{\tau_{1}}^{1,n} + \tilde{u}_{\tau_{2}}^{2,n} \right), \, \sigma_{\Gamma}^{n+1} = \sigma_{\Gamma}^{n} + \frac{1}{2} \left( \sigma_{\mathbf{n_{1}n_{1}}}(\tilde{\mathbf{u}}^{1,n}) + \sigma_{\mathbf{n_{2}n_{2}}}(\tilde{\mathbf{u}}^{2,n}) \right).$$
 326

Remark 3. We found an algorithm with a mechanical meaning: Find the tangential 327 part of the normal stress and the normal displacement at the interface so that the normal part of the normal stress and the tangential displacement on the interface match. 329 This is very similar to the original Neumann-Neumann algorithm, which means that 330 the implementation effort of the new algorithm from an existing Neumann-Neumann 331 is negligible (the same type of quantities – displacement fields and efforts – are im- 332 posed at the interfaces), except that the new algorithm requires the knowledge of 333 some geometric quantities, such as normal and tangential vectors. Note also that, 334 with the adjustment of the definition of tangential quantities for d=3, the algorithm is the same, and is also similar to the results in [8].

7 Conclusion 337

All algorithms and interface conditions are derived for problems posed on the whole 338 space, since for the time being, this is the only way to treat from the algebraic point 339 of view these problems. The effect of the boundary condition on bounded domains 340 cannot be quantified with the same tools. All the algorithms are designed in the 341 PDE level and it is very important to choose the right discrete framework in order 342 to preserve the optimal properties. For example, in the case of linear elasticity a 343 good candidate would be the TDNNS finite elements that can be found in [14]. The 344 implementation and the impact of the discretizations on the algorithms is an ongoing 345 work. 346

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Scalable Domain Decomposition Algorithms is	for
Contact Problems: Theory, Numerical Exper	iments,
and Real World Problems	

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Summary. We review our results related to the development of theoretically supported scalable algorithms for the solution of large scale contact problems of elasticity. The algorithms 11 combine the Total FETI/BETI based domain decomposition method adapted to the solution of 12 2D and 3D multibody contact problems of elasticity, both frictionless and with friction, with 13 our in a sense optimal algorithms for the solution of resulting quadratic programming and 14 OPOC problems. Rather surprisingly, the theoretical results are qualitatively the same as the 15 classical results on scalability of FETI/BETI for linear elliptic problems. The efficiency of the method is demonstrated by results of parallel numerical experiments for contact problems of 17 linear elasticity discretized by more than 11 million variables in 3D and 40 million variables 18 in 2D.

1 Introduction 20

Contact problems are in the heart of mechanical engineering. Solving large multi- 21 body contact problems of linear elastostatics is complicated by the inequality bound- 22 ary conditions, which make them strongly non-linear, and, if the system of bodies 23 includes "floating" bodies, by the positive semi-definite stiffness matrices resulting 24 from the discretization of such bodies. Observing that the classical Dirichlet and 25 Neumann boundary conditions are known only after the solution has been found, it 26 is natural to assume the solution of contact problems to be more costly than the so- 27 lution of a related linear problem with the classical boundary conditions. Since the 28 cost of the solution of any problem increases at least linearly with the number of the 29 unknowns, it follows that the development of a scalable algorithm for contact prob- 30 lems is a challenging task which requires to identify the contact interface in a sense 31 for free.

The first promising results, at least for the frictionless problems, were obtained 33 by the researchers who tried to modify the methods that were known to be scalable 34

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for linear problems, in particular multigrid and domain decomposition. Experimental 35 evidence of scalability was achieved with the monotonic multigrid (see [11] and the 36 references therein). In spite of these nice results, the necessity to keep the coarse 37 grid away from the contact interface prevented the authors to prove the optimality 38 results similar to the classical results for linear problems. However, such result was 39 obtained by Schöberl who has developed an approximate variant of the projection 40 method using a domain decomposition preconditioner and a linear multigrid solver 41 on the interior nodes. An experimental evidence of scalability for the frictionless 42 problems was presented by Avery and Farhat [1]. The point of this paper is to report 43 our optimality results for contact problems of linear elasticity, both frictionless and 44 with friction.

The results are based on a combination of several ingredients. The first one is the 46 application of the TFETI (Total FETI) [8] or TBETI (Total BETI) [14] methods, vari- 47 ants of the duality based domain decomposition methods introduced by Farhat and 48 Roux [9] (finite elements) and Langer and Steinbach [13] (boundary elements). Since 49 the TFETI/TBETI methods treat all the subdomains as "floating", the kernels of the 50 stiffness matrices of the subdomains are a priori known. This makes the method very 51 flexible and simplifies implementation of the multiplication of a vector by a gener- 52 alized inverse of the stiffness matrix. As any duality based method, TFETI/TBETI 53 reduces general inequality constraints to special separable ones.

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The second ingredient is the "natural coarse grid preconditioning" introduced for 55 linear problems by Farhat, Mandel, and Roux [10] and Langer and Steinbach [13]. 56 This preconditioned cost function has the spectrum of the Hessian confined to a pos- 57 itive interval independent of the discretization parameter h and the decomposition 58 parameter H provided the ratio H/h is uniformly bounded. Since our preconditioning uses a projector to the subspace with the solution, it follows that its application 60 to the solution of variational inequalities does not turn the separable constraints into 61 general constraints and can be interpreted as a variant of the multigrid method with 62 the coarse grid on the interface. This unique feature, as compared with the standard 63 multigrid preconditioning for the primal problem, reduces the development of scal- 64 able algorithms for the solution of variational inequalities to the solution of bound 65 and equality constrained quadratic programming or QPQC (quadratic programming 66 with quadratic constraints) problems with the rate of convergence in terms of bounds 67 on the spectrum.

The resulting QP and QPQC problems, arising in the solution of the frictionless 69 contact problems and the problems with the Tresca friction (an auxiliary problem for 70 Coulomb friction), respectively, are solved by our algorithms with the rate of convergence in terms of the bounds on the spectrum, the third ingredient of our development 72 (see [7]). Putting the three ingredients together with a few simple observations, we 73 get theoretically supported algorithms for contact problems. The theoretical results 74 are illustrated by the results of numerical experiments which show that both numeri- 75 cal and parallel scalability can be observed in practice. Finally we report the solutions 76 of some real world problems. More details can be found in Dostál et al. [3–5], and 77 Sadowská et al. [14].

## 2 Dual Formulation of Frictionless Contact Problems

To simplify our presentation, let us assume that the bodies are assembled from  $N_s$  80 subdomains  $\Omega^{(s)}$  which are "glued" together by suitable equality constraints. After 81 the standard finite element discretization, the equilibrium of the system is described 82 as a solution u of the problem

$$\min J(v)$$
 subject to  $\sum_{s=1}^{N_s} B_N^{(s)} v^{(s)} \le g_N$  and  $\sum_{s=1}^{N_s} B_E^{(s)} v^{(s)} = o$ , (1)

where o denotes the zero vector and J(v) is the energy functional defined by

$$J(v) = \sum_{s=1}^{N_s} \frac{1}{2} v^{(s)T} K^{(s)} v^{(s)} - v^{(s)T} f^{(s)},$$
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 $v^{(s)}$  and  $f^{(s)}$  denote the admissible subdomain displacements and the subdomain vector of prescribed forces,  $K^{(s)}$  is the subdomain stiffness matrix,  $B_N^{(s)} \in \mathbb{R}^{m_C \times n}$  and 87  $B_E^{(s)} \in \mathbb{R}^{m_E \times n}$  are the blocks of the matrix  $B = \begin{bmatrix} B_N^T, B_E^T \end{bmatrix}^T$  that correspond to  $\Omega^{(s)}$ , 88 and  $g_N$  is a vector collecting the normal gaps between the bodies in the reference 89 configuration. The matrix  $B_N$  and the vector  $g_N$  arise from the nodal or mortar de-90 scription of the non-penetration conditions, while  $B_E$  describes the "gluing" of the subdomains into the bodies and the Dirichlet boundary conditions. Recall that if the 92 problem is discretized by the TBETI method, then we get the potential energy minimization problem of the very same structure as (1), where all the objects correspond 94 only to the boundaries  $\Gamma^{(s)}$  of  $\Omega^{(s)}$  except the term with the prescribed volume forces 95 (if there is some); see [14] for more details. By contrast with TFETI, when the ma- 96 trices  $K^{(s)}$  are sparse, in the case of TBETI these are fully populated.

To simplify the presentation of basic ideas, we can describe the equilibrium in 98 terms of the global stiffness matrix K, the vector of global displacements u, and the vector of global loads f. In the TFETI/TBETI methods, we have

$$K = \operatorname{diag}(K^{(1)}, \dots, K^{(N_s)}), \quad u = \begin{bmatrix} u^{(1)} \\ \vdots \\ u^{(N_s)} \end{bmatrix}, \quad \text{and} \quad \mathbf{f} = \begin{bmatrix} f^{(1)} \\ \vdots \\ f^{(N_s)} \end{bmatrix},$$

where  $K^{(s)}$ ,  $s = 1, ..., N_s$ , is a positive semidefinite matrix. The energy function reads 102

$$j(v) = \frac{1}{2}v^T K v - f^T v \tag{103}$$

and the vector of global displacements u solves

$$\min j(v)$$
 s.t.  $B_N v \le g_N$  and  $B_E v = o$ .

Alternatively, the global equilibrium may be described by the Karush–Kuhn– 106 Tucker conditions (see, e.g., [6]) 107

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$$Ku = f - B^T \lambda, \quad \lambda_N \ge o, \quad \lambda^T (Bu - g) = o,$$
 (2)

where  $g = [g_N^T, o^T]^T$  and  $\lambda = [\lambda_N^T, \lambda_E^T]^T$  denotes the vector of Lagrange multipliers 108 which may be interpreted as the reaction forces. The problem (2) differs from the 109 linear problem by the non-negativity constraint on the components of reaction forces 110  $\lambda_N$  and by the complementarity condition.

We can use the first equation of (2) to eliminate the displacements. We shall get 112 the problem to find 113

min 
$$\Theta(\lambda)$$
 s.t.  $\lambda_N \ge o$  and  $R^T(f - B^T \lambda) = o$ , (3)

where

min 
$$\Theta(\lambda)$$
 s.t.  $\lambda_N \ge o$  and  $R^T(f - B^T \lambda) = o$ , (3)
$$\Theta(\lambda) = \frac{1}{2} \lambda^T B K^+ B^T \lambda - \lambda^T (B K^+ f - g) + \frac{1}{2} f K^+ f,$$
 (4)

 $K^+$  denotes a generalized inverse that satisfies  $KK^+K = K$ , and R denotes the full 115 rank matrix whose columns span the kernel of K. The action of  $K^+$  can be evaluated at the cost comparable with that of Cholesky's decomposition applied to the 117 regularized K (see [2]). Denoting  $\mathscr{F} = ||BK^+B^T||$ ,

$$F = \mathscr{F}^{-1}BK^+B^T, \quad e = SR^Tf, \quad G = SR^TB^T, \quad \widetilde{d} = \mathscr{F}^{-1}(BK^\dagger f - g),$$

with S denoting a nonsingular matrix that defines the orthonormalization of the rows 120 of  $R^TB^T$ , we can modify (3) to 121

$$\min \widetilde{\theta}(\lambda)$$
 s.t.  $\lambda_N \ge 0$  and  $G\lambda = e$ , (5)

where

$$\widetilde{\theta}(\lambda) = \frac{1}{2} \lambda^T F \lambda - \lambda^T \widetilde{d}. \tag{6}$$

Our next step is to replace the equality constraint in (5) by a homogeneous one. 123 To this end, it is enough to find any  $\lambda$  such that 124

$$G\lambda = e,$$
 125

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denote  $\lambda = \mu + \widetilde{\lambda}$ , and substitute into (5). We get

$$\widetilde{\theta}(\lambda) = \frac{1}{2}\mu^T F \mu - \mu^T (\widetilde{d} - F\widetilde{\lambda}) + const.$$

After returning to the old notation, problem (5) is reduced to

$$\min \frac{1}{2} \lambda^T F \lambda - \lambda^T d \quad \text{s.t.} \quad G \lambda = o \quad \text{and} \quad \lambda_N \ge \ell_N$$
 (7)

with  $\ell = -\widetilde{\lambda}$  and  $d = \widetilde{d} - F\widetilde{\lambda}$ . Since G has orthonormal rows, we can use the least square solution 130

$$\widetilde{\lambda} = G^T e. \tag{8}$$

## 3 Dual Formulation of Contact Problems with Tresca Friction

If the Tresca friction is prescribed on the contact interface, then the equilibrium of u the system is described as a solution u of the problem

$$\min J_T(v)$$
 subject to  $\sum_{s=1}^{N_s} B_N^{(s)} v^{(s)} \le g_N$  and  $\sum_{s=1}^{N_s} B_E^{(s)} v^{(s)} = o$ , (9)

where  $J_T(v)$  is the energy functional defined by

$$J_T(v) = J(v) + j(v), \quad j(v) = \sum_{i=1}^{m_C} \Psi_i \|T_i u\|,$$
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 $\Psi_i$  denotes an a priori defined slip bound at node i, and  $T_i u$  denotes the jump of the tangential displacement due to the displacement u. Using the standard procedure to modify the non-differentiable term j (see [3, 5]), we get

$$j(v) = \sum_{i=1}^{m_C} \Psi_i \|T_i u\| = \sum_{i=1}^{m_C} \max_{\tau_i \| \le \Psi_i} \tau_i^T T_i u,$$
 139

where  $\tau_i$  can be considered as Lagrange multipliers. We assume that  $B_N$ ,  $B_E$ , and  $T_{140}$  are full rank matrices.

Let  $\overline{d}$  denote the spatial dimension and let us introduce the Lagrangian with 142 three types of Lagrange multipliers, namely  $\lambda_N \in \mathbb{R}^{m_C}$  associated with the non-143 interpenetration condition,  $\lambda_E \in \mathbb{R}^{m_E}$  associated with the "gluing" and prescribed 144 displacements, and

$$au = [ au_1^T, au_2^T, \dots, au_{m_C}^T]^T \in \mathbb{R}^{(\overline{d}-1)m_C}$$
 146

which regularizes the non-differentiability. The Lagrangian associated with problem 147 (1) reads

$$L(u, \lambda_N, \lambda_E, \tau) = J(u) + \tau^T T u + \lambda_N^T (B_N u - c_N) + \lambda_E^T (B_E u - c_E).$$
 (10)

Using the convexity of the cost function and constraints, we can use the classical duality theory [6] to reformulate problem (9) to get

$$\min_{\substack{u \\ \lambda_E \in \mathbb{R}^{m_E}, \ \lambda_N \geq \mathbf{o} \\ \|\tau_i\| \leq \Psi_i, \ i=1,\dots,m_C}} L(u,\lambda_N,\lambda_E,\tau) = \max_{\substack{\lambda_E \in \mathbb{R}^{m_E}, \ \lambda_N \geq o \\ \|\tau_i\| \leq \Psi_i, \ i=1,\dots,m_C}} \min_{\substack{u \\ u = 1,\dots,m_C}} L(u,\lambda_N,\lambda_E,\tau).$$
 151

To simplify the notation, we denote

$$\lambda = \begin{bmatrix} \lambda_E \\ \lambda_N \\ \tau \end{bmatrix}, \quad B = \begin{bmatrix} B_E \\ B_N \\ T \end{bmatrix}, \quad c = \begin{bmatrix} c_E \\ c_N \\ o \end{bmatrix},$$
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and 154

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$$\Lambda(\boldsymbol{\Psi}) = \left\{ (\boldsymbol{\lambda}_E^T, \boldsymbol{\lambda}_N^T, \boldsymbol{\tau}^T)^T \in \mathbb{R}^{m_E + \overline{d}m_C} : \ \boldsymbol{\lambda}_N \geq o, \|\boldsymbol{\tau}_i\| \leq \boldsymbol{\Psi}_i, \ i = 1, \dots, m_C \right\},$$
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so that we can write the Lagrangian briefly as

$$L(u,\lambda) = \frac{1}{2}u^T K u - f^T u + \lambda^T (Bu - c)$$
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and problem (9) is equivalent to the saddle point problem

$$L(\widehat{u},\widehat{\lambda}) = \max_{\lambda \in \Lambda(\Psi)} \min_{u} L(u,\lambda). \tag{11}$$

Similarly to the frictionless case, we eliminate the primal variables from (11) and 159 carry out the homogenization to reduce the minimization problem to 160

$$\min \frac{1}{2} \lambda^T F \lambda - \lambda^T d \quad \text{s.t.} \quad G \lambda = o \quad \text{and} \quad \lambda \in \Lambda(\Psi)$$
 (12)

with the notation of Sect. 2. Notice that we minimize exactly the same type of the cost 161 function as in the frictionless case, but with some additional quadratic constraints.

# 4 Preconditioning by Projector

Our final step is based on the observation that both the frictionless contact problem 164 and the contact problem with Tresca friction are equivalent to 165

$$\min \theta(\lambda)$$
 s.t.  $\lambda \in \Omega$ , (13)

where

$$heta(\lambda) = rac{1}{2} \lambda^T (PFP + \overline{
ho}Q) \lambda - \lambda^T P \, d, \quad Q = G^T (GG^T)^{-1} G, \quad P = I - Q,$$
 167

 $\overline{\rho} > 0$ , and  $\Omega = \{\lambda : G\lambda = o \text{ and } \lambda_N \ge o\}$  (without friction) or  $\Omega = \{\lambda : G\lambda = o \text{ 168}\}$  $\lambda \in \Lambda(\Psi)$ } (Tresca). A good choice of the regularization parameter is given 169 by 170

$$\overline{\rho} = \|PFP\|,\tag{171}$$

as this is the largest value for which

$$||PFP|| > ||PFP + \overline{\rho}O||.$$

Problem (13) turns out to be a suitable starting point for development of an ef- 174 ficient algorithm for variational inequalities due to the following classical estimates 175 [10] of the extreme eigenvalues. 176

**Theorem 1.** If the decompositions and the discretizations of given contact problems 177 are sufficiently regular, then there are constants  $C_1 > 0$  and  $C_2 > 0$  independent of 178 the discretization parameter h and the decomposition parameter H such that 179

$$C_1 \frac{h}{H} \le \lambda_{\min}(PFP|\operatorname{Im}P) \quad and \quad \lambda_{\max}(PFP|\operatorname{Im}P) = ||PFP|| \le C_2,$$
 (14)

where  $\lambda_{\min}$  and  $\lambda_{\max}$  denote the extremal eigenvalues of the corresponding matrices. 180

**5** Optimality 181

Theorem 1 states that if we fix the regularization parameter  $\overline{\rho}$  and keep H/h uniformly bounded, then problem (13) resulting from the application of various discretizations and decompositions has the spectrum of the Hessian matrices confined 184 to a positive interval. It follows that to develop a scalable algorithm for the contact 185 problems, it is enough to find an algorithm that is able to find an approximate so- 186 lution of (13) in a number of matrix-vector multiplications uniformly bounded in 187 terms of bounds on the spectrum of the cost function.

Here we propose to use SMALSE (semi-monotonic augmented Lagrangian 189 method for separable and equality constraints), our variant of the augmented La- 190 grangian method [7]. SMALSE enforces the equality constraints by the Lagrange 191 multipliers generated in the outer loop, while the auxiliary QPQC problems with sep- 192 arable constraints are solved approximately in the inner loop by the MPGP algorithm 193 proposed by Dostál and Kozubek [7]. MPGP is an active set based algorithm which uses the conjugate gradient method to explore the current face, the fixed steplength 195 gradient projection to change the active set, and the adaptive precision control for 196 the solution of auxiliary linear problems. The unique feature of SMALSE with the 197 inner loop implemented by MPGP when used to (13) is the bound on the number of 198 iterations whose cost is proportional to the number of variables, so that it can return 199 an approximate solution for the cost proportional to the number of variables. It fol- 200 lows that SMALSE/MPGP is a scalable algorithm for the solution of (13) provided 201 the cost of decomposition of K and application of the projectors P and Q is not too 202 large.

**Theorem 2.** If the decompositions and the discretizations of a given contact prob- 204 lem are sufficiently regular, then there is a constant C > 0 independent of the discretization parameter h and the decomposition parameter H such that the algorithm 206 SMALSE/MPGP (or SMALBE/MPRGP for the frictionless problems) with fixed parameters specified in [7] can find the solution of (13) in a number of iterations 208 bounded by C provided the initial approximation satisfies

$$\|\lambda^0\| \le c\|Pd\|,$$
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where c > 0 is an a priori chosen constant.

# **6 Numerical Experiments**

The algorithms reported in this paper were implemented into our MatSol software 213 [12] and tested with the aim to verify their optimality and capability to solve the real 214 world problems.

#### 6.1 Scalability of TFETI: 2D Cantilever Beams with Tresca Friction

We first tested the scalability on a 2D problem of Fig. 1 with varying discretiza- 217 tions and decompositions using structured grids. We kept the ratio H/h of the 218 decomposition and the discretization parameters approximately constant so that the 219 assumptions of Theorem 1 were satisfied.

The results of computations carried out to the relative precision  $10^{-4}$  are in 221 Table 1. We can observe that the number of matrix-vector multiplications varies only 222 mildly with the increasing dimension of the problem in agreement with the theory. 223 We conclude that the scalability can be observed in practice.

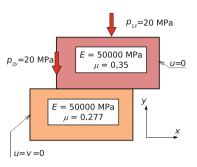


Fig. 1. Geometry of 2D cantilever beams

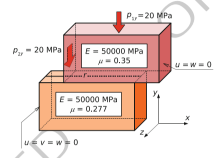


Fig. 2. Geometry of 3D cantilever beams

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Table 1. Numerical scalability of TFETI: 2D cantilever beams.

Number of subdomains	1936	4096	7744
Number of CPUs	48	48	48
Primal variables	10,071,072	21,307,392	40,284,288
Dual variables	384,473	817,793	1,551,089
Null space dimension	5808	12,288	23,232
SMALSE-M iterations	8	8	8
Hessian multiplications	119	134	180
Solution time [s]	839	1665	7825

## 6.2 Scalability of TFETI/TBETI: 3D Cantilever Beams with Tresca Friction

The second problem was a 3D alternative to the previous example (see Fig. 2). The 226 results of computations carried out for both TFETI and TBETI methods are in Ta- 227 bles 2 and 3, respectively. We can see that the number of matrix-vector multiplica- 228 tions again varies only mildly with the increasing problem size as predicted by the 229 theory.

**Table 2.** Numerical scalability of TFETI: 3D cantilever beams.

Number of subdomains	108	500	1372	2916
	108	300	13/2	2910
Number of CPUs	48	48	48	48
Primal variables	431,244	1,996,500	5,478,396	11,643,588
Dual variables	88,601	444,927	1,261,493	2,728,955
Null space dimension	648	3000	8232	17,496
SMALSE-M iterations	3	4	4	4
Hessian multiplications	78	97	93	119
Solution time [s]	60	374	1663	7745

Table 3. Numerical scalability of TBETI: 3D cantilever beams.

Number of subdomains	108	500	1372	2916
Number of CPUs	48	48	48	48
Primal variables	195,045	903,000	2,477,830	5,266,300
Dual variables	88,601	444,927	1,261,493	2,728,955
Null space dimension	648	3000	8232	17,496
SMALSE-M iterations	7	8	9	9
Hessian multiplications	160	161	160	260
Solution time [s]	46	301	2211	7949

## 6.3 Applications of TFETI/TBETI to Real World Problems

We have also tested our algorithms on real world problems. First we consider the 232 analysis of the stress in the roller bearings of Fig. 3. The problem is difficult because 233 it consists of 73 bodies in mutual contact and only one is fixed in space. The solution 234 of the problem discretized by 2,730,000/459,800 primal/dual variables and decom- 235 posed into 700 subdomains required 4,270 matrix-vector multiplications. The von 236 Mises stress distribution is in Fig. 3.

Second we consider the analysis of the yielding clamp connection of steel arched 238 supports depicted in Fig. 4. This type of construction is used to support the min- 239 ing openings. It is a typical multibody contact, where the yielding connection plays 240 the role of the mechanical protection against destruction, i.e., against the total de- 241 formation of the supporting arches. We consider contact with the Coulomb friction, 242 where the coefficient of friction was  $\mathscr{F} = 0.5$ . The problem was decomposed into 243 250 subdomains using METIS and discretized by 1,592,853 and 216,604 primal and 244 dual variables, respectively. The total displacements for both TFETI and TBETI are 245 depicted in Fig. 4. The solution required 1,922 matrix-vector multiplications.

#### 7 Comments and Conclusions

The TFETI method turns out to be a powerful engine for the solution of contact prob- 248 lems of elasticity. The results of numerical experiments comply with the theoretical 249 results and indicate high efficiency of the method reported here. Future research will 250 include adaptation of the standard preconditioning strategies.

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Fig. 3. Frictionless roller bearing of wind generator

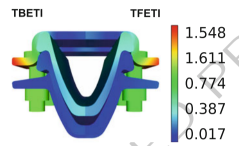


Fig. 4. Steel support with Coulomb friction

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# **Robust Coarsening in Multiscale PDEs**

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1 Introduction 5

Consider a variationally-posed second-order elliptic boundary value problem

$$a(u,v) \equiv \int_{\Omega} \mathcal{A}(\mathbf{x}) \nabla u \cdot \nabla v = \int_{\Omega} f(\mathbf{x}) v(\mathbf{x}), \text{ for all } v \in H_0^1(\Omega),$$
 (1)

with solution  $u \in H_0^1(\Omega)$  and domain  $\Omega \subset \mathbb{R}^d$ , d = 2,3, where the coefficient ten-7 sor  $\mathcal{A}(\mathbf{x})$  is highly heterogeneous (possibly in a spatially complicated way). We assume that  $\mathcal{A}(\mathbf{x})$  is symmetric, uniformly positive definite and mildly anisotropic, i.e. 9  $\lambda_{\min}(\mathscr{A}(\mathbf{x})) \gtrsim \lambda_{\max}(\mathscr{A}(\mathbf{x}))$  uniformly in  $\mathbf{x}$ . We are particularly interested in the case 10 when the contrast  $\max_{\mathbf{x},\mathbf{y}\in\Omega} \lambda_{\max}(\mathscr{A}(\mathbf{x}))/\lambda_{\max}(\mathscr{A}(\mathbf{y}))$  is large. Many examples of 11 this type arise in subsurface flow modelling or in material science. The space  $H_0^1(\Omega)$  12 is the usual Sobolev space of functions with vanishing trace on  $\partial\Omega$  and  $f\in H^{-1}(\Omega)$ . 13 For simplicity we assume for the remainder that  $\mathscr{A}(\mathbf{x}) = \alpha(\mathbf{x})I$ , i.e. a scalar diffusion 14 coefficient.

Let  $\mathscr{T}_h$  be a simplicial triangulation of  $\Omega$  and let (1) be discretised in  $V_h \subset H_0^1(\Omega)$ , 16 the space of continuous, piecewise linear FE functions with respect to  $\mathcal{T}_h$  that vanish 17 on  $\partial \Omega$ . For simplicity let  $\mathcal{I}_h$  be quasi-uniform. The a-orthogonal projection of u to 18  $V_h$  is denoted by  $u_h$ . In the usual nodal basis  $\{\varphi_i\}_{i=1}^n$  for  $V_h$ , the problem of finding  $u_h$  reduces to the  $n \times n$  linear system

$$A\mathbf{u} = \mathbf{b} \tag{2}$$

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with stiffness matrix  $A = (a(\varphi_i, \varphi_j))_{i,j=1}^n$ . Since the matrix A depends on  $\alpha$  only 21 through element averages, we can assume (w.l.o.g.) that  $\alpha$  is piecewise constant with 22 respect to  $\mathcal{T}_h$ . For simplicity we assume that  $\alpha$  is piecewise constant with respect to 23 some non-overlapping partitioning of  $\Omega$  into open, connected Lipschitz polyhedra 24 (polygons)  $\{\mathscr{Y}_m\}_{m=1}^{M}$  and set  $\alpha_m = \alpha|_{\mathscr{Y}_m}$ .

Especially for d=3 and for problems where  $\alpha$  varies on a small length scale 26  $\varepsilon \ll \operatorname{diam}(\Omega)$ , and thus the mesh size h needs to be very fine, multilevel itera- 27 tive solvers (multigrid, domain decomposition, etc.) are usually essential to solve 28

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this problem efficiently. Their scalability and robustness with respect to mesh re- 29 finement, as well as other discretisation parameters has been studied extensively. 30 Here we will focus on their robustness with respect to coefficient variation. We will 31 show that coefficient robustness is inherently linked to a judicious choice of coarse 32 space  $V_H$  (related to some coarse mesh  $\mathcal{T}_H$  with resolution H). If  $\varepsilon \gtrsim H$  and if we can choose a coarse mesh such that all coefficient jumps are aligned with the mesh, 34 then the coefficient robustness of standard coarse spaces has been analysed in the 35 1990s (cf. [3, 4, 10, 16, 21, 22, 25] and the references therein). For certain methods 36 the robustness may depend on the quasi-monotonicity of the coefficient with respect 37 to the coarse mesh (in the sense of [3]). Substructuring-type ("exotic") coarse spaces 38 are usually used to achieve uniform coefficient robustness. A certain amount of ro- 39 bustness can be recovered for standard piecewise linear coarse spaces by using the 40 multilevel solver as a preconditioner within CG (e.g. [24]). The key tool in all these 41 analyses is the weighted  $L_2$ -projection of Bramble and Xu [1]. It requires a piece- 42 wise constant weight with respect to the coarse mesh, an assumption that is often far 43 too stringent in real applications. We want to move away from this and crucially here 44 make no assumptions that the underlying coarse grids resolve the coefficients.

A lot of effort in the last 25 years has gone into the development of algebraic 46 methods to construct coarse spaces, such as algebraic multigrid (AMG), rather than 47 analytic/geometric ones. It has been confirmed numerically that AMG methods are 48 in practice robust to coefficient variation when applied to (2) (i.e. the number of 49 iterations is unaffected), and they are therefore extremely popular. However, they are 50 built on several heuristics and so a rigorous analysis of their coefficient-robustness 51 is difficult (see [22] for a review of existing theoretical results). Nevertheless, the 52 key principle of these algebraic coarse spaces, namely energy minimisation [11], 53 also underlies many other coarse spaces. To obtain rigorous coefficient-independent 54 convergence results we will need to work in the following energy and weighted  $L_2$ - 55 norms on  $D \subset \Omega$ ,  $||v||_{a,D} = \int_D \alpha |\nabla v|^2$  and  $||v||_{0 \alpha D} = \int_D \alpha v^2$ 

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respectively. When  $D = \Omega$  we will usually not specify the domain explicitly.

A convenient framework to analyse most multilevel methods is the Schwarz or 59 subspace correction framework [21, 23]. We restrict attention to the two-level over- 60 lapping additive Schwarz method and focus on the robustness of various coarse 61 spaces for this method. We review some recent papers on the topic mainly by the 62 author (jointly with co-workers), as well as by Efendiev et al. All the results ap- 63 ply immediately also to multiplicative, hybrid and non-overlapping versions of the 64 Schwarz method (see [9, 18] for some explicit comments). Many of the results can 65 be extended to a multilevel theory [5, 18].

# 2 Schwarz Framework and Abstract Coarse Spaces

Let us assume that  $\{\Omega_k\}_{k=1}^K$  is an overlapping partitioning of  $\Omega$  and let  $\Omega_k^{\circ}$  be the overlap of subdomain  $\Omega_k$ , i.e. the set of points  $\mathbf{x} \in \Omega_k$  that are contained in at least one

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other subdomain. We assume that  $\mathcal{T}_h$  is aligned with this partitioning. Furthermore, 70 let  $\{\chi_k\}_{k=1}^K \subset V_h$  be an arbitrary partition of unity (POU) of FE functions subordinate 71 to  $\{\Omega_k\}_{k=1}^K$  such that  $\|\chi_k\|_{\infty} \lesssim 1$  and  $\|\nabla \chi_k\|_{\infty} \leq \delta_k^{-1}$ , for all  $k=1,\ldots,K$ . Note that 72 (due to quasi-uniformity of  $\mathscr{T}_h$ ) we always have  $\delta_k \gtrsim h$ , and there is a partition of 73 unity such that  $\delta_k$  is proportional to the (minimal) width of  $\Omega_k^{\circ}$ . We assume as usual 74 that each point  $\mathbf{x} \in \Omega$  is contained in at most  $N_0$  subdomains (finite covering).

We associate with each  $\Omega_k$  the space  $V_k = \{v \in V_h : \operatorname{Supp}(v) \subset \overline{\Omega}_k\}$  and assume 76 that we have an additional coarse space

$$V_0 = V_H = \text{span}\{\Phi_i \in V_h : j = 1, \dots, N\} \subset V_h.$$
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Let  $\omega_i = \operatorname{interior}(\operatorname{Supp}(\Phi_i))$  and set  $H_i = \operatorname{diam}(\omega_i)$ . Then  $H = \max_i H_i$  is the coarse mesh size associated with  $V_H$ . 80

The two-level additive Schwarz preconditioner is now simply

$$M_{\text{AS}}^{-1} = R_0^T A_0^{-1} R_0 + \sum_{k=1}^K R_k^T A_k^{-1} R_k \quad \text{with} \quad A_k = R_k A R_k^T.$$

 $R_k$  is the matrix representation of a restriction operator from V to  $V_k$ : the simple 83 injection operator for  $k \ge 1$ , and for k = 0 induced by the coarse space basis  $\{\Phi_i\}_{i=1}^N$ so that the coarse space stiffness matrix is  $A_0 = (a(\Phi_i, \Phi_\ell))_{i,\ell}^N$ . 85

The following result can be proved in the same way as [18, Theorem 2.5]. Since 86 it is instructive, we give an outline of the proof. 87

**Theorem 1.** If there exists an operator  $\Pi: V_h \to V_0$  such that for all  $v \in V_h$ 88

$$\|\Pi v\|_a^2 \le C_1 \|v\|_a^2$$
 and  $\sum_{k=1}^K \|(v - \Pi v)\nabla \chi_k\|_{0,\alpha}^2 \le C_2 \|v\|_a^2$ , (3)

then  $\kappa(M_{AS}^{-1}A) \lesssim C_1 + C_2$ . The hidden constant depends on  $N_0$ .

*Proof.* Let  $v_0 = \Pi v$  be such that (3) holds and choose  $v_k = I_h(\chi_k(v - v_0))$ , where  $I_h$  90 is the standard nodal interpolant on  $V_h$ . This interpolant is stable for all piecewise 91 quadratic functions in the energy norm and in the weighted  $L_2$ -norm (independently 92 of  $\alpha$ ) (cf. [18, Lemma 2.3]), and so we get

$$\sum_{k=0}^{K} \|v_k\|_a^2 \lesssim \|v_0\|_a^2 + \sum_{k=1}^{K} \|\chi_k(v - v_0)\|_a^2$$
$$\lesssim \|v_0\|_a^2 + \sum_{k=1}^{K} \|\chi_k\|_{\infty}^2 \|v - v_0\|_{a,\Omega_k}^2 + \|(v - v_0)\nabla\chi_k\|_{0,\alpha}^2.$$

Now, the boundedness of the POU functions, the finite cover assumption, as well as 94 (3) lead to the stability estimate  $\sum_{k=0}^{K} \|v_k\|_a^2 \lesssim (C_1 + C_2) \|v\|_a^2$ . Since  $v = \sum_{k=0}^{K} v_k$ , the 95 result follows from the abstract Schwarz theory (cf. [21]). 96

This result shows the importance of the choice of coarse space. Provided we have 97 a good coarse space approximation in the weighted L<sub>2</sub>-norm that is moreover stable 98 in the energy norm, independently of variations in  $\alpha$ , then the bound on the condition 99 number for two-level additive Schwarz is also robust with respect to these variations. 100 Note that it is crucial to use the weighted  $L_2$  and the energy norm here to achieve 101

coefficient-robustness, and that we only require weak  $L_2$ -approximation in regions 102 where  $\nabla \chi_k \neq 0$ .

Several approaches have been studied in [2, 5–9, 17–19] to provide constants 104 in (3) that are independent of  $\alpha$  (or at least of the contrast in  $\alpha$ ) for various coarse 105 spaces. However, in most cases the constants are not independent of  $\frac{H}{\varepsilon}$ , where  $\varepsilon$ is the minimal length scale at which lpha varies in the regions where  $abla \chi_k 
eq 0$ , So 107 unfortunately in general, to be also independent of  $\frac{H}{s}$ , restrictions on the coarse mesh 108 size are needed, at least locally.

Let us discuss the assumptions (3) a bit further. Let  $\Pi v = \sum_{i} f_{i}(v) \Phi_{i}$ , where  $f_i: V_h \to \mathbb{R}$  is a suitable functional. Then

$$\|\Pi v\|_a = \|\sum_j f_j(v) \Phi_j\|_a \le \sum_j |f_j(v)| \|\Phi_j\|_a.$$

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We see that a set of coarse basis functions with bounded energy (independent of  $\alpha$ ) 113 is beneficial. The first approaches in [8, 9, 17] attacked this target directly and aimed 114 at bounding  $\|\Phi_i\|_a$ . In that case, it suffices to use the standard quasi-interpolant. 115 Alternatively, a weighted quasi-interpolant with  $f_j(v) = \int_{\omega_i} \alpha v / \int_{\omega_i} \alpha$  can be used. 116 For certain (locally quasi-monotone) coefficients  $\alpha$  this leads to a constant  $C_1$  that is independent of the contrast in  $\alpha$ , even if the energy of the basis functions is not 118 bounded (see below).

Similar comments can be made about the second assumption in (3). Note that

$$\|(v - \Pi v)\nabla \chi_{k}\|_{0,\alpha}^{2} \leq \begin{cases} \|\alpha|\nabla \chi_{k}|^{2}\|_{\infty}\|v - \Pi v\|_{0,\Omega_{k}^{\circ}}^{2}, & \text{or} \\ \|\nabla \chi_{k}\|_{\infty}^{2}\|v - \Pi v\|_{0,\alpha,\Omega_{k}^{\circ}}^{2}. \end{cases}$$

We can either try to choose a partition of unity  $\{\chi_k\}$  such that  $\|\alpha|\nabla\chi_k\|^2\|_{\infty}$  is bounded 122 independently of  $\alpha$ , which is again related to energy minimisation, or we can try to 123 bound  $||v - \Pi v||_{0,\alpha,\Omega_0^s}$  directly. As above, it is possible for certain (locally quasimonotone) coefficients to achieve this and to obtain a constant  $C_2$  that does not depend on the contrast in  $\alpha$  (see below).

When the coefficient is not locally quasi-monotone, then it is in general necessary 127 to enrich the coarse space, by either refining the coarse mesh locally, or by choosing 128 more than one basis function per subdomain  $\Omega_k$ , with the key tool to achieve coarse 129 space robustness being again energy minimisation.

To highlight some of the key issues we will use a number of representative model 131 problems shown in Fig. 1. For the rest of the paper, we will only focus on cases, such 132 as Fig. 1c, h, where it is impossible or impractical that the subdomains  $\{\Omega_k\}$  and 133 the supports  $\{\omega_i\}$  of the coarse basis functions resolve the coefficient jumps. The 134 resolved cases in Fig. 1a, b have already been studied extensively, see e.g. [3, 4, 10, 135] 16, 21, 22, 24, 25].

# 3 Analysis of Coefficient–Robustness

We present three possible approaches to try and prove coefficient robustness rigor- 138 ously and thus to design robust coarse spaces. For simplicity, we assume that for 139 each j = 1, ..., N, there exists a k = 1, ..., K such that  $\omega_j \subset \Omega_k$ . 140

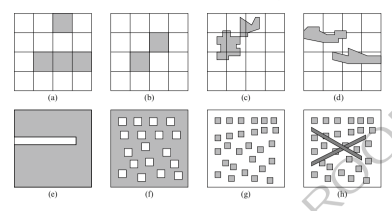


Fig. 1. Typical coefficient distributions (a) resolved; (b) not quasi-monotone; (c) neither quasimonotone nor resolved; (d) channelised; (e) flow barriers; (f) low permeability inclusions; (g) high permeability inclusions; (h) high permeability inclusions and channels

## 3.1 Standard Quasi-interpolant and Energy Minimisation

The first approach makes use of the standard quasi-interpolant

$$\Pi v = \sum_{j=1}^{N} \overline{v}_{\omega_j} \Phi_j, \quad \text{where} \quad \overline{v}_{\omega_j} = \frac{1}{|\omega_j|} \int_{\omega_j} v.$$

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Let  $\{\Phi_j\}_{i=1}^N$  be a set of bounded coarse basis functions that form a partition 144 of unity, except in a boundary layer of width  $\mathcal{O}(H)$  near  $\partial\Omega$ . Since each support 145  $\omega_i \subset \Omega_k$ , for some k, the supports have finite overlap. The constants  $C_1$  and  $C_2$  can 146 now be bounded independent of the contrast in  $\alpha$ , if either

$$\gamma_{2}(\alpha,\{\Phi_{j}\}) = \max_{j=1}^{N} H_{j}^{2-d} \|\Phi_{j}\|_{a}^{2} \quad \text{and} \quad \gamma_{\infty}(\alpha,\{\chi_{k}\}) = \max_{k=1}^{K} \delta_{k}^{2} \|\alpha^{1/2} \nabla \chi_{k}\|_{\infty}^{2}$$

(the so-called coarse space and partitioning robustness indicators) can be bounded 149 independent of  $\alpha$ , for some choice of the partition of unity  $\{\chi_k\}_{k=1}^K$  subordinate to 150  $\{\Omega_k\}_{k=1}^K$  (cf. [8]), or if  $\gamma_{\infty}(\alpha, \{\Phi_j\})$  can be bounded independent of  $\alpha$  (cf. [17]). 151 As mentioned above, this leads to the aim to construct coarse basis functions with 152 minimal or bounded energy. It is also at the heart of matrix-dependent prolongation 153 operators in multigrid methods.

For certain binary coefficient distributions, e.g. for high-permeability inclusions 155 in a low-permeability medium as depicted in Fig. 1g, it was then possible in [8] to 156 show (rigorously) under the assumption  $lpha \gtrsim 1$  that multiscale FEs (w.r.t. some coarse 157 mesh  $\mathcal{T}_H$ ) can provide such a basis  $\{\Phi_i\}$ , and that the indicators can be bounded 158 independent of the contrast in  $\alpha$ . However, they depend on  $H/\varepsilon$ , where  $\varepsilon$  is the 159 minimum width of any island/gap.

Similarly, it was possible in [17] to show (again assuming  $\alpha \gtrsim 1$ ) that aggregation based on a strong connection criterion (originally designed for AMG methods) 162

leads to a coarse basis  $\{\Phi_i\}$  for which the robustness indicators can be bounded independent of the contrast in  $\alpha$ . Here the bounds depend on H/h, since the overlap 164 between any two supports is only  $\mathcal{O}(h)$ .

However, this approach to analyse robustness fails even for the simpler, reverse situation of a high-permeability medium with low-permeability inclusions (e.g. 167 Fig. 1f), since in this case  $\gamma_i(\alpha, \{\Phi_i\})$  and  $\gamma_{\infty}(\alpha, \{\Phi_i\})$  depend on the contrast in  $\alpha$  168 for any choice of  $\{\Phi_i\}$ . Clearly a different quasi-interpolant  $\Pi$  is needed in general. 169

# 3.2 Weighted Quasi-interpolant and Poincaré's Inequality

The next approach to try to prove the assumptions in Theorem 1 makes use of the 171 weighted quasi-interpolant 172

$$\Pi v = \sum_{j=1}^{N} \overline{v}_{\omega_{j}}^{\alpha} \Phi_{j}, \quad \text{where} \quad \overline{v}_{\omega_{j}}^{\alpha} = \int_{\omega_{j}} \alpha v / \int_{\omega_{j}} \alpha.$$

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We describe this approach for one of the simplest coarse spaces, the piecewise 174 linear one. The following is taken from [18] (see also [6] for earlier results). Let 175  $V_H$  be the continuous, piecewise linear FE space associated with a shape-regular 176 simplicial triangulation  $\mathcal{T}_H$  of  $\Omega$ , such that  $\mathcal{T}_h$  is a refinement of  $\mathcal{T}_H$ . The functions  $\{\Phi_j\}_{j=1}^N$  are the standard nodal basis for  $V_H$ . For simplicity, we assume that  $\{\Omega_k\}_{k=1}^K=\{\omega_j\}_{j=1}^N$ , and choose  $\chi_k=\Phi_k$  (suitably modified near  $\partial\Omega$ ), so that the assumptions on  $\{\chi_k\}$  are satisfied with  $\delta_k \sim H_k$ .

The key observation in [18] is now that one further assumption suffices to fully 181 describe the dependency of the constants  $C_1$  and  $C_2$  in (3) on  $\alpha$ : 182

**Assumption 1** Let  $\omega_T = \bigcup_{\{k: \omega_k \cap T \neq \emptyset\}} \omega_k$  and  $H_T = \text{diam}(\omega_T)$ , for  $T \in \mathscr{T}_H$ , and assume that there exists a  $C_T^* > 0$  such that, for all  $v \in V_h$ , either 183 184

$$\inf_{c \in \mathbb{R}} \int_{\omega_T} \alpha (v - c)^2 d\mathbf{x} \lesssim C_T^* H_T^2 \int_{\omega_T} \alpha |\nabla v|^2 d\mathbf{x}, \quad \text{or} \quad (4)$$

$$\inf_{c \in \mathbb{R}} \int_{\omega_T} \alpha (v - c)^2 d\mathbf{x} \lesssim C_T^* H_T^2 \int_{\omega_T} \alpha |\nabla v|^2 d\mathbf{x}, \quad \text{or}$$

$$\partial \omega_T \cap \partial \Omega \neq \emptyset \quad \text{and} \quad \int_{\omega_T} \alpha v^2 d\mathbf{x} \lesssim C_T^* H_T^2 \int_{\omega_T} \alpha |\nabla v|^2 d\mathbf{x}.$$
(5)

**Proposition 1.** Let Assumption 1 hold. Then 
$$C_1 + C_2 \lesssim C^* = \max_{T \in \mathscr{T}_H} C_T^*$$
.

*Proof.* Let  $v \in V_h$  and  $v_0 = \sum_{j=1}^N \overline{v}_{\omega_j}^{\alpha} \Phi_j$ . By the Cauchy-Schwarz inequality we have 187  $|\bar{v}_{\omega_i}^{\alpha}|^2 \leq \int_{\omega_i} \alpha v^2 / \int_{\omega_i} \alpha$ , and so, using the fact that  $\Phi_j \leq 1$ , 188

$$\int_T lpha v_0^2 \leq \sum_{j:\omega_j \cap T 
eq \emptyset} rac{\int_{\omega_j} lpha v^2}{\int_{\omega_j} lpha} \int_T lpha \Phi_j^2 \leq \int_{\omega_T} lpha v^2,$$
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which also implies  $\int_T \alpha (v - v_0)^2 \lesssim \int_{\omega_T} \alpha v^2$ . Now, multiplying the left hand side by  $|\nabla \chi_k|_T^2$  (which is a constant  $\sim H_T^{-2}$ ) and summing over  $k \ge 1$ , we get 191

$$\sum_{k=1}^{K} \| (v - v_0) \nabla \chi_k \|_{0,\alpha,T}^2 \lesssim H_T^{-2} \int_{\omega_T} \alpha v^2.$$
 (6)

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If  $\{\Phi_i\}$  forms a partition of unity on all of  $\omega_T$  (i.e. if  $\partial \omega_T \cap \partial \Omega = \emptyset$ ), we can replace v in (6) by  $\hat{v} = v - c$ , for any  $c \in \mathbb{R}$ , without changing the integral on the left hand side. Otherwise we set  $\hat{v} = v$ . In both cases, by Assumption 1 194

$$\int_{\omega_T} \alpha \hat{v}^2 \lesssim C_T^* H_T^2 \int_{\omega_T} \alpha |\nabla v|^2. \tag{7}$$

Combining (6) and (7) and summing over all  $T \in \mathcal{T}_H$  gives the bound for  $C_2$ . The bound for  $C_1$  can be established in a similar way (cf. [18, Lemma 4.1]).

Assumption 1 postulates the existence of a discrete weighted Poincaré/ Friedrichs-type inequality on each  $\omega_T$ . It always holds, but in general the constants  $C_T^*$ will not be independent of  $\alpha|_{\omega_T}$  and  $H_T/h$ . As described in detail in [18, Sect. 3] 199 (see also [13–15]), to obtain independence of  $\alpha$ , we require a certain local quasi- 200 monotonicity of  $\alpha$  on each of the regions  $\omega_T$ .

Weighted Poincaré Inequalities. Let us consider a generic coarse element  $T \in \mathcal{T}_H$  202 and define the following subsets of  $\omega_T$  where  $\alpha$  is constant: 203

$$\omega^m = \omega_T \cap \mathscr{Y}_m, \qquad m = 1, ..., M.$$

By  $\mathscr{I}_T \subset \{1, ..., M\}$  we denote the index set of all regions  $\omega^m$  that are non-empty. 205 Let us assume w.l.o.g. that each of these subregions is connected. We generalise 206 now the notion of quasi-monotonicity coined in [3] by considering the following 207 three (two) directed combinatorial graphs  $\Gamma^{(k)} = (\mathbf{N}, \mathscr{E}^{(k)}), \ 0 \le k \le d-1$ , where 208  $\mathbf{N} = \{ \boldsymbol{\omega}^m : m \in \mathscr{I}_T \}$  and the edges are ordered pairs of vertices. We distinguish 209 between three (two) different types of connections.

**Definition 1.** Suppose that  $\gamma^{m,m_2} = \overline{\omega}^m \cap \overline{\omega}^{m_2}$  is a non-empty manifold of dimension 211 k, for  $0 \le k \le d-1$ . The ordered pair  $(\omega^m, \omega^{m_2})$  is an edge in  $\mathcal{E}^{(k)}$ , if and only if 212  $\alpha_m \lesssim \alpha_{m_2}$ . The edges in  $\mathcal{E}^{(k)}$  are said to be of type-k. 213

In addition, for  $1 \le k \le d - 1$ , we assume that

- meas $(\gamma^{m,m_2}) \sim \text{meas}(\omega^m \cup \omega^{m_2})^{k/d}$ , and
- $\gamma^{m,m_2}$  is sufficiently regular, i.e. it is a finite union of shape-regular k-dimensional 216 simplices of diameter  $\sim \text{meas}(\gamma^{m,m_2})^{1/k}$ . 217

Quasi-monotonicity is related to the connectivity in  $\Gamma^{(k)}$ . Let  $m_* \in \mathscr{I}_T$  be the 218 index of the region  $\omega^{m_*}$  with the largest coefficient:  $\alpha_{m_*} = \max_{m \in \mathscr{I}_T} \alpha_m$ . 219

**Definition 2.** The coefficient  $\alpha$  is type-k quasi-monotone on  $\omega_T$ , if there is a path in 220  $\Gamma^{(k)}$  from any vertex  $\omega^m$  to  $\omega^{m_*}$ . 221

The following lemma summarises the results in [13–15]. The existence of a 222 benign constant  $C_T^*$  that is independent of  $\alpha$  is directly linked to quasi-monotonicity, 223 the way in which  $C_T^*$  depends on  $H_T/h$  to the type. 224

**Lemma 1.** Let  $\omega_T \subset \mathbb{R}^d$ , d = 2,3. If  $\alpha$  is type-k quasi-monotone on  $\omega_T$ , then (4) 225 holds with 226

$$C_T^* = \begin{cases} 1, & \text{if } k = d - 1, \\ 1 + \log\left(\frac{H_T}{h}\right), & \text{if } k = d - 2, \\ \frac{H_T}{h}, & \text{if } k = d - 3. \end{cases}$$
 (8)

A similar result can also be established in the case where  $\partial \omega_K \cap \partial \Omega \neq \emptyset$ , i.e. the 227 case of Friedrichs inequality (5), see e.g. [18, Sect. 3] for details.

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Quasi-monotonicity is crucial. If the coefficient is not quasi-monotone, e.g. the 229 situation in Fig. 1d, then  $C^*$  cannot be bounded independent of  $\alpha$ . See [18, Exam-230] ple 3.1] for a counter example. If the coarse mesh is not adjusted in certain critical 231 areas of  $\Omega$ , then  $V_H$  is in general not robust. The numerical results in [18] show 232 that this is indeed the case and that quasi-monotonicity is necessary and sufficient. 233 However, a few simple adjustments suffice, namely  $\mathcal{T}_H$  has to be sufficiently fine in 234 certain "critical" areas of  $\Omega$ :

- 1. Choose  $H_T \leq \varepsilon_m$ , for all  $T \in \mathscr{T}_H$  that intersect a region  $\mathscr{Y}_m$  that is bordered by 236 two regions  $\mathscr{Y}_{m'}$  and  $\mathscr{Y}_{m''}$  with  $\alpha_{m'}\gg\alpha_m$  and  $\alpha_{m''}\gg\alpha_m$ . Here  $\varepsilon_m$  denotes the width of  $\mathscr{Y}_m$  at its narrowest point. This ensures that  $\alpha$  is quasi-monotone on all 238 regions  $\omega_T$  that intersect  $\mathscr{Y}_m$ .
- 2. Choose  $H_T \lesssim h$ , near any point or edge where  $\alpha$  is only type-(d-2) or type- 240 (d-3) quasi-monotone, i.e. near any cross point.

Usually a logarithmic growth  $C^* \sim \max_T \log(H_T/h)$  is acceptable, and so even re-242 gions where the coefficient is type-(d-2) quasi-monotone do not require any par- 243 ticular attention.

For an arbitrary piecewise constant coefficient function  $\alpha$  there will often only be 245 a relatively small (fixed) number of regions  $\omega_T$  where  $\alpha$  is not quasi-monotone (see 246 e.g. Fig. 1b, e). Therefore it is very easy to ensure through some local refinement of 247  $\mathcal{T}_H$  near these regions that  $C^* \sim 1$  (or  $C^* \sim \log(H/h)$ ). Note that crucially, this local 248 refinement does not mean that  $\mathcal{T}_H$  has to be aligned with coefficient jumps anywhere 249 in  $\Omega$ . The coarse grid merely has to be sufficiently fine in regions where  $\alpha$  is not 250 quasi-monotone. Ideas on how to adapt  $\mathcal{I}_H$  in such a way are suggested in [18].

"Exotic" coarse spaces. Substructuring-type ("exotic") coarse spaces (as suggested 252 in [3, 4, 16]) can be analysed in a similar way. Here the coarse basis functions are 253 constructed as a-harmonic extensions of face, edge or vertex "cut" functions associ- 254 ated with a non-overlapping decomposition  $\mathcal{T}_H$  of the domain. This decomposition 255 may be related to the overlapping partitioning  $\{\Omega_k\}$ , or it may come from a separate 256 coarse grid (not necessarily simplicial). If the coefficient does not vary along any of 257 the edges/faces of  $\mathcal{T}_H$ , then the space can be analysed like the piecewise linear one 258 above, using in addition the energy minimising property of the a-harmonic exten- 259 sion (cf. [14]). If the coefficient does vary along an edge/face, then special weighted 260 Poincaré inequalities for functions with vanishing weighted averages across edges/- 261 faces are required. These have recently been introduced in the context of FETI-DP 262 methods in [12], which also analyses the robustness of the "cut" functions. An ex- 263 plicit analysis in the context of overlapping Schwarz does not yet exist.

#### 3.3 Abstract Minimisation with Functional Constraints

An alternative to refining the coarse mesh in regions where  $\alpha$  is not type-(d-1) 266 or type–(d-2) quasi-monotone, is to associate more than one basis function (with 267 possibly identical supports) with each subdomain  $\Omega_k$ . Let

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$$V_0 = \text{span}\{\Phi_{k,j} = I_h(\chi_k \Psi_{k,j}) : j = 1, \dots, N_k, k = 1, \dots, K\},$$

where  $\Psi_{k,j}$ ,  $j=1,\ldots,N_k$ , are suitable FE functions in  $V_h(\overline{\Omega}_k)$  (that do not vanish on 270  $\partial \Omega_k$ ) such that the functions  $\{\Phi_{k,j}\}\subset V_h$  are linearly independent. Good choices for 271 the functions  $\Psi_{k,j}$  are the lowest modes of local eigenproblems, or more generally, 272 energy minimising functions that satisfy suitable constraints. The following analysis 273 is from [19] (see [2, 7] for related work). 274

In particular, let us assume that, for every  $\Omega_k$ , we have a collection of linear 275 functionals  $\{f_{k,j}\}_{i=1}^{N_k} \subset V_h(\overline{\Omega}_k)'$  and let 276

$$\Psi_{k,j} = \arg\min_{v \in V_h(\overline{\Omega}_k)} |v|_a^2$$
, subject to  $f_{k,l}(\Psi_{k,j}) = \delta_{jl}$   $j,l = 1,...,N_k$ . (9)

Now, for any  $v \in V_h$ , choose the following quasi-interpolant

$$\Pi v = \sum_{k=1}^{K} I_h \left( \chi_k \Pi_{\Omega_k} v \right), \quad \text{where} \quad \Pi_{\Omega_k} v = \sum_{j=1}^{N_k} f_{k,j} (v|_{\Omega_k}) \Psi_{k,j},$$

i.e. a linear combination of the basis functions  $\Phi_{k,j}$  with weights  $f_{k,j}(v|_{\Omega_k})$ . Then 279 the bounds on  $C_1$  and  $C_2$  in Theorem 3 depend only on the stability and on the local 280  $L_2$ -approximation properties of  $\Pi_{\Omega_k}$  on each  $\Omega_k$ .

**Theorem 1.** For all 
$$k = 1, ..., K$$
 and for all  $v \in V_h(\overline{\Omega}_k)$ , let

$$\|\Pi_{\Omega_k} v\|_{a,\Omega_k}^2 \le \|v\|_{a,\Omega_k}^2 \text{ and } \|v - \Pi_{\Omega_k} v\|_{0,\alpha,\Omega_k}^2 \lesssim \text{diam}(\Omega_k)^2 \|u\|_{a,\Omega_k}^2.$$
 (10)

Then 
$$C_1 = \mathcal{O}(1)$$
 and  $C_2 \lesssim (\operatorname{diam}(\Omega_k)/\delta_k)^2$ .

Note that the minimisation problems in (9) are local to each subdomain. There are 285 suitable choices for the functionals  $f_{k,i}$  that guarantee (10) and that lead to practical 286 algorithms to construct the functions  $\Psi_{k,j}$ ,  $j = 1, ..., N_k$ : 287

•  $f_{k,j}(v) = (\Psi_{k,j}, v)_{0,\alpha,\Omega_k}$  where  $\Psi_{k,j}$  is the jth eigenfunction corresponding to the variational eigenproblem: Find  $\eta \in V_h(\overline{\Omega}_k)$  and  $\lambda \geq 0$ , such that 289

$$a(\eta, w) = \lambda(\eta, w)_{0,\alpha,\Omega_k}, \quad \text{for all} \quad w \in V_h(\overline{\Omega}_k).$$
 (11)

This has first been suggested and analysed in [7].

- $f_{k,j}(v) = (\Psi_{k,j}, v)_{0,\alpha,\partial\Omega_k}$  where  $\Psi_{k,j}$  is the *j*th eigenfunction corresponding to a 291 variational eigenproblem similar to (11), but with  $(\eta, w)_{0,\alpha,\partial\Omega_k}$  instead of 292  $(\eta, w)_{0,\alpha,\Omega_k}$  on the right hand side of (11), i.e. an eigenproblem of Steklov- 293 Poincaré type. This has been analysed in [2]. 294
- $f_{k,j}(v) = \overline{v}_{D_{k,j}}^{\alpha}$  where  $\{D_{k,j}\}_{j=1}^{N_k}$  is a suitable non-overlapping partitioning of  $\Omega_k$  295 such that the weighted Poincaré inequality (4) holds on each  $D_{k,j}$  (e.g.  $D_{k,j}$  =  $\Omega_k \cap \mathscr{Y}_i$ ). The construction of  $\{\Psi_{k,i}\}$  requires the solution of  $N_k$  local saddle 297 point systems and was suggested and analysed in [19]. 298

It has been shown in [2, 7] how (10) can be proved (directly) in the first two cases, es- 299 sentially based on the observation that the coarse space consists of the lowest modes 300 corresponding to the operator pencil associated to the energy and to the weighted 301  $L_2$ -norm. But the assumptions can be proved for a much wider class of functionals 302 using the following abstract approximation result in [19]. This result is related to the classical Bramble-Hilbert lemma.

Abstract Approximation Result. Consider an abstract symmetric and continuous 305 bilinear form  $a(\cdot,\cdot):V\times V\mapsto \mathbb{R}$ , as well as a collection of linear functionals 306  $\{f_l\}_{l=1}^m \subset V', \text{ where } V \subset \mathcal{H} \text{ and } \mathcal{H} \text{ is a Hilbert space with norm } \|\cdot\|.$  We make 307 the following assumptions on  $a(\cdot,\cdot)$ , V,  $\mathcal{H}$ ,  $\|\cdot\|$  and  $\{f_l\}$ :

 $a(\cdot,\cdot)$  is positive semi-definite and defines a semi-norm  $[\cdot]_a$  on V, i.e. 309

$$|v|_a^2 = a(v, v) \ge 0$$
, for all  $v \in V$ .

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In addition, for  $v \in V$ , the expression  $\sqrt{\|v\|^2 + |v|_a^2}$  defines a norm on V.

Let  $c_q$  be a generic constant. For all  $\mathbf{q} \in \mathbb{R}^m$  there exists a  $v_{\mathbf{q}} \in V$  with **A2.** 

$$f_l(v_{\mathbf{q}}) = q_l$$
, and  $\|v_{\mathbf{q}}\| \lesssim c_q \|\mathbf{q}\|_{l^2(\mathbb{R}^m)}$ .

There are two constants  $c_a$  and  $c_f$  such that **A3.** 

$$||v||^2 \le c_a |v|_a^2 + c_f \sum_{l=1}^m |f_l(v)|^2$$
, for all  $v \in V$ . (12)

Now, as in the specific case above, define for all  $v \in V$ ,

$$\pi v = \sum_{l=1}^m f_l(v) \psi_l$$
, where  $\psi_l = \arg\min_{v \in V} |v|_a^2$ , subject to  $f_l(\psi_j) = \delta_{jl}$ .

Then the following inequalities hold; see [19, Theorem 3.3].

**Theorem 3.** Let Assumptions A1-A3 be satisfied. Then, for all  $u \in V$ : 318

$$|\pi u|_a \le |u|_a$$
 and  $||u - \pi u|| \le \sqrt{c_a} |u|_a$ . (13)

(Note that they are independent of the constants  $c_q$  and  $c_f$  in A2 and A3.) 319

In the specific case considered above, on an arbitrary subdomain  $\Omega_k$ , Assumption **A1** is naturally satisfied with  $\mathscr{H} = L_2(\Omega_k)$  and  $\|\cdot\| = \|\cdot\|_{0,\alpha,\Omega_k}$ . Assumption 321 A2 merely ensures that the linear functionals are linearly independent. Thus, the 322 question of coarse space robustness is reduced to verifying Assumption A3. For one 323 functional, i.e. for m = 1, this reduces to the weighted Poincaré inequality in Sect. 3.2 324 and to the restrictions on the coefficients made there. For more than one functional, 325 it opens the possibility to get coefficient robustness even in the case of non-quasi- 326 monotone coefficients, such as those depicted in Fig. 1b, d and even h. See [2, 7, 19] 327 for the complete analysis and some numerical experiments that confirm the robustness for the functionals defined on the previous page. See also [20] for a more recent 329 extension to systems of elliptic PDEs (such as linear elasticity). 330

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# **Multi-level Decompositions of Electronic Wave Functions**

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1 Introduction 7

The approximation of high-dimensional functions, whether they be given explicitly or implicitly as solutions of differential equations, represents one of the grand 9
challenges of applied mathematics. High-dimensional problems arise in many fields 10
of application such as data analysis and statistics, but first of all in the sciences. 11
One of the most notorious and complicated problems of this type is the Schrödinger 12
equation. The Schrödinger equation forms the basis of quantum mechanics and is 13
of fundamental importance for our understanding of atoms and molecules. It links 14
chemistry to physics and describes a system of electrons and nuclei that interact by 15
Coulomb attraction and repulsion forces. As proposed by Born and Oppenheimer in 16
the nascency of quantum mechanics, the slower motion of the nuclei is mostly separated from that of the electrons. This results in the electronic Schrödinger equation, 18
the problem to find the eigenvalues and eigenfunctions of the Hamilton operator

$$H = -\frac{1}{2} \sum_{i=1}^{N} \Delta_i - \sum_{i=1}^{N} \sum_{\nu=1}^{K} \frac{Z_{\nu}}{|\mathbf{x}_i - \mathbf{a}_{\nu}|} + \frac{1}{2} \sum_{\substack{i,j=1 \ i \neq i}}^{N} \frac{1}{|\mathbf{x}_i - \mathbf{x}_j|}.$$
 (1)

It acts on functions with arguments  $\mathbf{x}_1, \dots, \mathbf{x}_N \in \mathbb{R}^3$ , which are associated with the positions of the considered electrons. The  $\mathbf{a}_V$  are the fixed positions of the nuclei and the values  $Z_V$  the charges of the nuclei in multiples of the absolute electron charge.

The high dimensionality of the equation immediately rules out classical discretization methods for partial differential equations as numerical analysts are 24 familiar with. To overcome this curse of dimensionality, procedures like the Hartree-25 Fock method and its many variants and successors or density functional theory based 26 methods have been developed over the decades. They are used with much success and 27 form the basis of a steadily expanding branch of chemistry. See [6] for an overview 28 on the present state of the art in quantum chemistry, and [3, 10], and [11] for mathe-29 matically oriented expositions. All these methods suffer, however, either from a priori 30 modeling errors or from the fact that it is not clear how the accuracy can be system-31 atically improved without the effort truly exploding for larger numbers of electrons. 32

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It is therefore rather surprising that simple sparse grid-like multi-level expansions of 33 the electronic wave functions can be constructed whose convergence rate, measured 34 in terms of the number of basis functions involved, is independent of the number of 35 electrons and does not much differ from that for a two- or even one-electron system. 36 The purpose of this note is to explain these results and the effects behind them. For 37 details we refer to the references.

# 2 Regularity and Decay of the Wave Functions

The at least asymptotically, in relation to the high space dimension rapid conver- 40 gence of these expansions is based on very particular properties of the solutions of 41 the electronic Schrödinger equation: their regularity, that surprisingly increases with 42 the number of electrons, the decay behavior of their mixed derivatives, and their 43 antisymmetry enforced by the Pauli principle.

The solution space of the electronic Schrödinger equation is first the Hilbert 45 space  $H^1$  that consists of the square integrable functions

$$u: (\mathbb{R}^3)^N \to \mathbb{R}: (\mathbf{x}_1, \dots, \mathbf{x}_N) \to u(\mathbf{x}_1, \dots, \mathbf{x}_N)$$
 (2)

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with square integrable first-order weak derivatives; the dimension of their domain 47 increases with the number N of electrons. The norm  $\|\cdot\|_1$  on  $H^1$  is composed of the 48  $L_2$ -norm  $\|\cdot\|_0$  induced by the  $L_2$ -inner product and the  $L_2$ -norm of the gradient. In 49 the language of physics, the space  $H^1$  is the space of the wave functions for which 50 the total position probability remains finite and the expectation value of the kinetic 51 energy can be given a meaning. It can be shown that the second-order differential op- 52 erator (1) induces a bounded bilinear form on  $H^1$  that satisfies a Garding inequality. 53 The mathematically precise formulation of the eigenvalue problem is therefore the 54 corresponding weak form of the equation on the space  $H^1$ , the same kind of weak 55 form that one knows from the finite element method. The physically admissible so- 56 lutions are components  $u(\mathbf{x}) = \psi(\mathbf{x}, \boldsymbol{\sigma})$  of a full, spin-dependent wave function. By 57 the Pauli principle, they are therefore antisymmetric with respect to the exchange of 58 the positions  $\mathbf{x}_i$  of electrons of the same spin  $\sigma_i = \pm 1/2$ .

To describe the regularity properties of the eigenfunctions, we need to introduce 60 a scale of norms that are defined in terms of Fourier transforms. We first introduce 61 the polynomials

$$P_{\text{iso}}(\boldsymbol{\omega}) = 1 + \sum_{i=1}^{N} |\boldsymbol{\omega}_i|^2, \quad P_{\text{mix}}(\boldsymbol{\omega}) = \prod_{i=1}^{N} (1 + |\boldsymbol{\omega}_i|^2).$$
 (3)

The  $\boldsymbol{\omega}_i \in \mathbb{R}^3$  forming together the variable  $\boldsymbol{\omega} \in (\mathbb{R}^3)^N$  can be associated with the 63 momentums of the electrons. The expressions  $|\boldsymbol{\omega}_i|$  are their euclidean norms. The 64 norms describing the smoothness of the solutions are now given by

$$|||u|||_{\vartheta,m}^2 = \int P_{\text{iso}}(\boldsymbol{\omega})^m P_{\text{mix}}(\boldsymbol{\omega})^{\vartheta} |\widehat{u}(\boldsymbol{\omega})|^2 d\boldsymbol{\omega}.$$
(4)

They are defined on the Hilbert spaces  $H_{\rm mix}^{\vartheta,m}$  that consist of the square integrable 66 functions (2) for which these expressions remain finite. For nonnegative integer 67 values m and  $\vartheta$ , the norms measure the  $L_2$ -norm of weak partial derivatives. The 68 parameter m measures the isotropic smoothness that does not distinguish between 69 different directions, and the parameter  $\vartheta$  the mixed smoothness in direction of the 70 three-dimensional coordinate spaces of the electrons. The spaces  $L_2$  and  $H^1$  are special cases of such spaces.

It has been proved in [12] and [13] that the physically admissible eigenfunctions u 73 of the electronic Schrödinger operator (1) are at least contained in  $H_{\rm mix}^{\vartheta,1}$  for  $\vartheta=1/2$ . 74 Recently we were able to improve this result substantially. We have shown in [9] that 75 the eigenfunctions u of the electronic Schrödinger operator are, independent of their 76 symmetry properties, contained in

$$H_{\text{mix}}^{1,0} \cap \bigcap_{\vartheta < 3/4} H_{\text{mix}}^{\vartheta,1}. \tag{5}$$

The bound 3/4 is optimal and can, except for special cases, neither be reached nor 78 improved further. The proof is based on a representation of the eigenfunctions that 79 has been derived in [15] and for the two-electron case in [1]. It has been shown in 80 [15] that the eigenfunctions can be written as products

$$u(\mathbf{x}) = \exp\left(\sum_{i < j} \phi(\mathbf{x}_i - \mathbf{x}_j)\right) v(\mathbf{x})$$
 (6)

of more regular functions  $v \in H^{1,1}_{mix}$  and a universal factor that covers their singularities. This kind of splitting can be traced back to the work of Hylleraas [8] in the early years of quantum mechanics. It has been used in [4] and [7] to study the Hölder regularity of the eigenfunctions. There is a lot of freedom in the choice of the function  $\phi$ . 85 It needs only to be of the form

$$\phi(\mathbf{x}) = \widetilde{\phi}(|\mathbf{x}|), \quad \widetilde{\phi}'(0) = \frac{1}{2}, \tag{7}$$

where  $\phi:[0,\infty)\to\mathbb{R}$  is an infinitely differentiable function behaving sufficiently 87 well at infinity. The regularity is therefore determined by that of the explicitly known 88 factor from (6) that describes the behavior of the solutions at the singular points of 89 the electron-electron interaction potential.

The splitting (6) is of independent interest since it is obviously possible to obtain 91 better convergence rates for the regular part of the solutions than for the solutions 92 themselves. We will restrict ourselves, however, here to the direct approximation 93 of the eigenfunctions. The domain of the eigenfunctions is infinitely extended. The 94 eigenfunctions are, however, strongly localized. It is known for a long time that an 95 eigenfunction u for an eigenvalue below the ionization threshold of the given atom or 96 molecule decays exponentially in the  $L_2$ -sense. That means there is a constant  $\gamma > 0$  97 such that the function 98

$$\mathbf{x} \to \exp\left(\gamma \sum_{i=1}^{N} |\mathbf{x}_i|\right) u(\mathbf{x}),$$
 (8)

is square integrable. This constant depends on the distance of the eigenvalue under 99 consideration to the bottom of the essential spectrum. More details and references to 100 the literature can be found in [14]. It has been shown in [15] that these exponentially 101 weighted eigenfunctions admit the same kind of representation (6) as the eigenfunc- 102 tions themselves. Thus they share with them the described regularity properties [9]. 103 The convergence analysis is based on this observation.

# 3 Sparse Grids and Antisymmetry

To explain the meaning of these results for the approximation of the solutions of the Schrödinger equation, we consider a simple model problem, the approximation of 107 functions u of the variables  $x_1, \ldots, x_d$  that are odd and  $2\pi$ -periodic in every coordinate direction on the cube  $Q = [0, \pi]^d$  by tensor products

$$\phi(\mathbf{k}, \mathbf{x}) = \prod_{i=1}^{d} \phi_{k_i}(x_i)$$
 (9)

of the one-dimensional trigonometric polynomials

$$\phi_{k_i}(\xi) = \sqrt{\frac{2}{\pi}} \sin(k_i \xi) \tag{10}$$

labeled by the components  $k_i = 1, 2, \dots$  of the multi-indices k. Our presentation 111 closely follows [14]. Functions of the given kind that are square integrable over Q 112 can be expanded into a multivariate Fourier series 113

$$u(\mathbf{x}) = \sum_{\mathbf{k}} \widehat{u}(\mathbf{k}) \phi(\mathbf{k}, \mathbf{x}), \tag{11}$$

where the expansion coefficients are given by

$$\widehat{u}(\mathbf{k}) = \int_{Q} u(\mathbf{x})\phi(\mathbf{k}, \mathbf{x}) \, d\mathbf{x}. \tag{12}$$

We measure the speed of convergence of this series in the sense of the  $L_2$ -norm which 115 reads in terms of the expansion coefficients 116

$$||u||_0^2 = \sum_{\mathbf{k}} |\widehat{u}(\mathbf{k})|^2. \tag{13}$$

The speed of convergence of the series is therefore determined by the speed with which the expansion coefficients decay. Assume that all partial derivatives of u of 118 order s exist and are square integrable. This implies that

$$|u|_s^2 = \sum_{\mathbf{k}} |\mathbf{k}|^{2s} |\widehat{u}(\mathbf{k})|^2$$
 (14)

remains finite, where  $|\mathbf{k}|$  is defined by

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$$|\mathbf{k}|^2 = \sum_{i=1}^d k_i^2. \tag{15}$$

Consider now the finite part  $u_{\varepsilon}$  of the series (11) that extends over the multi-indices **k** 121 inside the ball of radius  $1/\varepsilon$  around the origin, for which

$$|\mathbf{k}| < \frac{1}{\varepsilon}.\tag{16}$$

Due to the orthonormality of the functions (9),  $u_{\varepsilon}$  is the best approximation of u by 12: a linear combination of the selected basis functions. It holds

$$\|u - u_{\varepsilon}\|_{0}^{2} \leq \varepsilon^{2s} \sum_{\mathbf{k}} |\mathbf{k}|^{2s} |\widehat{u}(\mathbf{k})|^{2} = \varepsilon^{2s} |u|_{s}^{2}.$$
 (17)

The number n of these basis functions grows like

$$n \sim \frac{1}{\epsilon^d} \tag{18}$$

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as  $\varepsilon$  goes to zero. This is out of every reach for higher space dimensions d, the 126 curse of dimensionality. It can only be broken if one restricts oneself to a class of 127 functions whose smoothness increases sufficiently fast with the space dimension d. 128 At this place the mixed regularity comes into play. Consider functions u that possess 129 corresponding weak partial derivatives and set

$$|u|_{1,\text{mix}}^2 = \int_{\mathcal{O}} \left| \frac{\partial^d u}{\partial x_1 \dots \partial x_d} \right|^2 d\mathbf{x}$$
 (19)

or, in terms of the expansion coefficients,

$$|u|_{1,\text{mix}}^2 = \sum_{\mathbf{k}} \left( \prod_{i=1}^d k_i \right)^2 |\widehat{u}(\mathbf{k})|^2.$$
 (20)

Let  $u_{\varepsilon}^*$  be the function represented by the finite part of the series (11) that extends over the multi-indices **k** inside the hyperboloid given by

$$\prod_{i=1}^{d} k_i < \frac{1}{\varepsilon},\tag{21}$$

instead of the ball (16). The  $L_2$ -error can then be estimated as

$$||u - u_{\varepsilon}^*||_0 \le \varepsilon |u|_{1,\text{mix}} \tag{22}$$

and tends like  $\mathcal{O}(\varepsilon)$  to zero. The dimension n of the space spanned by the functions 135 (9) for which (21) holds, now increases, however, only like 136

$$n \sim |\log \varepsilon|^{d-1} \varepsilon^{-1}. \tag{23}$$

This shows that a comparatively slow growth of the smoothness can help to reduce 137 the complexity substantially, an observation that forms the basis of the sparse grid or 138 hyperbolic cross techniques; see [2] for an overview. Due to the presence of the logarithmic term, the applicability of such methods is, however, still limited to moderate 140 space dimensions.

The rescue comes from the symmetry properties of the wave functions enforced 142 by the Pauli principle. They represent a possibility to escape from this dilemma without forcing up the smoothness requirements further, which has first been noted by 144 Hackbusch [5]. Consider functions u that are antisymmetric with respect to the exchange of their variables, i.e., that

$$u(\mathbf{P}\mathbf{x}) = \operatorname{sign}(\mathbf{P})u(\mathbf{x}) \tag{24}$$

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holds for all permutation matrices P. It is not astonishing that such symmetry properties are immediately reflected in the expansion (11). Let 148

$$\widetilde{\phi}(\mathbf{k}, \mathbf{x}) = \frac{1}{\sqrt{d!}} \sum_{\mathbf{P}} \operatorname{sign}(\mathbf{P}) \phi(\mathbf{k}, \mathbf{P}\mathbf{x})$$
 (25)

be the renormalized, antisymmetric parts of the functions (9), where the sums extend 149 over the d! permutation matrices **P** of order d. The antisymmetrized functions (25) 150 can be written as determinants

$$\frac{1}{\sqrt{d!}} \begin{vmatrix} \phi_{k_1}(x_1) & \dots & \phi_{k_d}(x_1) \\ \vdots & \ddots & \vdots \\ \phi_{k_1}(x_d) & \dots & \phi_{k_d}(x_d) \end{vmatrix}$$
(26)

and evaluated in this way. For the functions u in the given symmetry class, many 152 terms in the expansion (11) can be combined. It finally collapses into 153

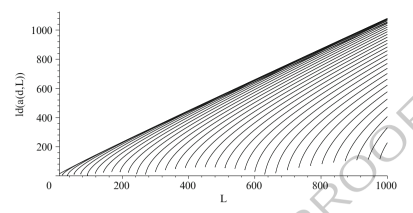
$$u(\mathbf{x}) = \sum_{k_1 > \dots > k_d} \left( u, \widetilde{\phi}(\mathbf{k}, \cdot) \right) \widetilde{\phi}(\mathbf{k}, \mathbf{x}), \tag{27}$$

where the expansion coefficients are the  $L_2$ -inner products of u with the corresponding functions (25). The number of basis functions needed to reach a given accuracy 155 is reduced by more than the factor d!, a very significant gain for larger dimensions d. 156

It remains to count the number of the sequences  $k_1 > k_2 > ... > k_d$  of natural 157 numbers that satisfy the condition (21) and with that also the number of basis func- 158 tion (25) needed to reach the accuracy  $\mathcal{O}(\varepsilon)$ . To study the asymptotic behavior of 159 the number of these sequences in dependence of the dimension d and the accuracy  $\varepsilon$ , 160 it suffices when we restrict ourselves to the case  $\varepsilon = 1/2^L$ , with positive integers L. 161 That is, we have to give bounds for the number of sequences  $k_1 > ... > k_d$  for which 162

$$\prod_{i=1}^{d} k_i \le 2^L. \tag{28}$$

The problem to estimate this number has to do with the prime factorization of integers. To simplify this problem, we group the numbers  $k_i$  into levels and decompose the space of the trigonometric polynomials correspondingly. Let



**Fig. 1.** The numbers  $a^*(L)$  and a(d,L) for d = 10, 15, 20, ..., 175

$$\ell(k_i) = \max \left\{ \ell \in \mathbb{Z} \mid 2^{\ell} \le k_i \right\}. \tag{29}$$

An upper bound for the number of these sequences is then the number a(d,L) of the sequences  $k_1 > k_2 > ... > k_d$  of natural numbers for which 167

$$\prod_{i=1}^{d} 2^{\ell(k_i)} \le 2^L. \tag{30}$$

The numbers a(d, L) can be calculated recursively; see [14] for details. A crude estimate yields a(d,L) = 0 if L+1 < d. Thus 169

$$a^*(L) := \max_{d \ge 1} a(d, L) = \max_{d \le L+1} a(d, L).$$
 (31)

Figure 1 shows, in logarithmic scale, how the a(d,L) behave compared to their joint 170 least upper bound  $a^*(L)$ . It becomes obvious from this picture that this upper bound exceeds the actual dimensions for larger d by many orders of magnitude, the more 172 the more the number d of variables increases. The joint least upper bound that is 173 independent of d for the number of the sequences  $k_1 > ... > k_d$  of natural numbers 174  $k_i$  for which (28) holds grows at least like  $\sim 2^L$  since already for the case d=1, 175 there are  $2^L$  such "sequences", namely those with values  $k_1 = 1, ..., 2^L$ . Figure 1 176 suggests conversely that the upper bound (31) for the number of these sequences 177 does not grow much faster than  $\sim 2^L$ . This is in fact the case since the number of the 178 decreasing infinite sequences  $k_1 \ge k_2 \ge k_3 \ge \dots$  of natural numbers for which

$$\prod_{i=1}^{\infty} 2^{\ell(k_i)} \le 2^L,\tag{32}$$

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with L a given nonnegative integer, is bounded by

$$\sum_{\ell=0}^{L} p(\ell) 2^{\ell},\tag{33}$$

where  $p(\ell)$  denotes the partition number of  $\ell$ , the number of possibilities of representing  $\ell$  as sum of nonnegative integers without regard to the order. To show this, we 182 observe that the number of these sequences is bounded by the number of sequences 183  $k_1, k_2, k_3, \ldots$  of natural numbers for which at least their levels  $\ell(k_1), \ell(k_2), \ldots$  decrease and that satisfy (32). We show that the expression (33) counts the number of 185 these sequences. Let the integers  $\ell_i = \ell(k_i)$  first be given. As there are  $2^{\ell_i}$  natural 186 numbers  $k_i$  for which  $\ell(k_i) = \ell_i$ , namely  $k_i = 2^{\ell_i}, \dots, 2^{\ell_i+1} - 1$ , there are

$$\prod_{i=1}^{\infty} 2^{\ell_i} = 2^{\ell}, \quad \ell = \sum_{i=1}^{\infty} \ell_i, \tag{34}$$

sequences  $k_1, k_2, k_3, \ldots$  for which the  $\ell(k_i)$  attain the prescribed values  $\ell_i$ . The problem thus reduces to the question how many decreasing sequences of nonnegative integers  $\ell_i$  exist that sum up to values  $\ell \leq L$ , i.e., for which

$$\sum_{i=1}^{\infty} \ell_i = \ell. \tag{35}$$

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This number is by definition the partition number  $p(\ell)$  of the nonnegative integer  $\ell$ . 191 Every sequence  $k_1 > k_2 > ... > k_d$  of natural numbers for which (28) holds can 192 obviously be expanded to an infinite, decreasing sequence  $k_1 \ge k_2 \ge k_3 \ge \dots$  of 193 natural numbers that satisfies the condition (32) by setting all  $k_i = 1$  for i > d. The sum (33) represents therefore also an upper bound for the number of these sequences. 195

The partition number plays a big role in combinatorics. Hardy and Ramanujan 196 have shown that it behaves asymptotically like

$$p(\ell) \sim \frac{\exp\left(\pi\sqrt{2\ell/3}\right)}{\ell} \tag{36}$$

as  $\ell$  goes to infinity. We conclude that the upper bound (31) for the number of determinants needed to reach an error  $\leq 2^{-L}|u|_{1,\text{mix}}$  behaves like 199

$$a^*(L) = (2^L)^{1+\delta(L)}, \quad 0 \le \delta(L) \le cL^{-1/2},$$
 (37)

where c is a constant that depends neither on L nor on the space dimension d or the 200 function u. Using the representation of  $a^*(L)$  from (31) and the recursively calculated 201 values a(d,L), the exponents  $1+\delta(L)$  can be calculated exactly. They decay for L 202 ranging from 10 to 1,000 monotonely from 1.406 to 1.079. For  $L = 100, 1 + \delta(L) = 203$ 1.204. In other words, the error tends faster to zero in the number n of determinants 204 than 205

$$\sim \frac{1}{n^{1-\vartheta}} \tag{38}$$

for any given  $\vartheta$  in the interval  $0 < \vartheta < 1$ . Not only does the convergence rate de-206 teriorate neither with the dimension nor the number of variables, it behaves asymp- 207 totically almost as in the one-dimensional case. Similar results hold for partially 208 antisymmetric functions as they occur in quantum mechanics.

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# 4 Eigenfunction and Wavelet Expansions

The constructions sketched in the previous section transfer to the more complicated case of the expansion of the solutions of the electronic Schrödinger equation into correspondingly antisymmetrized tensor products of three-dimensional Hermite functions or other eigenfunctions of three-dimensional Schrödinger-like operators as in [14] or wavelets as in [16]. Indeed, it finally turns out that the convergence rate measured in terms of the number of basis functions involved does not deteriorate with the number of electrons and comes close to that for the two- or even one-particle case. We do not explicate the partly technical details here but explain how one can utilize the intermediate smoothness of the exponentially weighted solutions (8) to obtain optimal convergence rates.

Let  $e^{\psi}$  be exponential factor in (8). The argumentation starts from functions v 221 whose exponentially weighted counterparts  $e^{\psi}v$  are located in  $H_{\text{mix}}^{1,1}$ , that is, have in 222 contrast to the solutions of the Schrödinger equation full mixed regularity. The essential observation is that the norm  $\|e^{\psi}v\|_{1,1}$  can be estimated by the sum of the 224 weighted  $L_2$ -norms  $\|e^{\psi}D^{\alpha}v\|_0$  of the involved derivatives  $D^{\alpha}v$  of v and vice versa. 225 This comes from the special structure of the function  $\psi$ . The norm  $\|e^{\psi}v\|_{1,1}$  measures therefore the exponentially weighted  $L_2$ -norms of the involved derivatives of v. 227 It is therefore reasonable to start from a sequence  $T_n: H^1 \to H^1$ ,  $n=1,2,\ldots$ , of 228 linear approximation operators that are uniformly  $H^1$ -bounded and to require that

$$\|\mathbf{v} - T_n \mathbf{v}\|_1 \lesssim n^{-q} \|\mathbf{e}^{\Psi} \mathbf{v}\|_{1,1}$$
 (39)

for all functions  $v \in H^1$  for which  $e^{\psi}v \in H^{1,1}_{mix}$ . The constant q > 0 is an unspecified 230 convergence rate also depending on what n means. These assumptions form a proper 231 framework for sparse grid-like approximation methods as those mentioned above 232 modeled after the example from the last section. Another example is the expansion 233 into tensor products of three-dimensional functions with given angular parts; see 234 [14]. The range of the  $T_n$  is in this case infinite dimensional. The exponential factor 235 is the tribute paid to the infinite extension of the domain. The assumption (39) implies 236 for the functions  $u \in H^1$  for which  $e^{\psi}u \in H^{\vartheta,1}_{mix}$  for some  $0 < \vartheta < 1$ , the error estimate 237

$$||u - T_n u||_1 \lesssim n^{-\vartheta q} |||e^{\Psi} u||_{\vartheta, 1}.$$
 (40)

The proof utilizes that the spaces  $H_{\rm mix}^{\vartheta,1}$ ,  $0 < \vartheta < 1$ , are interpolation spaces between 238 the spaces  $H^1 = H_{\rm mix}^{0,1}$  and  $H_{\rm mix}^{1,1}$ .

We conclude that for the case of the solutions u of the Schrödinger equation 240 the  $H^1$ -error  $||u - T_n u||_1$  tends faster to zero as  $n^{-\vartheta q}$  for any  $\vartheta < 3/4$ . An estimate 241 directly based on an estimate of their K-functional even shows that

$$\|u - T_n u\|_1 \le \sqrt{\ln(n)} \, n^{-3/4q} \tag{41}$$

so that up to the logarithmic term only the factor 3/4 gets lost compared to the case 243 of full mixed regularity. The estimate is optimal, at least up to the logarithmic factor, 244 and can in general not be improved further. 245

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# A Substructuring Preconditioner for **Three-Dimensional Maxwell's Equations**

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Summary. We propose a new nonoverlapping domain decomposition preconditioner for 12 the discrete system arising from the edge element discretization of the three-dimensional 13 Maxwell's equations. This preconditioner uses the simplest coarse edge element space in- 14 duced by the coarse triangulation. We will show that the rate of the PCG convergence with 15 this substructuring preconditioner is quasi-optimal, and is independent of large variations of 16 the coefficients across the local interfaces.

1 Introduction 18

When the time-dependent Maxwell's equations is solved numerically, we need to 19 solve the following **curlcurl**-system at each time step [4, 6, 8, 12]: 20

$$\mathbf{curl}(\alpha \mathbf{curlu}) + \beta \mathbf{u} = \mathbf{f} \quad \text{in} \quad \Omega \tag{1}$$

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where  $\Omega$  is assumed to be an open polyhedral domain in  $\mathbb{R}^3$ , and the coefficients 21  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  are two positive bounded functions in  $\Omega$ . We shall complement the 22 Eq. (1) with the perfect conductor condition  $\mathbf{u} \times \mathbf{n} = 0$  on  $\partial \Omega$ , where  $\mathbf{n}$  is the unit 23 outward normal vector on  $\partial \Omega$ . 24

Edge finite element methods have been widely applied in the numerical solution 25 of the system (1), see, for example, [5, 6, 8, 11]. Compared to the standard nodal 26 finite element methods, the discrete systems resulting from the edge element dis- 27 cretization are essentially different in nature. The non-overlapping domain decom- 28 position preconditioners have been well developed for the nodal element systems for 29 the standard second order elliptic problems in the past two decades, and proved both 30 numerically and theoretically to perform nearly optimally in terms of the fine mesh 31 size and subdomain size; see, e.g., the monograph [15]. But these preconditioners, 32

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or their natural generalizations turn out to perform mostly very poorly for the edge 33 element systems for the **curlcurl**-system (1), especially in three dimensions.

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A lot of important efforts have been made in the construction of effective do- 35 main decomposition methods for the system (1). A substructuring type method was 36 analysed in [16] for two dimensions, and in [2] for three dimensions with two sub- 37 domains. In [7], a novel substructuring type method was proposed for general two- 38 dimensional multiple subdomains with quite irregular boundaries, and it was proved 39 to be nearly optimal in terms of a variety of mesh decompositions and distributions 40 of physical material properties. However, it has been a challenge how to construct an 41 efficient non-overlapping domain decomposition preconditioner for the Maxwell's 42 equations in three dimensions with general multiple subdomains. A first important 43 attempt to this problem was made in [9] where a wire basket type algorithm was pro- 44 posed and analysed. Then a substructuring preconditioner and a dual-primal FETI 45 algorithm were introduced and fully analysed for three dimensions in [10] and [14], 46 respectively. These three methods have their respective advantages and disadvan- 47 tages: the algorithms in [9] and [14] both involve smaller coarse solvers but they are 48 difficult to implement; the method in [10] is easier to implement but it involves a 49 relatively large coarse solver.

This work intends to construct a new substructuring type preconditioner for the 51 three-dimensional **curlcurl**-system (1) for general multiple subdomains. In this pre- 52 conditioner, the coarse space is chosen to be the edge element space induced by 53 the coarse triangulation, so the resulting coarse solver is very cheap and simple to 54 implement. It is shown that the rate of the PCG convergence with this substructur- 55 ing preconditioner is quasi-optimal, and more importantly, independent of the large 56 variations of the coefficients in the system (1) across the local interfaces.

# 2 Domain Decompositions and Discretizations

This section introduces the non-overlapping domain decomposition of domain  $\Omega$ , 59 the weak form of the system (1) and the edge element spaces. 60

# 2.1 Initial Domain Decomposition Based on the Distribution of the Coefficients 61

We assume that the entire domain  $\Omega$  is decomposed into  $N_0$  open convex polyhedral 62 subdomains  $D_1, D_2, \cdots, D_{N_0}$  such that  $\bar{\Omega} = \bigcup_{r=1}^{N_0} \bar{D}_r$  and  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  are positive 63 constants on each subdomain  $D_r$ , namely for  $r = 1, 2, ..., N_0$ ,

$$\alpha(\mathbf{x}) = \alpha_r, \quad \beta(\mathbf{x}) = \beta_r \quad \forall \mathbf{x} \in D_r.$$

Clearly such a decomposition is always possible when the domain  $\Omega$  is occupied by multiple media. In fact, if for some medium we have an irregular nonconvex subre- 67 gion in  $\Omega$ , we can further split each nonconvex medium subregion into smaller convex subdomains. This means that our assumption does cover many practical cases, 69 especially considering the fact that the domain  $\Omega$  on which we solve the original 70 Maxwell system (1) by a finite element method is often obtained by approximating 71 the original physical domain by a polyhedral domain. Note that  $N_0$  typically is a *fixed* 72 constant in applications, so  $diam(D_r) = O(1)$ .

Let  $F_{nm}$  denote the common face of two neighboring subdomains  $D_n$  and  $D_m$ , and set  $D_{nm} = D_n \cup D_m \cup F_{nm}$ . For simplicity of the analysis, we assume

$$\beta_r \lesssim \alpha_r \lesssim d^{-2}\alpha_r, \quad r = 1, \dots, N_0.$$
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#### 2.2 Domain Decomposition

For a number  $d \in (0, 1)$ , let each polyhedron  $D_l$  be decomposed into the union 77 of some non-overlapping tetrahedra (or hexahedra)  $\{\Omega_k\}$  of size d (see [3, 15] and 78 [18]), which results in a non-overlapping domain decomposition for  $\Omega$ :  $\bar{\Omega} = \bigcup_{k=1}^N \bar{\Omega}_k$ . 79 Naturally we further assume that  $\Omega_i \cap \Omega_j = \emptyset$  when  $i \neq j$ ; if  $i \neq j$  and  $\partial \Omega_i \cap \partial \Omega_j \neq \emptyset$ , 80  $\partial \Omega_i \cap \partial \Omega_j$  is a common face (or edge or vertex) of  $\Omega_i$  and  $\Omega_j$ . Now the subdomains 81  $\Omega_1, \cdots, \Omega_N$  constitute our desired *coarse* triangulation  $\mathcal{T}_d$  of  $\Omega$ . The faces and vertices of the subdomains are always denoted by  $\Gamma$  and  $\Gamma$ 0, while the common (open) 83 face of the subdomains  $\Omega_i$  and  $\Omega_j$  are denoted by  $\Gamma_{ij}$ , and the union of all such common faces by  $\Gamma$ 1, i.e.,  $\Gamma = \bigcup \bar{\Gamma}_{ij}$ 1.  $\Gamma$ 2 will be called the interface. By  $\Gamma_i$ 3 we denote the 85 intersection of  $\Gamma$ 3 with the boundary of the subdomain  $\Omega_i$ 4. So we have  $\Gamma_i = \partial \Omega_i$ 6 if 86  $\Omega_i$ 8 is an interior subdomain of  $\Omega$ 2. We shall set  $\Omega_{ij} = \Omega_i \cup \Omega_j \cup \Gamma_{ij}$ 5.

#### 2.3 Weak Formulation

Let  $H(\mathbf{curl}; \Omega)$  be the Sobolev space consisting of all square integrable functions 89 whose  $\mathbf{curl}$ 's are also square integrable in  $\Omega$ , and  $H_0(\mathbf{curl}; \Omega)$  be a subspace of 90  $H(\mathbf{curl}; \Omega)$  of all functions whose tangential components vanish on  $\partial \Omega$ . Then by 91 writing the scalar product in  $(L^2(\Omega))^3$  as  $(\cdot, \cdot)$ , we can state the variational problem 92 for system (1) as follows:

Find  $\mathbf{u} \in H_0(\mathbf{curl}; \Omega)$  such that

$$\mathscr{A}(\mathbf{u}, \mathbf{v}) = (\mathbf{f}, \mathbf{v}), \quad \forall \mathbf{v} \in H_0(\mathbf{curl}; \Omega)$$
 (3)

where  $\mathcal{A}(\cdot,\cdot)$  is a bilinear form given by

$$\mathscr{A}(\mathbf{u}, \mathbf{v}) = (\alpha \operatorname{\mathbf{curl}} \mathbf{u}, \operatorname{\mathbf{curl}} \mathbf{v}) + (\beta \mathbf{u}, \mathbf{v}), \quad \mathbf{u}, \mathbf{v} \in H(\operatorname{\mathbf{curl}}; \Omega).$$

#### 2.4 Fine Triangulation and Their Associated Finite Element Spaces

We further divide each  $\Omega_k$  into smaller tetrahedral elements of size h so that elements from two neighboring subdomains have an intersection which is either empty or a single nodal point or an edge or a face on the interface  $\Gamma$ . Let  $\mathcal{T}_h$  be the resulting triangulation of the domain  $\Omega$ , which we assume is quasi-uniform. Then we introduce the Nédélec edge element space of the lowest order defined on  $\mathcal{T}_h$  (cf. [12] and [13]):

$$V_h(\Omega) = \left\{ \mathbf{v} \in H_0(\mathbf{curl}; \Omega); \ \mathbf{v} \mid_K \in R(K), \ \forall K \in \mathscr{T}_h \right\},$$

where R(K) is a subset of all linear polynomials on the element K of the form:

$$R(K) = \left\{ \mathbf{a} + \mathbf{b} \times \mathbf{x}; \ \mathbf{a}, \mathbf{b} \in \mathbf{R}^3, \ \mathbf{x} \in K \right\}.$$
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In an analogous way, we can define the coarse edge element space  $V_d(\Omega) \subset V_h(\Omega)$ , 107 associated with the *coarse* triangulation  $\mathscr{T}_d$ .

It is well-known that for any  $\mathbf{v} \in V_h(\Omega)$ , its tangential components are continuous on all edges of each element in the triangulation  $\mathcal{T}_h$ . Moreover, each edge element function  $\mathbf{v}$  in  $V_h(\Omega)$  is uniquely determined by its moments on each edge e of  $\mathcal{T}_h$ :

$$\left\{\lambda_e(\mathbf{v}) = \int_e \mathbf{v} \cdot \mathbf{t}_e ds; \ e \in \mathscr{E}_h\right\},$$

where  $\mathscr{E}_h$  denotes the set of the *fine* edges from the triangulation  $\mathscr{T}_h$ , and  $\mathbf{t}_e$  denotes the unit vector on the edge e.

By  $Z_h(\Omega)$  we denote the continuous piecewise linear finite element subspace of  $H^1_0(\Omega)$  associated with the triangulation  $\mathscr{T}_h$ . Similarly, let  $Z_d(\Omega)$  denote the continuous piecewise linear finite element subspace of  $H^1_0(\Omega)$  associated with the triangulation  $\mathscr{T}_d$ .

#### 2.5 Discrete Variational Problem

Using the edge element space  $V_h(\Omega)$ , the system (3) may be approximated as follows: Find  $\mathbf{u}_h \in V_h(\Omega)$  such that

$$(\alpha \mathbf{curl} \ \mathbf{u}_h, \mathbf{curl} \ \mathbf{v}_h) + (\beta \mathbf{u}_h, \mathbf{v}_h) = (\mathbf{f}, \mathbf{v}_h), \quad \forall \mathbf{v}_h \in V_h(\Omega). \tag{4}$$

Define the operator  $A: V_h(\Omega) \to V_h(\Omega)$  by

$$(A\mathbf{u}_h, \mathbf{v}_h) = (\alpha \mathbf{curl} \ \mathbf{u}_h, \mathbf{curl} \ \mathbf{v}_h) + (\beta \mathbf{u}_h, \mathbf{v}_h), \quad \forall \mathbf{u}_h, \mathbf{v}_h \in V_h(\Omega),$$
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Then, (4) can be written in the operator form

$$A\mathbf{u}_h = \mathbf{f}_h. \tag{5}$$

# 3 A Nearly Optimal Preconditioner for A

#### 3.1 Construction of the Preconditioner

We first introduce some useful sets and subspaces.

 $\mathcal{E}_h$ : the set of all edges from the triangulations  $\mathcal{T}_h$ ;

 $\mathscr{E}_{\Gamma,h}$ : the set of edges which belong to  $\mathscr{E}_h$  and have two endpoints on the interface 129

 $\mathscr{E}_d$ : the set of all (coarse) edges from the triangulations  $\mathscr{T}_d$ ;

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 $\mathcal{W}_{E}$ : the union of all the coarse edges  $E' \in \mathcal{E}_{d}$ , which have a common endpoint 132 with the coarse edge  $E \in \mathcal{E}_d$ . And  $\mathcal{W}_E$  is called E-basket. 133

 $\mathscr{E}^b_{\mathrm{E},h}$ : the set of all (fine) edges which belong to  $\mathscr{E}_h$  and have at least one endpoint on  $\mathscr{W}_{E}$ ; 135

Let D be either a subdomain  $\Omega_i$  or a subdomain  $\Omega_k$  or a subdomain  $\Omega_{ij}$  or a sub-136 domain  $D_{mn}$ . The restrictions of  $V_h(\Omega)$  (resp.  $Z_h(\Omega)$ ) on D is denoted by  $V_h(D)$  (resp. 137  $Z_h(D)$ ). The following local subspaces of  $V_h(D)$  will be important to our analysis: 138

$$V_h^0(D) = \left\{ \mathbf{v} \in V_h(D); \ \mathbf{v} \times \mathbf{n} = 0 \ \text{on} \ \partial D \right\},$$
 139

and

$$Z_h^0(D) = \Big\{ \varphi \in Z_h(\Omega); \ supp \ \varphi \subset D \Big\}.$$

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We define subspaces of  $V_h(\Omega)$ :

$$V_h^H(\Omega) = \Big\{ \mathbf{v} \in V_h(\Omega); \ \mathbf{v} \ ext{is the discrete $A$-extension of } \mathbf{v}|_{\partial\Omega_k} \ ext{in each } \Omega_k \Big\},$$

$$V_h^H(\Omega_{ij}) = V_h^H(\Omega) \bigcap V_h^0(\Omega_{ij}),$$

and for  $E \in \mathcal{E}_d$ ,

$$V_h^{\mathrm{E}}(\Omega) = \left\{ \mathbf{v} \in V_h^H(\Omega); \ \lambda_e(\mathbf{v}) = 0 \text{ for each } e \in \mathscr{E}_{\Gamma,h} \backslash \mathscr{E}_{\mathrm{E},h}^b \right\}.$$

It is well known that a suitable *coarse* subspace plays a key role in the construction of an effective domain decomposition preconditioner, and it is generally rather technical and problem-dependent to choose such a coarse subspace. Surprisingly we are going to choose the coarse subspace to be the simplest one, namely the subspace 151  $V_d(\Omega)$  induced by the coarse triangluation  $\mathcal{T}_d$ . 152

It is easy to see that the space  $V_h(\Omega)$  has the (non-direct sum) decomposition

$$V_h(\Omega) = V_d(\Omega) + \sum_{k=1}^{N} V_h^0(\Omega_k) + \sum_{E} V_h^E(\Omega) + \sum_{F_i} V_h^H(\Omega_{ij}). \tag{6}$$

Next, we define the corresponding solvers on the subspaces  $V_h^0(\Omega_k)$ ,  $V_h^{\rm E}(\Omega)$ ,  $V_h^H(\Omega_{ij})$  and  $V_d(\Omega)$ . 155

As usual, we denote the restriction of A on  $V_h^0(\Omega_k)$  by  $A_k$ , i.e., 156

$$(A_k \mathbf{v}, \mathbf{u})_{\Omega_k} = (A \mathbf{v}, \mathbf{u}) = \mathscr{A}(\mathbf{v}, \mathbf{u}), \ \mathbf{v} \in V_h^0(\Omega_k), \ \forall \mathbf{u} \in V_h^0(\Omega_k).$$
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Let  $B_k: V_h^0(\Omega_k) \to V_h^0(\Omega_k), B_d: V_d(\Omega) \to V_d(\Omega)$  and  $B_{ij}: V_h^H(\Omega_{ij}) \to V_h^H(\Omega_{ij})$ 158 be the symmetric and positive definite operators such that 159

$$(B_k \mathbf{v}, \mathbf{v}) \cong (A_k \mathbf{v}_k, \mathbf{v}_k)_{\Omega_k}, \quad \forall \mathbf{v} \in V_h^0(\Omega_k),$$

where  $\mathbf{v}_k = \mathbf{v}|_{\Omega_k}$  for  $k = 1, 2, \dots, N$ , and 161

$$(B_d \mathbf{v}_d, \mathbf{v}_d) \cong \mathscr{A}(\mathbf{v}_d, \mathbf{v}_d), \quad \forall \mathbf{v}_d \in V_d(\Omega),$$
  
 $(B_{ij} \mathbf{v}, \mathbf{v}) \cong \mathscr{A}(\mathbf{v}, \mathbf{v}), \quad \forall \mathbf{v} \in V_h^H(\Omega_{ij}).$ 

The symbol ≈ above means each of the two quantities involved is bounded by the 162 other up to a constant independent of h, d and functions involved in the two quantities.

The local solvers on  $V_h^{\rm E}(\Omega)$  should be solvable in an efficient manner, and their 165 constructions are much more tricky and technical than the others. To do so, we introduce more notation.

For any face F from the triangulations  $\mathcal{I}_d$ , we use  $F_b$  to denote the union of all 168  $\mathcal{T}_h$ -induced (closed) triangles on F, which have either one single vertex or one edge 169 lying on  $\partial F$ , and  $F_{\partial}$  to denote the open set  $F \setminus F_b$ . For any subdomain  $\Omega_k$ , define 170

$$\Delta_k = \bigcup_{\mathrm{F} \subset \Gamma_k} \mathrm{F}_b, \quad k = 1, \cdots, N.$$

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We will also need the so-called tangential divergence  $\operatorname{div}_{\tau}\Phi = \operatorname{curl}_{S}\Phi$  for  $\Phi \in {}_{172}$  $V_h(\Gamma_k)$ , which is defined here as in [1, 2]. Then we can introduce our local solver 173  $B_{\rm E}: V_h^{\rm E}(\Omega) \to V_h^{\rm E}(\Omega)$  as follows: 174

$$(B_{E}\mathbf{v},\mathbf{u}) = h[1 + \log(d/h)] \sum_{k=1}^{N} \left\{ \alpha_{k} \langle \operatorname{div}_{\tau}(\mathbf{v} \times \mathbf{n}) |_{\varGamma_{k}}, \operatorname{div}_{\tau}(\mathbf{u} \times \mathbf{n}) |_{\varGamma_{k}} \rangle_{\Delta_{k}} + \beta_{k} \langle \mathbf{v} \times \mathbf{n}, \mathbf{u} \times \mathbf{n} \rangle_{\Delta_{k}} \right\}, \qquad \mathbf{v} \in V_{h}^{E}(\Omega), \ \forall \mathbf{u} \in V_{h}^{E}(\Omega).$$
(7)

For convenience, we call  $B_{\rm E}$  an

Let  $Q_k:V_h(\Omega) o V_h^0(\Omega_k), \ Q_d:V_h(\Omega) o V_d(\Omega), \ Q_{\mathrm E}:V_h(\Omega) o V_h^{\mathrm E}(\Omega)$  and  $Q_{ij}:V_h(\Omega)\to V_h^H(\Omega_{ij})$  be the standard the standard  $L^2$ -projections. Then we are ready to propose our new preconditioner for A as follows: 178

$$B^{-1} = B_d^{-1} Q_d + \sum_{k=1}^N B_k^{-1} Q_k + \omega \sum_{\mathcal{E}} B_{\mathcal{E}}^{-1} Q_{\mathcal{E}} + \sum_{\Gamma_{ij}} B_{ij}^{-1} Q_{ij}, \tag{8}$$

where  $\omega$  is a (constant) relaxation parameter, which is introduced to obtain a balance 179 between the local solvers  $B_{\rm E}$  and other remaining solvers. 180

#### 3.2 Algorithm Based on the New Preconditioner and Main Results

The action of the preconditioner  $B^{-1}$  which is needed in each PCG iteration can be 182 described in the following algorithm. 183

**Algorithm 4.1.** For  $\mathbf{g} \in V_h(\Omega)$ , we can compute  $\mathbf{u} = B^{-1}\mathbf{g}$  in five steps. 184

Step 1. Solve the system for  $\mathbf{u}_d \in V_d(\Omega)$ :

$$(B_d \mathbf{u}_d, \mathbf{v}_d) = (\mathbf{g}, \mathbf{v}_d), \quad \forall \mathbf{v}_d \in V_d(\Omega);$$

Step 2. Solve the following system for  $\mathbf{u}_k \in V_h^0(\Omega_k)$  in each subdomain in parallel:

$$(B_k \mathbf{u}_k, \mathbf{v}) = (\mathbf{g}, \mathbf{v}), \quad \forall \mathbf{v} \in V_b^0(\Omega_k), \quad k = 1, \cdots, N;$$

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Step 3. Solve the following system for  $\mathbf{u}_{ij} \in V_h^0(\Omega_{ij})$  in each subdomain  $\Omega_{ij}$  in 190 parallel:

$$(B_{ij}\mathbf{u}_{ij},\mathbf{v}) = (\mathbf{g},\mathbf{v}) - (A_i\mathbf{u}_i,\mathbf{v})_{\Omega_i} - (A_j\mathbf{u}_j,\mathbf{v})_{\Omega_j}, \quad \forall \mathbf{v} \in V_h^0(\Omega_{ij});$$

Step 4. Solve the system for  $\mathbf{u}_{\mathrm{E}} \in V_h^{\mathrm{E}}(\Omega)$ :

$$(B_{\mathrm{E}}\mathbf{u}_{\mathrm{E}},\mathbf{v})=(\mathbf{g},\tilde{\mathbf{v}})-\sum_{k=1}^{N}(A_{k}\mathbf{u}_{k},\tilde{\mathbf{v}}),\ \mathbf{v}\in V_{h}^{\mathrm{E}}(\Omega),$$
 194

where  $\tilde{\mathbf{v}} \in V_h(\Omega)$  is a natural extension of  $(\mathbf{v} \times \mathbf{n})|_{\Gamma}$  by zero.

Step 5. Set  $\Phi_h = (\sum_{\Gamma_{ij}} \mathbf{u}_{ij} + \sum_{\mathbf{E}} \mathbf{u}_{\mathbf{E}}) \times \mathbf{n}|_{\Gamma}$  and compute the A-extension A-extension

of  $\Phi_h$  on each  $\Omega_k$  to obtain  $\mathbf{u}^H \in V_h^H(\Omega)$ . This leads to

$$\mathbf{u} = \mathbf{u}_d + \sum_{k=1}^N \mathbf{u}_k + \mathbf{u}^H.$$
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Remark 1. For the local solver  $B_{ij}$  on each face  $\Gamma_{ij}$ , we may use the face extended domain formed by, e.g., one half of each of the two neighboring subdomains  $\Omega_i$  200 and  $\Omega_j$ . Such definition of  $B_{ij}$ 's can reduce the computational complexity in their numerical realization.

Let E denote a coarse edge of the subdomain  $D_r$ . Define

$$V_h^{\perp}(\Omega) = \{\mathbf{v}_h : \mathbf{v}_h \in V_h(\Omega), \ \int_E \mathbf{v}_h \cdot \mathbf{t}_E ds = 0 \text{ for each } E\}.$$

We shall use  $\kappa^{\perp}(B^{-1}A)$  to denote the *induced condition number* of the preconditioned system  $B^{-1}A$  associated with the subspace  $V_h^{\perp}(\Omega)$ , namely the condition number of  $B^{-1}A$  restricted on the subspace  $V_h^{\perp}(\Omega)$  (cf. [17]). At this moment we are able to establish only the following estimate of the induced condition number. As the estimate is quite lengthy and technical, we cannot include it here due to the page limitation.

**Theorem 1.** Under the assumptions (2), the preconditioner B given in (8) is nearly optimal in the sense that

$$\kappa^{\perp}(B^{-1}A) \le C[1 + \log(d/h)]^2 [1 + \log(1/h)]^2 \tag{9}$$

where the constant C is independent of h, d and the jumps of the coefficients.

As we see from the above theorem that the induced condition number grows 214 logarithmically with the degrees of freedom in each subdomain, but also with the 215

degrees of freedom of the entire fine mesh. We believe this is mainly due to the re- 216 striction of our current analysis technique, namely the estimate must be done for the 217 induced condition number in the subspace  $V_h^\perp(\Omega)$  associated with the coarse trian- 218 gulation formed by the material subdomains  $D_r$ . We expect the estimate should be 219 finally carried out directly in the entire edge element space  $V_h(\Omega)$ , that will remove 220 the logarithmic factor of 1/h in the estimate (9). This expectation has already been 221 confirmed by our three-dimensional numerical experiments; see the next section.

## 4 Numerical Experiments

In this section we shall conduct some numerical experiments to check the conver- 224 gence of the newly proposed preconditioner, and find out whether they are consistent 225 with the prediction of the convergence theory developed in the previous sections.

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In our experiments, we take the domain to be the unit cube  $\Omega = (0,1)^3$ , while 227 the right-hand side f of the system (1) is selected such that the exact solution  $\mathbf{u} = 228$  $(u_1, u_2, u_3)^T$  is given by 229

$$u_1 = xyz(x-1)(y-1)(z-1),$$

$$u_2 = \sin(\pi x)\sin(\pi y)\sin(\pi z),$$

$$u_3 = (1-e^x)(1-e^{x-1})(1-e^y)(1-e^{y-1})(1-e^z)(1-e^{z-1}),$$

when the coefficients  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  are both constant 1. This right-hand side  $\mathbf{f}$  is 230 then fixed in all our experiments, but the coefficients  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  may be taken 231 differently.

We then need to triangulate the domain  $\Omega$  into subdomains  $\{\Omega_k\}$ . For this, we 233 first partition the three edges of  $\Omega$  on x-, y- and z-axis into n equal subintervals from 234 which one can naturally generate  $n^3$  equal smaller cubes of size d = 1/n. This yields 235 the desired subdomain decomposition in our experiments.

Next, we further triangulate each subdomain  $\Omega_k$  to get a fine triangulation  $\mathcal{T}_h$  of 237 size h over the domain  $\Omega$ . To generate  $\mathcal{T}_h$ , we divide each subdomain into  $m^3$  equal 238 smaller cubes of size h = 1/(mn), in the same manner as done in the previous subdo-239 main generation. Then  $\mathcal{T}_h$  is obtained by triangulating each cube into six tetrahedra. 240 For easy identification, we may denote the triangulation  $\mathcal{T}_h$  as  $m^3(n^3)$  below.

The edge finite element space of the lowest order is used for the discretization 242 of (3). The resulting system (5) is solved by PCG method with the newly proposed 243 preconditioners B defined in Sect. 4. We shall choose the balancing parameter  $\omega$  in 244 front of the E-basket local solvers  $B_{\rm E}$  in (8) as  $\omega = 1$  or  $\omega = 2.5$ .

We consider various distributions of the coefficients  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  and report 246 the corresponding numbers of PCG iterations, and the condition numbers of  $B^{-1}A$  247 for some representative cases. The PCG iteration is terminated in our experiments 248 when the relative residual is less than  $10^{-6}$ .

Case (i): coefficients  $\alpha(\mathbf{x}) = \beta(\mathbf{x}) = 1$ , with no jumps. The PCG iterations and the 250 condition numbers (in brackets) for  $\omega = 2.5$  are listed in Table 1. 251

Page 80

	$\omega = 1.0$				$\omega = 2.5$						
$m \setminus n$	4	6	8	10	4	6	8	10			
4	34	33	32	32	31 (34.24)	31 (36.31)	31 (36.94)	30 (37.40)			
8	41	40	39	38	39 (52.15)	38 (53.78)	37 (54.21)	37 (54.61)			
12	48	47	44	42	43 (64.29)	43 (65.91)	41 (66.19)	41 (66.62)			
16	51	50	49	45	47 (74.40)	46 (75.69)	44 (75.82)	44 (76.39)			

Table 1. Iterations (and condition numbers) with smooth coefficients

We observe from the above table that the number of PCG iterations grows slowly 252 when m = d/h increases but n = 1/d is fixed, and that these numbers vary stably 253 when m is fixed but n increases. This justifies our early expection that the condition 254 number of the preconditioned system  $B^{-1}A$  should grow logarithmically with d/h 255 only, not with 1/h.

One important issue we like to draw the readers' attention to is the large-scale of 257 the discrete system we are solving. For instance, when m = 16 and n = 10, the total 258 number of degrees of freedom for the fine edge element system is about 28,672,000. 259

Case (ii): coefficients  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  have large jumps:

$$\alpha(\mathbf{x}) = \beta(\mathbf{x}) = \alpha_0$$
 in  $D$ ;  $\alpha(\mathbf{x}) = \beta(\mathbf{x}) = 1$  in  $\Omega \setminus D$ .

where  $D \subset \Omega$  is a union of several subdomains  $\Omega_k$ . We choose  $\alpha_0 = 10^{-5}$  or  $\alpha_0 =$ 10<sup>5</sup>, and consider two choices of D, where one does not have cross-points, while the 263 other has one *cross-point*. 264

Example 1: 265

$$D = \left[\frac{1}{4}, \, \frac{1}{2}\right]^3.$$

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t2.1 t2 2 t2 3 t2 4 t2.5

t2.7

Example 2 267

$$D = \left[\frac{1}{4}, \frac{1}{2}\right]^3 \bigcup \left[\frac{1}{2}, \frac{3}{4}\right]^3.$$
 268

The numerical results are given in Tables 2 and 3, from which we can make some 269 similar observations about the PCG convergence in terms of the mesh and subdomain 270 quantities d/h and d as we did for Case (i). 271

			Example 1		Example 2				
	$\omega = 1.0$		$\omega =$	2.5	$\omega = 1.0$		$\omega = 2.5$		
$m \setminus n$	4	8	4	8	4	8	4	8	
4	29	31	26 (32.00)	29 (35.97)	28	30	26 (35.51)	30 (35.97)	
8	35	38	32 (44.88)	37 (52.97)	35	38	32 (45.88)	37 (52.59)	
12	38	45	36 (56.02)	42 (64.96)	37	45	35 (55.66)	41 (63.81)	
16	40	49	37 (64.65)	45 (74.68)	40	49	37 (65.65)	45 (74.31)	

**Table 2.** Iterations (and condition numbers) with  $\alpha_0 = 10^{-5}$ 

			Example 1		Example 2				
	$\omega = 1.0$		$\omega = 2.5$			= 1.0	$\omega = 2.5$		
$m \setminus n$	4	8	4	8	4	8	4	8	
4	42	42	36 (40.47)	36 (42.71)	42	44	38 (40.55)	37 (42.72)	
8	49	48	45 (61.08)	44 (62.89)	52	51	46 (60.20)	45 (62.89)	
12	55	54	50 (74.04)	49 (76.28)	56	56	50 (76.24)	51 (76.28)	
16	59	57	54 (91.51)	52 (86.45)	59	59	53 (83.35)	54 (86.45)	

**Table 3.** Iterations (and condition numbers) with  $\alpha_0 = 10^5$ 

Case (iii): coefficients  $\alpha(\mathbf{x})$  and  $\beta(\mathbf{x})$  have large jumps:

$$\alpha(\mathbf{x}) = \begin{cases} \alpha_0, & \text{in } D \\ 1, & \text{in } \Omega \backslash D, \end{cases} \qquad \beta(\mathbf{x}) = \begin{cases} \beta_0, & \text{in } D \\ 1, & \text{in } \Omega \backslash D, \end{cases}$$
 273

t3.2

t3.6 t3.7

272

where  $D \subset \Omega$  is a union of several subdomains  $\Omega_k$ . We choose  $\alpha_0 = 10^{-5}$  or  $\alpha_0 = 274$   $10^5$ , but  $\beta_0 \neq \alpha_0$ . We still consider two different regions D from Examples 1 and 2 in 275 the previous Case (ii), but choose the balancing parameter  $\omega$  in front of the E-basket 276 local solvers  $B_{\rm E}$  in (8) as  $\omega = 2.5$ .

The numerical results are given in Tables 4 and 5. Again, we can make similar observations about the PCG convergence in terms of the mesh and subdomain quantities d/h and d as we did for Case (i).

		Exar	nple	1	Example 2				
	$\beta_0$ :	$= \alpha_0 \times 10^2$	$\beta_0$ :	$= \alpha_0 \times 10^{-2}$	$\beta_0$ :	$= \alpha_0 \times 10^2$	$\beta_0$ =	$= \alpha_0 \times 10^{-2}$	
$m \setminus n$	4	8	4	8	4	8	4	8	
4	30	36	46	47	30	36	45	47	
8	39	43	56	56	39	45	56	56	
16	49	52	65	65	49	52	63	65	

**Table 4.** Iterations with  $\alpha_0 = 10^{-5}$ 

		Exan	nple	1	Example 2				
	$\beta_0 = \alpha_0 \times 10^2$		$\beta_0$ =	$= \alpha_0 \times 10^{-2}$	$\beta_0$ :	$= \alpha_0 \times 10^2$	$\beta_0 = \alpha_0 \times 10^{-2}$		
$m \setminus n$	4	8	4	8	4	8	4	8	
4	31	37	38	41	31	37	39	46	
8	37	47	46	49	37	47	53	58	
16	48	56	55	57	48	56	66	73	

**Table 5.** Iterations with  $\alpha_0 = 10^5$ 

We may also observe from the previous numerical experiments that appropriate 281 choices of the parameter  $\omega$  can significantly improve the efficiency of the preconditioner B. It is important to see that the choices of  $\omega$  seem independent of the fine 283

and coarse meshsizes h and d, so we may determine  $\omega$  by solving some small scale 284

systems, e.g., a system with $m = n = 4$ .	285
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# A Two-Level Schwarz Preconditioner for **Heterogeneous Problems**

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1 Introduction 11

Coarse space correction is essential to achieve algorithmic scalability in domain de- 12 composition methods. Our goal here is to build a robust coarse space for Schwarz 13 type preconditioners for elliptic problems with highly heterogeneous coefficients 14 when the discontinuities are not just across but also along subdomain interfaces, 15 where classical results break down [3, 6, 9, 15].

In previous work, [7], we proposed the construction of a coarse subspace based 17 on the low-frequency modes associated with the Dirichlet-to-Neumann (DtN) map 18 on each subdomain. A rigorous analysis was recently provided in [2]. Similar ideas 19 to build stable coarse spaces, based on the solution of local eigenvalue problems 20 on entire subdomains, can be found in [4], and even traced back to similar ideas 21 for algebraic multigrid methods in [1]. However, we will argue below that the DtN 22 coarse space presented here is better designed to deal with coefficient variations that 23 are strictly interior to the subdomain, being as robust as, but leading to a smaller 24 dimension than the coarse space analysed in [4].

The robustness result that we obtain, generalizes the classical estimates for over- 26 lapping Schwarz methods to the case where the coarse space is richer than just the 27 constant mode per domain [8], or other classical coarse spaces (cf. [15]). The analysis 28 is inspired by that in [4, 13] and crucially uses the framework of weighted Poincaré 29 inequalities, introduced in [10, 11] and successfully applied also to other methods in 30 [12, 14].

# 2 Two-Level Schwarz Method with DtN Coarse Space

We consider the variational formulation of a second order, elliptic boundary value 33 problem with Dirichlet boundary conditions: Find  $u^* \in H_0^1(\Omega)$ , for a given domain 34

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 $\Omega \subset \mathbb{R}^d \ (d=2 \text{ or } 3)$  and a source term  $f \in L_2(\Omega)$ , such that

$$a(u^*, v) \equiv \int_{\Omega} \alpha(x) \nabla u^* \cdot \nabla v = \int_{\Omega} fv \equiv (f, v), \quad \forall v \in H_0^1(\Omega),$$
 (1)

and the diffusion coefficient  $\alpha = \alpha(x)$  is a positive piecewise constant function that may have large variations within  $\Omega$ .

We consider a discretization of the variational problem (1) with continuous, 38 piecewise linear finite elements (FE). For a shape regular, simplicial triangulation 39  $\mathcal{T}_h$  of  $\Omega$ , the standard space of continuous and piecewise linear functions (w.r.t  $\mathcal{T}_h$ ) 40 is then denoted by  $V_h$ . The subspace of functions from  $V_h$  that vanish on the boundary of  $\Omega$  is denoted by  $V_{h,0}$ . The discrete FE problem that we want to solve is: Find 42  $u_h \in V_{h,0}$  such that

$$a(u_h, v_h) = (f, v_h), \quad \forall v_h \in V_{h,0}. \tag{2}$$

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Given the usual nodal basis  $\{\phi_i\}_{i=1}^n$  for  $V_{h,0}$  consisting of "hat" functions with n:=44 dim $(V_{h,0})$ , (2) can be compactly written as

$$A$$
**u** = **f**, with  $A_{ij} := a(\phi_j, \phi_i)$  and  $f_i = (f, \phi_i), i, j = 1, ..., n,$  (3)

where **u** and **f** are respectively the vector of coefficients corresponding to the un-  $\frac{1}{4}$  known FE function  $u_h$  in (2) and to the r.h.s function f.

Two-level Schwarz type methods for (2) are now constructed by choosing an 48 overlapping decomposition  $\{\Omega_j\}_{j=1}^J$  of  $\Omega$  with a subordinate partition of unity 49  $\{\chi_j\}_{j=1}^J$ , as well as a suitable coarse subspace  $V_H \subset V_{h,0}$ . In practice the overlapping subdomains  $\Omega_j$  can be constructed automatically given the system matrix A by 1 using a graph partitioner, such as METIS, and adding on a number of layers of fine 52 grid elements to the resulting nonoverlapping subdomains. A suitable partition of 53 unity can be constructed from the geometric information of the fine grid. For more 54 details see e.g. [15] or [2]. We assume that each point  $x \in \Omega$  is contained in at most 55  $N_0$  subdomains  $\Omega_j$ .

The crucial ingredient to obtain robust two-level methods for problems with heterogeneous coefficients is the choice of coarse space  $V_H \subset V_{h,0}$ . Let us assume for the smoment that we have such a space  $V_H$  and a restriction operator  $R_0$  from  $V_{h,0}$  to  $V_H$  so and define restriction operators  $R_j$  from functions in  $V_{h,0}$  to functions in  $V_{h,0}(\Omega_j)$ , or 60 from vectors in  $\mathbb{R}^n$  to vectors in  $\mathbb{R}^{\dim V_{h,0}(\Omega_j)}$ , by setting  $(R_j u)(x_i) = u(x_i)$  for every grid point  $x_i \in \Omega_j$ . The two-level overlapping additive Schwarz preconditioner for 62 (3) is then simply

$$M_{AS,2}^{-1} = \sum_{j=0}^{J} R_j^T A_j^{-1} R_j$$
 where  $A_j := R_j A R_j^T$ ,  $j = 0, ..., J$ . (4)

In the classical algorithm  $V_H$  consists simply of FEs on a coarser triangulation 64  $\mathcal{T}_H$  of  $\Omega$  and  $R_H$  is the canonical restriction from  $V_{h,0}$  to  $V_H$ , leading to a fully scal-65 able iterative method with respect to mesh/problem size (provided the overlap size is 66 proportional to the coarse mesh size H). However, unfortunately this preconditioner 67 is not robust to strong variations in the coefficient  $\alpha$ . We will now present a new, 68

completely local approach to construct a robust coarse space, as well as an associated restriction operator using eigenvectors of local Dirichlet-to-Neumann maps, 70 proposed in [7].

We start by constructing suitable local functions on each subdomain  $\Omega_j$  that will 72 then be used to construct a basis for  $V_H$ . To this end, let us fix  $j \in \{1, ..., J\}$  and 73 first consider at the continuous level the Dirichlet-to-Neumann map  $\mathrm{DtN}_j$  on the 74 boundary of  $\Omega_j$ . Let  $\Gamma_j := \partial \Omega_j$  and let  $v_\Gamma : \Gamma_j \to \mathbb{R}$  be a given function, such that 75  $v_\Gamma|_{\partial\Omega} = 0$  if  $\Gamma_j \cap \partial\Omega \neq \emptyset$ . We define

$$\mathrm{DtN}_{j}(\nu_{\Gamma}) := \left. \alpha \frac{\partial \nu}{\partial \nu_{j}} \right|_{\Gamma_{j}},$$

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where  $v_i$  is the unit outward normal to  $\Omega_i$  on  $\Gamma_i$ , and  $\nu$  satisfies

$$-\operatorname{div}(\alpha \nabla v) = 0 \text{ in } \Omega_i, \quad v = v_{\Gamma} \text{ on } \Gamma.$$
 (5)

The function v is the  $\alpha$ -harmonic extension of the boundary data  $v_{\Gamma}$  to the interior 79 of  $\Omega_i$ .

To construct the (local) coarse basis functions, we now find the low frequency 81 modes of the Dirichlet-to-Neumann operator  $DtN_j$  with respect to the weighted  $L_2$ – 82 norm on  $\Gamma_j$ , i.e. the smallest eigenvalues of

$$DtN_{j}(v_{\Gamma}^{(j)}) = \lambda^{(j)} \alpha v_{\Gamma}^{(j)}. \tag{6}$$

Then we extend each of these modes  $v_{\Gamma}^{(j)}$   $\alpha$ -harmonically to the whole domain and 84 let  $v^{(j)}$  be its extension. This is equivalent to the Steklov eigenvalue problem of 85 looking for the pair  $(v^{(j)}, \lambda^{(j)})$  which satisfies: 86

$$-\operatorname{div}(\alpha \nabla v^{(j)}) = 0 \text{ in } \Omega_j \quad \text{and} \quad \alpha \frac{\partial v^{(j)}}{\partial v_j} = \lambda \, \alpha v^{(j)} \text{ on } \Gamma_j. \tag{7}$$

The variational formulation of (7) is to find  $(v^{(j)}, \lambda^{(j)}) \in H^1(\Omega_j) \times \mathbb{R}$  such that

$$\int_{\Omega_{j}} \alpha \nabla v^{(j)} \cdot \nabla w = \lambda^{(j)} \int_{\Gamma_{j}} \operatorname{tr}_{j} \alpha v^{(j)} w, \ \forall w \in H^{1}(\Omega_{j}),$$
 (8)

where  $\operatorname{tr}_j \alpha(x) := \lim_{y \in \Omega_j \to x} \alpha(y)$ . To discretize this generalized eigenvalue problem, 88 we consider for all  $v, w \in H^1(\Omega_j)$  the bilinear forms

$$a_j(v,w) := \int_{\Omega_j} \alpha \nabla v \cdot \nabla w$$
 and  $m_j(v,w) := \int_{\Gamma_j} \operatorname{tr}_j \alpha v w$ 

and restrict (8) to the FE space  $V_h(\Omega_j)$ . The coefficient matrices associated with the 91 variational forms  $a_j$  and  $m_j$  are 92

$$A_{kl}^{(j)}:=\int_{\Omega_j} lpha 
abla \phi_k \cdot 
abla \phi_l \quad ext{and} \quad M_{kl}^{(j)}:=\int_{\Gamma_j} ext{tr}_j lpha \phi_k \, \phi_l,$$

where  $\phi_k$  and  $\phi_l$  are any two nodal basis functions for  $V_h(\Omega_j)$  associated with vertices 94 of  $\mathcal{T}_h$  contained in  $\overline{\Omega}_j$ . Then the FE approximation to (8) in matrix notation is 95

$$A^{(j)}\mathbf{v}^{(j)} = \lambda^{(j)}M^{(j)}\mathbf{v}^{(j)} \tag{9}$$

where  $\mathbf{v}^{(j)} \in \mathbb{R}^{n_j}$ ,  $n_j := \dim V_h(\Omega_j)$ , denotes the degrees of freedom of the FE approximation to  $v^{(j)}$  in  $V_h(\Omega_j)$ .

Let the  $n_j$  eigenpairs  $(\lambda_\ell^{(j)}, \mathbf{v}_\ell)_{\ell=1}^{n_j}$  corresponding to (9) be numbered in increasing 98 order of  $\lambda_\ell^{(j)}$ . Since  $M_{kl}^{(j)} \neq 0$  only if  $\phi_k$  and  $\phi_l$  are associated with the  $n_\Gamma$  vertices 99 of  $\mathscr{T}_h$  that lie on  $\Gamma_j$ , it is easy to see that at most  $n_\Gamma$  of the eigenvalues  $\lambda_\ell^{(j)}$  are 100 finite. Moreover, the smallest eigenvalue  $\lambda_1^{(j)} = 0$  with constant eigenvector and the 101 set of eigenvectors  $\{\mathbf{v}_\ell\}_{\ell=1}^{n_j}$  can be chosen so that they are  $A^{(j)}$ -orthonormal. The 102 local coarse space is now defined as the span of the FE functions  $v_\ell^{(j)} \in V_h(\Omega_j)$ , 103  $\ell \leq m_j \leq n_\Gamma$ , corresponding to the first  $m_j$  eigenpairs of (9). For each subdomain 104  $\Omega_j$ , we choose the value of  $m_j$  such that  $\lambda_\ell^{(j)} < \operatorname{diam}(\Omega_j)^{-1}$ , for all  $\ell \leq m_j$ , and 105  $\lambda_{m_j+1}^{(j)} \geq \operatorname{diam}(\Omega_j)^{-1}$ . We will see in the analysis in the next section why this is a 106 sensible choice.

Using the partition of unity  $\{\chi_j\}_{j=1}^J$ , we now combine the local basis functions 108 constructed in the previous section to obtain a conforming coarse space  $V_H \subset V_{h,0}$  on 109 all of  $\Omega$ . The new coarse space is defined as

$$V_H := \operatorname{span}\left\{I_h\left(\chi_j v_\ell^{(j)}\right) : 1 \le j \le J \text{ and } 1 \le \ell \le m_j\right\},\tag{10}$$

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where  $I_h$  is the standard nodal interpolant onto  $V_{h,0}(\Omega)$ . The dimension of  $V_H$  is 111  $\sum_{j=1}^J m_j$ . By construction each of the functions  $I_h(\chi_j v_\ell^{(j)}) \in V_{h_0}$ , so that as required 112  $V_H \subset V_{h,0}$ . The transfer operator  $R_0$  from  $V_{h_0}$  to  $V_H$  is defined in a canonical way by 113 setting  $R_0^T u_H(x_i) = u_H(x_i)$ , for all  $u_H \in V_H$  and for all vertices  $x_i$  of  $\mathcal{T}_h$ .

We will see in the next section that under some mild assumptions on the variability of  $\alpha$  this choice of coarse space leads to a scalable and coefficient-robust domain decomposition method with supporting theory.

# 3 Conditioning Analysis

To analyse this method let us first define the boundary layer  $\Omega_j^\circ := \{x \in \Omega_j : \chi_j(x) < 119 \}$  for each  $\Omega_j$  that is overlapped by neighbouring domains, i.e. We assume that this 120 layer is uniformly of width  $\geq \delta_j$ , in the sense that it can be subdivided into shape 121 regular regions of diameter  $\delta_j$ , and that the triangulation  $\mathscr{T}_h$  resolves it. This also 122 guarantees that it is possible to find a partition of unity such that  $|\chi_j| = \mathscr{O}(1)$  and 123  $|\nabla \chi_j| = \mathscr{O}(\delta_j^{-1})$ .

We now state the key assumption on the coefficient distribution  $\alpha(x)$ .

**Assumption 1** We assume that, for each  $j=1,\ldots,J$ , there exists a set  $X_j \subset \Gamma_j$  (not 126 necessarily connected) such that (i)  $\max_{x,y\in X_k} \frac{\alpha(x)}{\alpha(y)} = \mathcal{O}(1)$  and (ii) there exists a path 127

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 $P_y$  from each  $y \in \Omega_i$  to  $X_i$ , such that  $\alpha(x)$  is an increasing function along  $P_y$  (from y to  $X_i$ ). 129

Lemma 1 (weighted Poincaré inequality [10]). Let Assumption 1 hold.

$$\int_{\Omega_j^\circ} \alpha |v - \overline{v}^{X_j}|^2 \, \leq \, C_P \, \delta_j \, \int_{\Omega_j^\circ} \alpha |\nabla v|^2, \qquad \text{for all } \, v \in V_h(\Omega_j),$$

where 
$$\overline{v}^{X_j} := \frac{1}{|X_i|} \int_{X_i} v$$
.

Remark 1. Note that Assumption 1 is related to the classical notion of quasi-monotonicity coined in [3]. It ensures that the constant  $C_P$  in the Poincaré-type inequality 133 in Lemma 1, as well as all the other (hidden) constants below are independent of the values of the coefficient function  $\alpha(x)$ . The constants may however depend logarithmically or linearly on  $\delta_i/h$ . This depends on the geometry and shape of the paths  $P_v$ and on the size and shape of the set  $X_i$ . For more details see [2] and [10, 11].

The following proposition [2, Theorem 3.2] is the central result in our analysis. 138 It proves the stability and a weak approximation property for a local projection onto 139 the span of the first  $m_i$  eigenvectors. 140

**Proposition 1.** Let Assumption 1 hold, and for any  $u \in V_h(\Omega_i)$ , define the projection 141  $\Pi_{i}u := \sum_{\ell=1}^{m_{j}} a_{i}(v_{\ell}^{(j)}, u) v_{\ell}^{(j)}$ . Then 142

$$|\Pi_j u|_{a,\Omega_i} \le |u|_{a,\Omega_i} \quad and \tag{11}$$

$$||u - \Pi_j u||_{0,\alpha,\Omega_j^{\circ}} \lesssim \sqrt{c_j(m_j)} \,\delta_j \,|u|_{a,\Omega_j}. \tag{12}$$

$$||u - \Pi_{j}u||_{0,\alpha,\Omega_{j}^{\circ}} \lesssim \sqrt{c_{j}(m_{j})} \delta_{j} |u|_{a,\Omega_{j}}.$$

$$||u - \Pi_{j}u||_{0,\alpha,\Omega_{j}^{\circ}} \lesssim \sqrt{c_{j}(m_{j})} \delta_{j} |u|_{a,\Omega_{j}}.$$

$$(12)$$

$$where c_{j}(m_{j}) := C_{p}^{2} + \left(\delta_{j}\lambda_{m_{j}+1}^{(j)}\right)^{-1}.$$

As usual (cf. [15]), the following condition number bound can then be obtained 144 via abstract Schwarz theory by constructing a stable splitting.

**Theorem 1.** Let Assumption 1 be satisfied. Then the condition number of the two-146 level Schwarz algorithm with the coarse space  $V_H$  based on local DtN maps and 147 defined in (10) can be bounded by 148

$$\kappa(M_{AS,2}^{-1}A) \lesssim \max_{j=1}^{J} \{c_j(m_j)\} \lesssim C_P^2 + \max_{j=1}^{J} \left(\delta_j \lambda_{m_j+1}^{(j)}\right)^{-1}.$$
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The hidden constant is independent of h,  $\delta_i$ , diam( $\Omega_i$ ), and  $\alpha$ .

*Proof.* We construct a stable splitting for a function  $u \in V_{h,0}$  using the projections  $\Pi_i$ , i = 1, ..., J, in Proposition 1 to define the coarse quasi-interpolant 152

$$u_0 := I_h\left(\sum_{j=1}^J \chi_j \Pi_j u|_{\Omega_j}\right) \in V_H. \tag{13}$$

If we now choose  $u_i := I_h(\chi_i(u - \Pi_i u)) \in V_{h,0}(\Omega_i)$ , then 153

$$u = \sum_{j=0}^J u_j$$
 and  $\sum_{j=0}^J \int_{\Omega} \alpha |\nabla u_j|^2 \lesssim \max_{j=1}^J \{c_j(m_j)\} \int_{\Omega} \alpha |\nabla u|^2$ 

For details see the proof of [2, Theorem 3.5].

Remark 2. Note that by choosing the number  $m_i$  of modes per subdomain such that  $\lambda_{m,+1}^{(j)} \ge \operatorname{diam}(\Omega_j)^{-1}$ , as stated in Sect. 2, we have 157

$$\kappa(M_{AS-1}^{-1}A) \lesssim \left(C_P^2 + \max_i \operatorname{diam}(\Omega_i)/\delta_i\right).$$
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Hence, provided the constant  $C_P$  is uniformly bounded, independently of any jumps 159 in the coefficients, we retrieve the classical estimate for the two-level additive Schwarz method independently of any variations of coefficients across or along subdomain boundaries.

## **4 Numerical Results**

We choose  $\Omega = (0,1)^2$  and discretize (1) on a uniform grid with  $2m^2$  elements, 164 setting u = 0 on the left hand boundary and  $\frac{\partial u}{\partial v} = 0$  on the remainder. We use METIS 165 to split the domain into 16 irregular subdomains as shown in Fig. 1 and construct the 166 overlapping partition by extending each subdomain by one layer of fine grid elements 167 using Freefem++ [5].

As the coarse space we use the DtN coarse space described in Sect. 2 with  $m_i$  169 chosen such that  $\lambda_{m_j}^{(j)} < \operatorname{diam}(\Omega_j)^{-1} \le \lambda_{m_i+1}^{(j)}$ , for all  $j=1,\ldots,16$  (labelled D2N). 170 We compare this preconditioner with the one-level additive Schwarz method (la- 171 belled NONE) and the two-level method with partition of unity coarse space, i.e. 172 choosing  $m_j = 1$  for all j (labelled POU). To confirm in some sense the optimality 173 of our choice for  $m_i$ , we also include results with the DtN coarse space choosing 174  $m_i + 1$  and max $\{1, m_i - 1\}$  basis functions per subdomain (labelled D2N+ and D2N-, 175 respectively). We use the preconditioners within a conjugate gradient iteration with 176 tolerance  $10^{-1}$ .

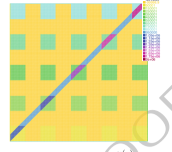
In the first test case (**Example 1**), we choose m = 160 and  $\alpha$  as depicted in 178 Fig. 2, i.e. 25 high permeability inclusions and one channel. In the second test case 179 (Example 2), we choose m = 80 and  $\alpha$  to be a realization of a log-normal distribution 180 with exponential covariance function (variance  $\sigma^2 = 4$  and correlation length  $\lambda = 181$ 4/m) and mean of log  $\alpha$  equal 3 (cf. Fig. 3).

In Fig. 4 we plot  $||u - \bar{u}||_{\infty}$  for Example 1 against the iteration count, where  $\bar{u}$  is 183 the solution of (3) obtained via a direct solver. Clearly both the one-level and the 184 two-level preconditioner with POU coarse space are not robust. The POU coarse 185 space seems to have hardly any influence at all (520 versus 619 iterations), whereas the new DtN coarse space leads to a robust convergence and a significantly reduced 187 number of iterations of 64.

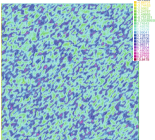
Finally, in Table 1 we compare the different preconditioners and show that the 189 criterion for the number  $m_i$  of eigenmodes that we select in each subdomain is in 190 some sense optimal. Adding one more functions has hardly any impact on the performance while removing one has a strong negative impact. See [2] for more extensive 192 numerical experiments.



Fig. 1. Partition into 16 subdomains



**Fig. 2.** Example 1 (max<sub>x,y</sub>  $\frac{\alpha(x)}{\alpha(y)}$  $= 2 \cdot 10^6$ )



**Fig. 3.** Example 2 (max<sub>x,y</sub>  $\frac{\alpha(x)}{\alpha(y)}$ 

Fig. 4. Convergence history (Example 1)

	Co	pace siz	$V_H$	# PCG Iterations (tol = $10^{-7}$ )						
	NONE	POU	D2N-	D2N	D2N+	NONE	POU	D2N-	D2N	D2N+
Example 1	0	16	32	46	62	619	520	446	64	37
Example 2	0	16	82	98	114	89	92	50	38	36

Table 1. Comparison of DtN coarse space against simple POU coarse space and no coarse space, as well as demonstration of "optimality" of automatic criterion for choosing  $\{m_i\}$ .

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t1.1 t1.2 t1.3 t1.4

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# **Heterogeneous Domain Decomposition Methods** for Eddy Current Problems

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Summary. The usual setting of an eddy current problem distinguishes between a conducting region and an air region (non-conducting) surrounding the conductor. For the numerical 8 approximation of this heterogeneous problem it is very natural to use iterative substructur- 9 ing methods based on transmission conditions at the interface. We analyze the convergence 10 of the Dirichlet-Neumann iterative method for two different formulations of the eddy current problem: the one that consider as main unknown the electric field and the one based on the 12 magnetic field.

1 Introduction 14

To model the electromagnetic phenomena concerning alternating currents at low frequencies it is often used the time-harmonic eddy current model (see e.g. [2]). The 16 main equations of this model are Faraday's law 17

$$\operatorname{curl} \mathbf{E} = -i\omega \mu \mathbf{H} \quad \text{in } \Omega \,, \tag{1}$$

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and Ampère's law

$$\operatorname{curl} \mathbf{H} = \sigma \mathbf{E} + \mathbf{J}_e \quad \text{in } \Omega \,, \tag{2}$$

where  $\mathbf{E}$ ,  $\mathbf{H}$  and  $\mathbf{J}_e$  denote the electric field, the magnetic field and the applied current density respectively. For the sake of simplicity we assume that the computational do- 20 main  $\Omega\subset\mathbb{R}^3$  is a simply connected Lipschitz polyhedron with connected boundary 21 that contains a conducting region  $\Omega_C \subset\subset \Omega$  and that both  $\Omega_C$  and its complement 22  $\Omega_I := \Omega \setminus \overline{\Omega_C}$  are connected Lipschitz polyhedra. Let us denote  $\Gamma := \overline{\Omega_C} \cap \overline{\Omega_I}$ . The 23 magnetic permeability  $\mu$  is assumed to be a symmetric uniformly positive definite 24  $3 \times 3$  matrix with entries in  $L^{\infty}(\Omega)$ , whereas the electric conductivity  $\sigma$  is supposed 25 to be a bounded symmetric positive definite matrix in the conducting regions, and to 26 be null in non-conducting regions. The real scalar constant  $\omega \neq 0$  is a given angular frequency. In  $\partial\Omega$  suitable boundary conditions must be assigned. Most often the 28 tangential component of either the electric field  $\mathbf{E} \times \mathbf{n}$  or the magnetic field  $\mathbf{H} \times \mathbf{n}$  are 29 given (here **n** denotes the unit outward normal vector on  $\partial \Omega$ ).

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Let us introduce some notations that will be used in the following. The space 31  $H(\operatorname{curl};\Omega)$  indicates the set of real or complex vector valued functions  $\mathbf{v}\in(L^2(\Omega))^3$  32 such that  $\operatorname{curl}\mathbf{v}\in(L^2(\Omega))^3$  and  $H^0(\operatorname{curl};\Omega)$  its subspace constituted by  $\operatorname{curl}$ -free 33 functions. Given a certain subset  $\Lambda\subset\partial\Omega$ , we denote by  $H_{0,\Lambda}(\operatorname{curl};\Omega)$  the subspace of functions in  $H(\operatorname{curl};\Omega)$  such that their tangential trace is null on  $\Lambda$ , and in 35 particular we write  $H_0(\operatorname{curl};\Omega):=H_{0,\partial\Omega}(\operatorname{curl};\Omega)$ .

We recall the spaces  $H^{-1/2}(\operatorname{curl}_{\tau};\partial\Omega):=\left\{(\mathbf{n}\times\mathbf{v}\times\mathbf{n})_{|\partial\Omega}\,|\,\mathbf{v}\in H(\operatorname{curl};\Omega)\right\}$ , 37 and  $H^{-1/2}(\operatorname{div}_{\tau};\partial\Omega):=\left\{(\mathbf{v}\times\mathbf{n})_{|\partial\Omega}\,|\,\mathbf{v}\in H(\operatorname{curl};\Omega)\right\}$ , (see [4]). These two spaces 38 are in duality and the following formula of integration by parts holds true

$$\int_{\Omega} \left( \mathbf{w} \cdot \operatorname{curl} \overline{\mathbf{v}} - \operatorname{curl} \mathbf{w} \cdot \overline{\mathbf{v}} \right) = \left\langle \mathbf{w} \times \mathbf{n}, \mathbf{n} \times \overline{\mathbf{v}} \times \mathbf{n} \right\rangle_{\partial \Omega} \quad \forall \, \mathbf{w}, \, \mathbf{v} \in \mathcal{H}(\operatorname{curl}; \Omega) \,. \tag{40}$$

## 2 One Field Formulations

First we notice that Eqs. (1) and (2) do not completely determine the electric field in  $\Omega_I$  and it is necessary to require the gauge condition 43

$$\operatorname{div}\mathbf{E}_{I} = 0 \text{ in } \Omega_{I}. \tag{3}$$

41

(Here and in the sequel, given any vector field  $\mathbf{v}$  defined in  $\Omega$ , we denote  $\mathbf{v}_L$  its 44 restriction to  $\Omega_L$ , L=C,L.) When imposing electric boundary conditions,  $\mathbf{E} \times \mathbf{n} = \mathbf{0}$  45 on  $\partial \Omega$ , in order to have a unique solution we need to impose the additional gauge 46 condition  $\int_{\Gamma} \mathbf{E}_I \cdot \mathbf{n} = 0$ .

From Faraday law  $\mu^{-1}$  curl  $\mathbf{E} = -i\omega\mathbf{H}$  and replacing in Ampère law one has 48 curl  $(\mu^{-1}\text{curl }\mathbf{E}) = -i\omega(\sigma\mathbf{E} + \mathbf{J}_e)$ . So the E-based formulation of the eddy current 49 problem with electric boundary conditions reads

$$\begin{array}{ll} \operatorname{curl} \left( \mu^{-1} \operatorname{curl} \mathbf{E} \right) + i \omega \sigma \mathbf{E} = -i \omega \mathbf{J}_{e} & \text{in } \Omega \\ \operatorname{div} \mathbf{E}_{I} = 0 & \text{in } \Omega_{I} \\ \int_{\Gamma} \mathbf{E}_{I} \cdot \mathbf{n} = 0 & \text{on } \partial \Omega \,. \end{array}$$

Since  $\sigma \equiv 0$  in the non-conducting region, the generator current has to satisfy the 52 compatibility conditions  $\operatorname{div} \mathbf{J}_{e,I} = 0$  in  $\Omega_I$  and, when imposing  $\mathbf{E} \times \mathbf{n} = 0$  on  $\partial \Omega$ , 53  $\int_{\Gamma} \mathbf{J}_{e,I} \cdot \mathbf{n} = 0$ .

Notice that the two gauge conditions  $\operatorname{div} \mathbf{E}_I = 0$  and  $\int_{\Gamma} \mathbf{E}_I \cdot \mathbf{n} = 0$  are equivalent 55 to  $\int_{\Omega_I} \mathbf{E}_I \cdot \nabla \overline{\phi}_I = 0$  for all  $\phi_I \in H^1_*(\Omega_I)$  being  $H^1_*(\Omega_I) = \{\phi_I \in H^1(\Omega_I) : \phi_{I|\partial\Omega} \equiv 56 \text{ and } \phi_{I|\Gamma} \text{ is constant}\}$ . Hence the weak form of the E-based formulation is

Find 
$$\mathbf{E} \in W$$
 such that
$$\int_{\Omega} (\mu^{-1} \operatorname{curl} \mathbf{E} \cdot \operatorname{curl} \overline{\mathbf{w}} + i\omega \sigma \mathbf{E} \cdot \overline{\mathbf{w}}) = -i\omega \int_{\Omega} \mathbf{J}_{e} \cdot \overline{\mathbf{w}}$$
for all  $\mathbf{w} \in W$ 

where 
$$W:=\{\mathbf{w}\in H_0(\mathrm{curl}\,;\Omega):\,\int_{\Omega_I}\mathbf{w}_I\cdot\nabla\overline{\phi}_I=0\,\,\forall\,\phi_I\in H^1_*(\Omega_I)\}.$$

Remark 1. The gauge conditions can be imposed by means of a Lagrange multiplier. 60 (See [2], Sect. 4.6.)

Due to the heterogeneous nature of the problem, it is natural to consider an it- 62 erative procedure by subdomains in order to deal with homogeneous problem. A 63 procedure of this kind is the following: 64

Given 
$$\lambda^{(0)} \in H^{-1/2}(\operatorname{curl}_{\tau}; \Gamma)$$
 for  $n \geq 0$   
find  $\mathbf{E}_{I}^{(n+1)} \in W_{I}$  such that
$$\mathbf{n} \times \mathbf{E}_{I}^{(n+1)} \times \mathbf{n} = \lambda^{(n)} \text{ on } \Gamma$$

$$\int_{\Omega_{I}} \mu^{-1} \operatorname{curl} \mathbf{E}_{I}^{(n+1)} \cdot \operatorname{curl} \overline{\mathbf{w}}_{I} = -i\omega \int_{\Omega_{I}} \mathbf{J}_{e,I} \cdot \overline{\mathbf{w}}_{I} \quad \forall \mathbf{w}_{I} \in W_{I} \cap H_{0}(\operatorname{curl}; \Omega_{I});$$
find  $\mathbf{E}_{C}^{(n+1)} \in H(\operatorname{curl}; \Omega_{C})$  such that
$$\int_{\Omega_{C}} (\mu^{-1} \operatorname{curl} \mathbf{E}_{C}^{(n+1)} \cdot \operatorname{curl} \overline{\mathbf{w}}_{C} + i\omega \sigma \mathbf{E}_{C}^{(n+1)} \cdot \overline{\mathbf{w}}_{C}) = -i\omega \int_{\Omega_{C}} \mathbf{J}_{e,C} \cdot \overline{\mathbf{w}}_{C}$$

$$-\langle \mu^{-1} \operatorname{curl} \mathbf{E}_{I}^{(n+1)} \times \mathbf{n}_{I}, \mathbf{n} \times \mathbf{w}_{C} \times \mathbf{n} \rangle_{\Gamma} \quad \forall \mathbf{w}_{C} \in H(\operatorname{curl}; \Omega_{C});$$
set

 $\boldsymbol{\lambda}^{(n+1)} = (1-\theta)\boldsymbol{\lambda}^{(n)} + \theta(\mathbf{n} \times \mathbf{E}_C^{(n+1)} \times \mathbf{n})|_{\Gamma},$ 

where  $W_I := \{ \mathbf{w}_I \in H_{0,\partial\Omega}(\operatorname{curl};\Omega_I) : \int_{\Omega_I} \mathbf{w}_I \cdot \nabla \overline{\phi}_I = 0 \ \forall \ \phi_I \in H^1_*(\Omega_I) \}, \ \mathbf{n}_I \ \text{denotes}$  66 the unit normal vector on  $\Gamma$  pointing outwards  $\Omega_I$  and  $\theta$  is a positive acceleration parameter.

Another possibility is to eliminate the electric field. Multiplying Faraday law by a function  $\mathbf{v} \in H_0(\text{curl }; \Omega)$  with  $\text{curl } \mathbf{v}_I = 0;$ 

$$i\omega \int_{\Omega} \mu \mathbf{H} \cdot \overline{\mathbf{v}} = -\int_{\Omega} \operatorname{curl} \mathbf{E} \cdot \overline{\mathbf{v}} = -\int_{\Omega} \mathbf{E} \cdot \operatorname{curl} \overline{\mathbf{v}}$$

$$= -\int_{\Omega_{C}} \sigma^{-1} (\operatorname{curl} \mathbf{H}_{C} - \mathbf{J}_{e,C}) \cdot \operatorname{curl} \overline{\mathbf{v}}_{C}.$$
<sup>71</sup>

Given  $\mathbf{g}_I \in (L^2(\Omega_I))^3$  let  $V(\mathbf{g}_I)$  denotes the space  $V(\mathbf{g}_I) := \{\mathbf{v} \in H_0(\operatorname{curl};\Omega) : 72 \operatorname{curl} \mathbf{v}_I = \mathbf{g}_I\}$ . The weak form of **H**-based formulation of the eddy current problem 73 with magnetic boundary conditions  $\mathbf{H} \times \mathbf{n} = \mathbf{0}$  on  $\partial \Omega$  reads

Find 
$$\mathbf{H} \in V(\mathbf{J}_{e,I})$$
 such that
$$\int_{\Omega_C} \sigma^{-1} \operatorname{curl} \mathbf{H} \cdot \operatorname{curl} \overline{\mathbf{v}} + i\omega \int_{\Omega} \mu \mathbf{H} \cdot \overline{\mathbf{v}} = \int_{\Omega_C} \sigma^{-1} \mathbf{J}_{e,C} \cdot \operatorname{curl} \overline{\mathbf{v}}_C$$
(4) for all  $\mathbf{v} \in V(\mathbf{0})$ .

Since  $\sigma \equiv 0$  in the non-conducting region, when imposing  $\mathbf{H} \times \mathbf{n} = 0$  on  $\partial \Omega$  the 75 generator current has to satisfy the compatibility conditions  $\operatorname{div} \mathbf{J}_{e,I} = 0$  in  $\Omega_I$  and 76  $\mathbf{J}_{e,I} \cdot \mathbf{n} = 0$  on  $\partial \Omega$ . Hence there exists  $\mathbf{H}_{e,I}^* \in H_{0,\partial\Omega}(\operatorname{curl};\Omega_I)$  such that  $\operatorname{curl} \mathbf{H}_{e,I}^* = 77$   $\mathbf{J}_{e,I}$ . Then we can write  $\mathbf{H}_I = \mathbf{H}_{e,I}^* + \mathbf{Z}_I$  with  $\mathbf{Z}_I \in H_{0,\partial\Omega}^0(\operatorname{curl};\Omega_I)$ . Let  $\mathbf{H}_e^*$  be a func-78 tion in  $H(\operatorname{curl};\Omega)$  such that  $\mathbf{H}_{e|\Omega_I}^* = \mathbf{H}_{e,I}^*$  and let us denote  $\mathbf{Z} := \mathbf{H} - \mathbf{H}_e^* \in V(\mathbf{0})$ . Mul-79 tiplying Eq. (4) by  $-i\omega^{-1}$  and setting  $\widehat{F}(\mathbf{v}) := \int_{\Omega} \mu \mathbf{H}_e^* \cdot \overline{\mathbf{v}} - i\omega^{-1} \int_{\Omega_C} \sigma^{-1} \operatorname{curl} \mathbf{H}_e^* \cdot \mathbf{s}$  ocurl  $\overline{\mathbf{v}}$ , we can consider the equivalent problem

Find  $\mathbf{Z} \in V(\mathbf{0})$  such that

$$\int_{\Omega} \mu \mathbf{Z} \cdot \overline{\mathbf{v}} - i\omega^{-1} \int_{\Omega_C} \sigma^{-1} \operatorname{curl} \mathbf{Z} \cdot \operatorname{curl} \overline{\mathbf{v}} = -i\omega^{-1} \int_{\Omega_C} \sigma^{-1} \mathbf{J}_{e,C} \cdot \operatorname{curl} \overline{\mathbf{v}}_C - \widehat{F}(\mathbf{v})$$
 so for all  $\mathbf{v} \in V(\mathbf{0})$ .

For the sake of simplicity we will assume that  $\mathbf{J}_{e,I} \cdot \mathbf{n} = 0$  on  $\Gamma$ . Then it is possible 83 to take  $\mathbf{H}_{e,I}^* \in H_0(\text{curl}; \Omega_I)$  and  $\mathbf{H}_{e,C}^*$  equal zero.

Remark 2. Notice that  $H^0_{0,\partial\Omega}(\operatorname{curl};\Omega_I) = \nabla H^1_{0,\partial\Omega}(\Omega_I) \oplus \mathscr{H}(\Omega_I)$  where  $\mathscr{H}(\Omega_I) := 85$  $\{\mathbf v_I \in H^0_{0 \partial \Omega}(\operatorname{curl};\Omega_I) : \operatorname{div} \mathbf v_I = 0 \text{ and } \mathbf v_I \cdot \mathbf n = 0 \text{ on } \Gamma\}$  that is a space of finite dimension. In this geometrical setting the dimension of  $\mathcal{H}(\Omega_I)$  coincides with the 87 first Betti number of  $\Omega_I$ . (See [2], Sect. 5.1.)

We propose an iterative procedure for the solution of the H-based formulation 89 that start from a data in the trace space

$$H_0^{-1/2}(\operatorname{curl}_{\tau};\Gamma) := \left\{ (\mathbf{n} \times \mathbf{w}_I \times \mathbf{n})_{|\Gamma} : \mathbf{w}_I \in H_{0,\partial\Omega}^0(\operatorname{curl};\Omega_I) \right\}. \tag{91}$$

It reads: 92

S: 
$$\begin{aligned} &\text{Given } \boldsymbol{\lambda}^{(0)} \in H_0^{-1/2}(\operatorname{curl}_{\tau}; \boldsymbol{\Gamma}) \text{ for } n \geq 0 \\ &\text{find } \mathbf{H}_C^{(n+1)} \in H(\operatorname{curl}; \boldsymbol{\Omega}_C) \text{ such that} \\ &\mathbf{n} \times \mathbf{H}_C^{(n+1)} \times \mathbf{n} = \boldsymbol{\lambda}^{(n)} \quad \text{on } \boldsymbol{\Gamma} \\ &\int_{\boldsymbol{\Omega}_C} (\boldsymbol{\mu} \mathbf{H}_C^{(n+1)} \cdot \overline{\mathbf{v}}_C - i\boldsymbol{\omega}^{-1} \boldsymbol{\sigma}^{-1} \operatorname{curl} \mathbf{H}_C^{(n+1)} \cdot \operatorname{curl} \overline{\mathbf{v}}_C) \\ &= -i\boldsymbol{\omega}^{-1} \int_{\boldsymbol{\Omega}_C} \boldsymbol{\sigma}^{-1} \mathbf{J}_{e,C} \cdot \operatorname{curl} \overline{\mathbf{v}}_C \quad \forall \mathbf{v}_C \in H_0(\operatorname{curl}; \boldsymbol{\Omega}_C); \end{aligned}$$
 find  $\mathbf{Z}_I^{(n+1)} \in H_{0,\partial\boldsymbol{\Omega}}^0(\operatorname{curl}; \boldsymbol{\Omega}_I) \text{ such that}$  
$$\int_{\boldsymbol{\Omega}_I} \boldsymbol{\mu} \mathbf{Z}_I^{(n+1)} \cdot \overline{\mathbf{v}}_I = i\boldsymbol{\omega}^{-1} \langle \boldsymbol{\sigma}^{-1}(\operatorname{curl} \mathbf{H}_C^{(n+1)} - \mathbf{J}_{e,C}) \times \mathbf{n}_C, \mathbf{n} \times \mathbf{v}_I \times \mathbf{n} \rangle_{\boldsymbol{\Gamma}} \\ &- \int_{\boldsymbol{\Omega}_I} \boldsymbol{\mu} \mathbf{H}_{e,I}^* \cdot \overline{\mathbf{v}}_I \quad \forall \mathbf{v}_{I,h} \in H_{0,\partial\boldsymbol{\Omega}}^0(\operatorname{curl}; \boldsymbol{\Omega}_I); \end{aligned}$$
 set 
$$\boldsymbol{\lambda}^{(n+1)} = (1-\boldsymbol{\theta}) \boldsymbol{\lambda}^{(n)} + \boldsymbol{\theta} (\mathbf{n} \times \mathbf{Z}_I^{(n+1)} \times \mathbf{n})_{|\boldsymbol{\Gamma}},$$

being  $\mathbf{n}_C$  the unit normal vector on  $\Gamma$  pointing outwards  $\Omega_C$  and  $\theta$  a positive acceleration parameter. 95

# 3 Convergence Analysis

Both the H-based formulation and the E-based formulation are of the form: find 97  $\mathbf{u} \in V \subset H(\text{curl}; \Omega)$  such that 98

$$a(\mathbf{u}, \mathbf{v}) = F(\mathbf{v}) \quad \forall \mathbf{v} \in V,$$
 (5)

96

where  $a(\cdot,\cdot)$  is a sesquilinear form continuous and coercive in  $V \times V$  and  $F(\cdot)$  99 is a continuous linear functional on the Hilbert space V. The proposed iterative 100

procedures are preconditioned Richardson methods for the Steklov-Poincare equa- 101 tion obtained in the following way (see e.g. [8]): for L = C, I let us define the 102 spaces  $V_L := \{\mathbf{v}_{|\Omega_L} : \mathbf{v} \in V\}, X := \{(\mathbf{n} \times \mathbf{v} \times \mathbf{n})_{\Gamma} : \mathbf{v} \in V\}$  and  $V_{L,0} := \{\mathbf{v}_L \in \mathsf{los}\}$  $V_L: (\mathbf{n} \times \mathbf{v}_L \times \mathbf{n})_{\Gamma} = \mathbf{0}$ ; the sesquilinear forms  $a_L(\cdot, \cdot): V_L \times V_L \to \mathbb{C}$  and the 104 linear functionals  $F_L: V_L \to \mathbb{C}$  such that  $a(\mathbf{v}, \mathbf{w}) = a_C(\mathbf{v}_C, \mathbf{w}_C) + a_I(\mathbf{v}_I, \mathbf{w}_I)$  and 105  $F(\mathbf{v}) = F_C(\mathbf{v}_C) + F_I(\mathbf{v}_I) \quad \forall \mathbf{v}, \mathbf{w} \in V.$  If the sesquilinear forms  $a_L(\cdot, \cdot)$  are continuous and coercive in  $V_{L,0}$  for both L=C,I we can define the extension operators 107  $\mathbf{R}_L: X \to V_L$  in the following way: for any  $\boldsymbol{\eta} \in X$ ,  $\mathbf{R}_L \boldsymbol{\eta}$  is the unique function in  $V_L$  108 such that

$$(\mathbf{n} \times \mathbf{R}_L \boldsymbol{\eta} \times \mathbf{n})|_{\Gamma} = \boldsymbol{\eta}$$
  
 $a_L(\mathbf{R}_L \boldsymbol{\eta}, \mathbf{v}_L) = 0 \quad \forall \mathbf{v}_L \in V_{L,0}.$ 

111

113

Let us consider the Steklov-Poincare operators  $S_L: X \to X'$  given by

$$\langle S_L \boldsymbol{\eta}, \boldsymbol{v} \rangle_{\Gamma} = a_L(\mathbf{R}_L \boldsymbol{\eta}, \mathbf{R}_L \boldsymbol{v}) \quad \forall \, \boldsymbol{\eta} \,, \, \boldsymbol{v} \in X \,.$$

Moreover we can define the functions  $\hat{\mathbf{u}}_L \in V_{L,0}$  such that

$$a_L(\hat{\mathbf{u}}_L, \mathbf{v}_L) = F_L(\mathbf{v}_L) \quad \forall \mathbf{v}_L \in V_{L,0}$$

and  $\chi_L \in X'$  given by  $\langle \chi_L, \eta \rangle_{\Gamma} = F_L(\mathbf{R}_L \eta) - a_L(\hat{\mathbf{u}}_L, \mathbf{R}_L \eta) \quad \forall \eta \in X$ . Let us denote 115  $\chi = \chi_I + \chi_C$ . The Steklov-Poincare equation reads: find  $\lambda \in X$  such that 116

$$(S_I + S_C)\lambda = \chi. \tag{6}$$

$$(S_I + S_C) \boldsymbol{\lambda} = \boldsymbol{\chi}. \tag{6}$$
If  $\boldsymbol{\lambda}$  is solution of (6) then  $\mathbf{u} = \begin{cases} \mathbf{R}_C \boldsymbol{\lambda} + \hat{\mathbf{u}}_C \text{ in } \Omega_C \\ \mathbf{R}_I \boldsymbol{\lambda} + \hat{\mathbf{u}}_I \text{ in } \Omega_I \end{cases}$  is solution of (5).

If for one of the two subdomains the sesquilinear form  $a_L(\cdot,\cdot)$  is also continuous 118 and coercive in  $V_L$  then for each  $\boldsymbol{\xi} \in X'$  there exist a unique  $\mathbf{F}_L \boldsymbol{\xi} \in V_L$  such that  $a_L(\mathbf{F}_L\boldsymbol{\xi},\mathbf{w}_L) = \langle \boldsymbol{\xi},\mathbf{n} \times \mathbf{w}_L \times \mathbf{n} \rangle_{\Gamma} \quad \forall \mathbf{w}_L \in V_L.$  It is easy to see that  $\langle S_L(\mathbf{n} \times \mathbf{F}_L\boldsymbol{\xi} \times 120) \rangle_{\Gamma}$  $|\mathbf{n}\rangle, \boldsymbol{\eta}\rangle_{\Gamma} = \langle \boldsymbol{\xi}, \boldsymbol{\eta}\rangle_{\Gamma}$  for all  $\boldsymbol{\eta} \in X$  hence  $S_L^{-1}(\boldsymbol{\xi}) = \mathbf{n} \times \mathbf{F}_L \boldsymbol{\xi} \times \mathbf{n}$ . It is well known that the Dirichlet-Neumann iterative method is equivalent to the preconditioned Richard- 122 son method for the Steklov-Poincare equation

$$oldsymbol{\lambda}^{(n+1)} = oldsymbol{\lambda}^{(n)} + heta S_L^{-1} \left[ oldsymbol{\chi} - (S_I + S_C) oldsymbol{\lambda}^{(n)} 
ight].$$
 124

In the **H**-based formulation the preconditioner is  $S_I$  while in the **E**-based formulation 125 the preconditioner is  $S_C$ .

We are interested in the finite element approximation of these problems using the 127 Nédélec curl-conforming edge elements of degree k,  $N_{L,h}^k \subset H(\text{curl}; \Omega_L)$  (see [7]) for 128 L = C, I. Let us denote  $\P_k$ ,  $k \ge 0$ , the space of polynomials of degree less than or equal k in the three variables  $x_1, x_2, x_3$ , and by  $\P_k$  the space of homogeneous polynomials of degree k. For  $k \ge 1$  we define the polynomial spaces  $M_k := \{ \mathbf{q} \in (\widetilde{\P}_k)^3 \mid \mathbf{q}(\mathbf{x}) \cdot \mathbf{x} = 0 \}$  131 and  $R_k := (\P_{k-1})^3 \oplus M_k$ . Let us consider a tetrahedral triangulation of  $\Omega$ ,  $\mathscr{T}_h$ , such that its restriction to  $\Omega_L$ ,  $\mathcal{I}_{Lh}$ , induces a triangulation of  $\Omega_L$ . Then 133

$$N_{L,h} := \{ \mathbf{w}_h \in H(\operatorname{curl}; \Omega_L) \, | \, \mathbf{w}_{h|K} \in R_k \quad \forall \, K \in \mathscr{T}_{L,h} \} \,. \tag{134}$$

We want to show that in the discrete setting the iterative procedure converges and 135 that the convergence rate is independent of h.

The discrete **H**-based formulation is stated in the space 137

$$V_h(\mathbf{0}) := \{ \mathbf{v}_h \in N_h^k : \mathbf{v}_{Lh} \in H_{0,\partial\Omega}^0(\operatorname{curl};\Omega_I) \} \subset V(\mathbf{0}).$$
 138

The space *X* for the Dirichlet-Neumann procedure is

$$\chi_h^0 = \{ (\mathbf{n} \times \mathbf{v}_h \times \mathbf{n})_{|\Gamma} : \mathbf{v}_h \in V_h(\mathbf{0}) \} \subset H_0^{-1/2}(\operatorname{curl}_{\tau}; \Gamma).$$
 140

In  $\Omega_C$  we use the standard Nédélec finite elements  $N_{C.h}^k$ , while in  $\Omega_I$  we have the 141 finite element space 142

$$V_{I,h}(\mathbf{0}) = N_{I,h}^k \cap H_{0,\partial\Omega}^0(\operatorname{curl};\Omega_I)$$
.

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144

Remark 3. Let  $L^k_{I,h}\subset H^1(\Omega_I)$  be the space of standard Lagrange finite elements of degree k and  $H_{I,h,0} = L_{I,h}^k \cap H_{0,\partial\Omega}^1(\Omega_I)$ . Then 146

$$V_{I,h}(\mathbf{0}) = \nabla H_{I,h,0} + \mathcal{H}_{I,h}$$

where  $\mathcal{H}_{I,h}$  is a space whose dimension coincides with  $n_{\Gamma}$ , the first Betti number of 148  $\Omega_I$ . More precisely, there exits a system of cutting surfaces  $\Xi_I$ ,  $I=1,\ldots,n_\Gamma$  with 149  $\partial \Xi_l \subset \Gamma$  such that every function  $\mathbf{v}_I \in H_{0,\partial\Omega}(\operatorname{curl};\Omega_I)$  restricted to  $\Omega_I \setminus \cup_{l=1}^{n_\Gamma} \Xi_l$  is 150 the gradient of a function belonging to  $H^1(\Omega_I \setminus \bigcup_{l=1}^{n_\Gamma} \Xi_l)$  (see e.g. [3, 5, 6]). If the triangulation  $\mathcal{I}_{I,h}$  induces a triangulation on each surface  $\Xi_l$  the space  $\mathscr{H}_{I,h}$  is the one generated by the  $(L^2(\Omega_I))^3$ -extension of the gradient of the piecewise linear 153 function taking value one at the node on one side of  $\Xi_l$  and value zero at all the other 154 nodes including those on the other side of  $\Xi_l$  (see [2], Sect. 5.4). 155

Concerning the E-based formulation, for its finite element approximation we 156 consider the space 157

$$W_h := \{ \mathbf{w}_h \in N_h^k : \int_{\Omega_I} \mathbf{w}_h \cdot \mathbf{\nabla} \overline{\phi}_{I,h} = 0 \quad orall \, \phi_{I,h} \in H_{I,h,*}^k \}$$
 158

where  $H_{I,h,*}^k = L_{I,h}^k \cap H_*^1(\Omega_I)$ . (Notice that  $W_h$  is not a subspace of W.) The space X where the Steklov-Poincare operators are defined is the space of discrete traces 159 160

$$\chi_h = \{ (\mathbf{n} \times \mathbf{w}_h \times \mathbf{n})_{|\Gamma} : \mathbf{w}_h \in N_h^k \} \subset H^{-1/2}(\operatorname{curl}_{\tau}; \Gamma).$$

Also in this case we use the standard Nédélec finite elements  $N_{C,h}^k$  in  $\Omega_C$  while in  $\Omega_I$ we consider the finite element space 163

$$W_{I,h} := \{ \mathbf{w}_{I,h} \in N_{I,h}^k : \int_{\Omega_I} \mathbf{w}_{I,h} \cdot \nabla \overline{\phi}_{I,h} = 0 \quad \forall \, \phi_{I,h} \in H_{I,h,*}^k \}.$$

In order to prove the convergence of the iterative procedure let us proceed as in 165 [1]. If  $k \in \mathbb{C}$  is an eigenvalue of the map  $T_L: X \to X$ ,  $T_L \boldsymbol{\eta} := \boldsymbol{\eta} - \theta S_L^{-1}(S_I + S_C) \boldsymbol{\eta}$  166 with L=I or L=C, then  $k=1-\theta\frac{\langle (S_I+S_C)\pmb{\eta},\pmb{\eta}\rangle_{\Gamma}}{\langle S_L\pmb{\eta},\pmb{\eta}\rangle_{\Gamma}}=(1-\theta)-\theta\frac{\langle S_M\pmb{\eta},\pmb{\eta}\rangle_{\Gamma}}{\langle S_L\pmb{\eta},\pmb{\eta}\rangle_{\Gamma}}$  for any 167 eigenvector  $\pmb{\eta}\in X$ . Here M=I or M=C but  $M\neq L$ . If

$$\operatorname{Re}[\langle S_{I}\boldsymbol{\eta},\boldsymbol{\eta}\rangle_{\Gamma}]\operatorname{Re}[\langle S_{C}\boldsymbol{\eta},\boldsymbol{\eta}\rangle_{\Gamma}] + \operatorname{Im}[\langle S_{I}\boldsymbol{\eta},\boldsymbol{\eta}\rangle_{\Gamma}]\operatorname{Im}[\langle S_{C}\boldsymbol{\eta},\boldsymbol{\eta}\rangle_{\Gamma}] \geq 0 \tag{7}$$

and  $0 \le \theta \le 1$  then

$$|k|^2 \leq (1-\theta)^2 + \theta^2 \frac{|\langle S_M \boldsymbol{\eta}, \boldsymbol{\eta} \rangle_{\varGamma}|^2}{|\langle S_L \boldsymbol{\eta}, \boldsymbol{\eta} \rangle_{\varGamma}|^2} \leq (1-\theta)^2 + \theta^2 \frac{\beta_M^2}{\alpha_L^2}$$
 170

being  $\beta_M$  the continuity constant of  $S_M$  and  $\alpha_L$  the coercivity constant of  $S_L$ . Choosing  $0 < \theta < \min\left(1, \frac{2\alpha_L^2}{\alpha_L^2 + \beta_M^2}\right)$  on has |k| < 1 for each k eigenvalue of T, hence in the discrete setting the Dirichlet-Neumann procedures converges and, if  $\alpha_L$  and  $\beta_M$  are independent of the mesh size, h, also the convergence rate is independent of h.

In the **H**-based formulation we have L = I and M = C. The sesquilinear form

$$a_C(\mathbf{v}_C, \mathbf{w}_C) := \int_{\Omega_C} \left( -i\omega^{-1} \sigma^{-1} \operatorname{curl} \mathbf{v}_C \cdot \operatorname{curl} \overline{\mathbf{w}}_C + \mu \mathbf{v}_C \cdot \overline{\mathbf{w}}_C \right)$$
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is clearly continuous and coercive in  $H(\operatorname{curl};\Omega_C)$  hence in  $N_{C,h}^k$ . In the insulator  $a_I(\mathbf{v}_I,\mathbf{w}_I):=\int_{\Omega_I}\mu\mathbf{v}_I\cdot\overline{\mathbf{w}}_I$  is continuous and coercive in  $H^0(\operatorname{curl};\Omega_I)$  then also in 178  $V_{I,h}^0$ . The coercivity of  $S_I$  with a constant  $\alpha_I$  independent of h follows from the coercivity of  $a_I(\cdot,\cdot)$  and the continuity of the trace operator while the continuity of  $S_C$  180 with a constant  $\beta_C$  independent of h follows from the continuity of  $a_C(\cdot,\cdot)$  and the existence of a continuous extension operator  $\mathscr{E}_{C,h}:\chi_h\to N_{C,h}^k$  with continuity constant 182 independent of h. Such an extension has been constructed in [1]. Moreover (7) clearly 183 holds because it reduces to  $\left(\int_{\Omega_C}\mu\mathbf{R}_C\boldsymbol{\eta}\cdot\overline{\mathbf{R}_C\boldsymbol{\eta}}\right)\left(\int_{\Omega_I}\mu\mathbf{R}_I\boldsymbol{\eta}\cdot\overline{\mathbf{R}_I\boldsymbol{\eta}}\right)\geq 0$ . Hence taking 184  $\theta$  small enough the iterative Dirichlet-Neumann procedure for the  $\mathbf{H}$ -based formulation converges with a rate independent of the mesh size.

On the other hand for the E-based formulation we have L=C and M=I. Again 187 the sesquilinear form

$$a_C(\mathbf{v}_C, \mathbf{w}_C) := \int_{\Omega_C} \left( \mu^{-1} \operatorname{curl} \mathbf{v}_C \cdot \operatorname{curl} \overline{\mathbf{w}}_C + i\omega \sigma \mathbf{v}_C \cdot \overline{\mathbf{w}}_C \right)$$
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is clearly continuous and coercive in  $H(\operatorname{curl};\Omega_C)$  hence in  $N_{C,h}^k$ . The coercivity of  $S_C$  190 (the preconditioner in this case) with a constant  $\alpha_C$  independent of h follows from the 191 uniform coercivity of  $a_C(\cdot,\cdot)$  and the continuity of the trace operator. In the insulator 192 we have  $a_I(\mathbf{v}_I,\mathbf{w}_I):=\int_{\Omega_I}\mu^{-1}\operatorname{curl}\mathbf{v}_I\cdot\operatorname{curl}\overline{\mathbf{w}}_I$  that is continuous in  $H(\operatorname{curl};\Omega_I)$ , hence 193 in  $W_{I,h}$ . Proceeding as in [2], Sect. 5.5, it can be proved that it is coercive in  $W_{I,h}\cap H_0(\operatorname{curl};\Omega_I)$ . In order to prove the continuity of  $S_I$  with a constant  $\beta_I$  independent 195 of h we need a continuous extension operator  $\mathscr{E}_{I,h}:\chi_h\to W_{I,h}\cap H_{0,\partial\Omega}(\operatorname{curl};\Omega_I)$ . We 196 know that there exists a continuous extension  $\widehat{\mathscr{E}}_{I,h}:\chi_h\to N_{I,h}^k\cap H_{0,\partial\Omega}(\operatorname{curl};\Omega_I)$  (see 197 again [1]). Given  $\eta_h\in\chi_h$  let  $\Phi_{I,h}\in H_{I,h,*}^k$  be such that

$$\int_{\Omega_I} \nabla \Phi_{I,h} \cdot \nabla \psi_{I,h} = \int_{\Omega_I} \widehat{\mathscr{E}}_{I,h} \boldsymbol{\eta}_h \cdot \nabla \psi_{I,h} \quad \forall \, \psi_{I,h} \in H^k_{I,h,*} \,. \tag{199}$$

Then  $\mathscr{E}_{I,h} \boldsymbol{\eta}_h := \widehat{\mathscr{E}}_{I,h} \boldsymbol{\eta}_h - \boldsymbol{\nabla} \boldsymbol{\Phi}_{I,h}$  is a continuous extension from  $\chi_h$  in the space  $W_{I,h} \cap 200$   $H_{0,\partial\Omega}(\operatorname{curl};\Omega_I)$  with continuity constant independent of h. Condition (7) reduce 201 in this case to  $\left(\int_{\Omega_C} \mu^{-1} \operatorname{curl} \mathbf{R}_C \boldsymbol{\eta} \cdot \operatorname{curl} \overline{\mathbf{R}_C \boldsymbol{\eta}}\right) \left(\int_{\Omega_I} \mu^{-1} \operatorname{curl} \mathbf{R}_I \boldsymbol{\eta} \cdot \operatorname{curl} \overline{\mathbf{R}_I \boldsymbol{\eta}}\right) \geq 0$  that 202 clearly holds true.

4 Conclusion 204

We proposed two iterative substructuring methods for two different formulations of the eddy current problem based on the electric field and magnetic field, respectively, and provided the convergence analysis. Both formulations use a constrained space 207 in the insulator. In the **E**-based formulation the constrain is imposed introducing a 208 Lagrange multiplier while in the **H**-based formulation a finite element approximation 209  $V_{I,h}(\mathbf{0})$  of the constrained space  $H_{0,\partial\Omega}(\text{curl};\Omega_I)$  is used. The dimension of  $V_{I,h}(\mathbf{0})$  is 210 equal to  $n_{\Gamma}$ , the dimension of the  $\mathscr{H}_{I,h}$ , plus the dimension of  $H_{I,h,0}$ , that is a space 211 of scalar functions. So the subproblem in the insulator is smaller for the **H**-based 212 formulation than for the **E**-based formulation. However the construction of a base of 213  $\mathscr{H}_{I,h}$  requires the determination of a system of cutting surfaces. This procedure can 214 be cumbersome in complex geometry configurations (for instance if the conductor is 215 a trefoil knot) and the **E** based formulation avoids this difficult.

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# Mesh Regularization in Bank-Holst Parallel hp-Adaptive Meshing

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1 Introduction 9

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In this work, we study mesh regularization in Bank-Holst parallel adaptive paradigm 10 when adaptive enrichment in both h (geometry) and p (degree) is used. The paradigm 11 was first introduced by Bank and Holst in [1-3] and later extended to hp-adaptivity 12 in [5]. In detail, the paradigm can be summarized in the following steps.

Step 1 – Load Balancing: The problem is solved on a coarse mesh, and available a posteriori error estimates are used to partition the mesh into subregions. The 15 partition is such that each subregion has approximately the same error although subregions may vary considerably in terms of number of elements, number of degrees 17 of freedom, and polynomial degree.

Step 2 – Adaptive Meshing: Each processor is provided with complete data for 19 the coarse problem and instructed to sequentially solve the entire problem, with the 20 stipulation that its adaptive enrichment (in h or p) should be limited largely to its 21 own subregion. The target number of degrees of freedom for each processor is the 22

Step 3 – Mesh Regularization: The local mesh on each processor is regularized 24 such that the mesh for the global problem described in Step 4 is conforming in both 25 h and p.

Step 4 – Global Solve: The final global problem consists of the union of the 27 refined partitions provided by each processor. A final solution is computed using 28 domain decomposition.

This paradigm is attractive as it requires low communication and allows existing sequential adaptive finite element codes to run in parallel environment without 31

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much effort in recoding. However, it also poses some challenges in mesh regularization (Step 3). Since the adaptive enrichment on each processor (Step 2) is completely independent of what happens on other processors, the global refined mesh, aconstructed from the meshes associated with the refined regions on each of the processors, is initially non-conforming along the interface system. Thus, we need to efficiently identify and resolve these nonconformities, and ultimately to establish flinks between degrees of freedom on the fine mesh interface system on a given processor and the corresponding degrees of freedom on other processors which share its interface. These tasks are challenging due to the fact that the meshes are unstructured in geometry (in h), have variable degree (variable p), no element refinement tree is 41 available, and nonconformity exists in both h and p.

2 Data Structures

In our implementation of Bank-Holst paradigm in PLTMG, a relaxed version of 44 longest edge bisection h-refinement and a rather flexible p-refinement strategy are 45 used for hp-refinement, see [7].

## 2.1 Boundary Edge Data Structure

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Each boundary edge is represented by a column in the  $6 \times NBF$  integer array IB- 48 NDRY, where NBF is the number of boundary edges. For the Ith column of IBNDRY, 49 four of the six entries contain information about the endpoint vertices, and indication of whether the edges is curved or straight, and a user-defined label. One entry indicates edge type (various boundary condition types, or internal interface), and the fifth entry, nonzero only for edges defining the interface system used in the parallel computation, encodes information which is used in the regularization process. This entry is described in more detail in Sect. 2.2 (Table 1).

Table 1. Boundary edge information

	First vertex number
	Second vertex number
IBNDRY(3,I)	Curved edge
IBNDRY(4,I)	
IBNDRY(5,I)	Parallel information
IBNDRY(6,I)	User label

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<sup>&</sup>lt;sup>3</sup> The term "interface" is used to refer to the system of edges that are shared by two subregions, and the term "boundary" is used to refer to the union of the physical boundary of the domain and the interface.

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## 2.2 Interface Edge Labeling

One approach to solve the nonconformities in the global refined mesh is to build and 57 store refinement trees for all elements. However, such trees lose some of their attractiveness if procedures such as mesh moving and edge flipping destroy some of their 59 properties. In addition, we only need information about the edges on the interface 60 system, which typically is a very small fraction of the total information describing 61 the mesh. Thus, instead of creating refinement trees for all elements, during the regularization phase we recover a refinement tree for each interface edge that defines the initial interface system. To insure that subregions remain geometrically conforming 64 on all processors, we forbid mesh moving and edge flipping for all vertices and edges 65 lying on the interface system.

Only minimal information needed to recover the edge refinement tree is stored 67 for each interface edge. In particular, for each interface edge E, we need the index of  $_{68}$ its original edge r(E) in the interface system of the broadcast coarse mesh (after Step 69 1) and its position in the refinement binary tree s(E). Because the original (interface) 70 edges are the same on all processors, we can first match them, and then their descendants based on their positions in the refinement tree. These two pieces of information 72 are combined to make a single integer, label(E), the parallel information for edge  $E_{73}$ stored in the fifth row of the IBNDRY array:

$$label(E) = r(E) + (s(E) - 1) * base.$$

Here base is an integer which is larger than the number of boundary edges NBF in 75 the broadcast coarse mesh. For edge  $E_{org}$  in the broadcast mesh,  $r(E_{org})$  is its number 76 in the IBNDRY system and  $s(E_{org}) = 1$ . When an edge E is refined into two children 77  $E_1$  and  $E_2$ , their labels are determined from label(E) and the following identities:

$$r(E_1) = r(E_2) = r(E)$$
  
 $s(E_1) = 2 * s(E)$   
 $s(E_2) = 2 * s(E) + 1$ 

For consistency,  $E_1$  and  $E_2$  are ordered in the counterclockwise traversal defined by 79 vertices of E. 80

#### 2.3 Interface Data Structure

When a boundary edge is refined, its entries in IBNDRY are replaced by those of 82 one of its children. Thus IBNDRY contains only refined boundary edges. To recover 83 the refinement trees of the interface edges, first all of the refined edges are sorted 84 in groups according to r(E). The refined edges in each group are then ordered in 85 a counterclockwise traversal of the interface based on their vertices (end points). 86 Edges in each group will be used to recover a refinement tree whose leaves and root 87 represent themselves and their original edge respectively.

In order to illustrate the construction of the refinement tree of edges sharing the 89 same ancestor, we consider the group of all refined edges associated with the original 90

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edge E as shown in Fig. 1. These edges have the same index r(E) and have been 91 ordered via a counterclockwise traversal. For simplicity, only positions of these edges 92 in the binary tree are shown. First, leaf nodes for the refined edges are created. Since 93 the two nodes with largest keys (nodes 15 and 14 in our example) are siblings, their 94 s(E) values are used to create the node of their parent (node 7). Then the parent 95 node for the two nodes with the next largest keys (nodes 10 and 11 in our example) 96 are created and so on. The process is completed when the root node (with key 1) is 97 created.

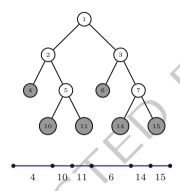


Fig. 1. Refinement tree associated with an original edge

Following the above procedure, we construct the interface data IPATH outlined 99 in Table 2. Each interface edge, including those associated with internal nodes in 100 refinement trees, is represented by a column with six entries in IPATH array. The 101 first entry contains the index r(E) if the edge is original (root) and zero otherwise. 102 When edges from the two sides of the interface are matched, this entry is updated 103 with the index of the corresponding edge. The second entry stores either the index 104 of the edge's first child or its number in IBNDRY array (with minus sign) if it has 105 no child. Sibling edges are put consecutively in IPATH array so storing the index for 106 the second child edge is not necessary. Depending on the stage in the construction 107 of IPATH array, the third and forth entries accommodate the indices of either edges, 108 vertices or degrees of freedom of the two ends of the edge. The fifth entry is either 109 the first or last (with minus sign) index of the interior degree(s) of freedom of the 110 edge. This information together with the degree of the edge stored in the last entry 111 are sufficient to recover all indices of the edge's interior degrees of freedom as they 112 are numbered consecutively. The sign of the fifth entry indicates if they are increase 113 or decrease along the counterclockwise traversal of the interface.

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tree section type root root/leaf internal leaf IPATH(1,\*) -1/n -1/n 0/n0/n IPATH(2,\*) child child -е -е e1/v1/d1 IPATH(3,\*)|e1/v1/d1v1/d1 v1/d1 IPATH(4,\*)|e2/v2/d2v2/d2 e2/v2/d2 v2/d2 IPATH(5,\*)+-d +-d +-d +-d IPATH(6,\*)degree degree degree degree l=label, n=neighbor, e=edge k, v=vertex, d=dof

**Table 2.** Interface data structure: tree section

## 3 Mesh Regularization

The regularization phase requires two all-to-all communication steps. The first de- 116 scribes the initial (non-conforming in h and p) interface system, and the second describes the final conforming system.

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### 3.1 Data Reordering

At the beginning of the regularization step, each processor reorders its data struc- 120 tures. For processor I, edges, vertices and degrees of freedom on the interface be- 121 tween subregion I and the rest of the domain (fine interface) appear first in their 122 respective arrays. These data are also arranged in a counterclockwise traversal of the 123 interface to aid in the creation of the parallel interface data structure IPATH. Next, 124 in all arrays, appears data corresponding to the interior of subregion I (fine interior); 125 typically this is the majority of the data on processor I. Then appears data corre- 126 sponding to the coarse part of the interface system on processor I (the interface not 127 bounding region I). Finally appears data corresponding to the interiors of subregions 128 other than I. Note that the first two blocks of this data (fine interface and fine interior) 129 represent the contribution of processor I to the global fine mesh.

The parallel interface data structure IPATH is arranged in two sections; at the 131 beginning is a pointer section with pointers for each processor's contribution to the 132 fine interface system, and then two special sets of pointers, one for the local coarse 133 interface system and one for the global fine mesh as a whole (see Table 3). The 134 second section contains the tree data for individual edges on the interface system. 135 After regularization, each processor has an IPATH array that contains complete data 136 of the two-sided global fine interface system appended with data of local coarse 137 interface system.

### 3.2 Fine Mesh Regularization

After reordering and a global exchange of interface data, each processor has com- 140 plete information of the fine interface system. Then each process matches its interface edges against those of it neighbors. First original coarse edges are matched 142

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**Table 3.** Interface data structure: pointer section

	pointer section: $1 \rightarrow p+2$
IPATH(1,I)	first interface tree entry for subregion I
IPATH(2,I)	last interface tree entry for subregion I
IPATH(3,I)	first interface vertex/dof for subregion I
IPATH(4,I)	last interface vertex/dof for subregion I
I = p +	- 1: pointers for local coarse system
I=p	+2: pointers for global fine system

based on their labels. Then their descendants are matched following the refinement 143 tree structures. We note here that for two neighboring processors, counterclockwise 144 traversals of the interface are in opposite directions. An example of descendants of 145 two original edges (from two different processors) is shown in Fig. 2.

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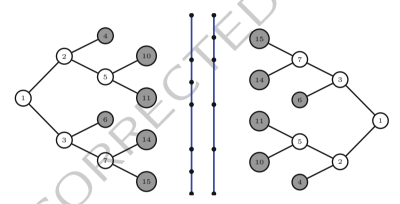


Fig. 2. Edge matching

When a pair of matching edges is determined, their first entries in IPATH are 147 updated to store the indices (also in IPATH array) of their neighbors (change status 148 from "-1" or "0" to "n" as in Table 2). If edges without corresponding neighbors 149 are found, this indicates nonconformity in h. This is resolved by the processor with the less refined interface; it executes appropriate steps of h-refinement to make its 151 interface match that of it neighbor. Although we must allow for arbitrary differences 152 in refinement, it is typical to see at most one level of refinement difference on the fine 153 portion of the interface. An example in Fig. 2 is edge 4 on the left that corresponds 154 to edge 7 on the right with two child edges 14 and 15. In this case, edge 4 on the left 155 will be h-refined one level.

When issues of h-conformity are resolved, the edges are re-examined to eliminate nonconformity in degree. Since the mesh is now h-conforming, each leaf edge 158 on the fine interface system should have exactly one matching neighbor (from an- 159 other processor). If the degrees of a matching pair are different, this nonconformity 160 is resolved by the processor with the edge of lower degree; it executes appropriate 161 steps of p-refinement in order to achieve the same degree as its neighbor on the interface edge. However, if red-green like refinement rules are applied as in [6], fixing 163 the degree for one interface edge might also change the degree of another interface 164 edge and cause further nonconformity. Thus, multiple communication steps might 165 be required to eliminate nonconformity in degree. This issue was the main motiva- 166 tion for us to find a more flexible p-refinement algorithm and more general nodal 167 basis functions for transition elements, allowing the mesh to be made both h and p 168 conforming with just one communication step. Such approach is described in [5, 7]. 169

When the global mesh is conforming, a second reordering as described above is 170 carried out locally on each processor, followed by a second all-to-all broadcast of the 171 new IPATH array. This time no nonconforming edges will be encountered during the 172 matching process.

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## 3.3 Coarse Mesh Regularization

The coarse part of the local mesh on processor I allows a complete conforming mesh 175 of the whole domain on each processor, thus avoiding otherwise necessary commu- 176 nication steps. Due to constraints of shape regularity, the coarse mesh will typically 177 be reasonably fine in areas near the fine subregion  $\Omega_I$  and become more coarse in 178 regions more distant from  $\Omega_I$ . However, in some special situations such as having 179 a singularity outside of  $\Omega_I$ , the coarse mesh on processor I might be refined [8]. In 180 very unusual circumstances, it is possible for the coarse mesh on some processors to 181 be more refined (in h or in p) than the global fine mesh in some areas. Although this 182 does not influence the global fine mesh solution directly, our DD solver assumes that 183 the coarse mesh on each processor is not more refined than the global fine mesh, see 184 [4, 9].

As described in Sect. 3.1, the IPATH array on each processor has a section for the coarse interface edges; this part of the data structure is local and different on every 187 processor. Following the second and final broadcast of the IPATH data structure, 188 each coarse interface edge is matched with one of the global fine edges. Here, the 189 matching is one-way from a coarse edge to a fine edge only. Based on this type of 190 matching, over-refined coarse edges are identified and then unrefined in either h or p. 191

We have also observed empirically [5, 9] that the convergence properties of our 192 DD solver are enhanced when elements in the coarse regions having edges on the 193 coarse interface system are more refined than those in the interior parts of the coarse region. To capture this effect, we also allow some limited refinement of elements 195 lying along the coarse interface. The level of refinement on the interface boundary of 196  $\Omega_I$  is determined by its distance from  $\Omega_I$ ; distance is measured in a graph in which the  $\Omega_I$  correspond to vertices and the edge between  $\Omega_I$  and  $\Omega_I$  is present if and only 198 if they have a shared interface boundary. The level of allowed refinement decays as 199  $2^{-K}$ , where K is the distance from  $\Omega_I$  to  $\Omega_J$ .

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# **Robust Parameter-Free Multilevel Methods** for Neumann Boundary Control Problems

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Summary. We consider a linear-quadratic elliptic control problem (LOECP). For the problem 8 we consider here, the control variable corresponds to the Neumann data on the boundary of 9 a convex polygonal domain. The optimal control unknown is the one for which the harmonic 10 extension approximates best a specified target in the interior of the domain. We propose a 11 multilevel preconditioner for the reduced Hessian resulting from the application of the Schur 12 complement method to the discrete LOECP. In order to derive robust stabilization parametersfree preconditioners, we first show that the Schur complement matrix is associated to a linear 14 combination of negative Sobolev norms and then propose preconditioner based on multilevel 15 methods. We also present numerical experiments which agree with the theoretical results.

1 Introduction 17

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The problem of solving linear systems is central in numerical analysis. Systems arising from the discretization of PDEs and control problems have received special attention since they appear in many applications, such as in fluid dynamics and structural 20 mechanics. Typically, as the dimension of the discrete space increases, the resulting 21 system becomes very ill-conditioned. To avoid the large cost of LU factorizations of 22 KKT saddle point linear systems, we consider instead the reduced Hessian systems. 23 To build efficient solvers, the spectral properties of these systems must be taken into 24 account. In this paper, we develop the mathematical tools necessary to analyze and to 25 design solvers for a model control problem. We believe that the proposed framework 26 can be extended to more complex control problems.

# 2 Setting Out the Problem

Consider the following LQECP:

$$\begin{split} \text{Minimize } J(u,\lambda) &:= \|u - u_*\|_{L^2(\Omega)}^2 + \frac{\alpha}{2} \, \|\lambda\|_{H^{-1/2}(\Gamma)}^2 + \frac{\beta}{2} \, \|\lambda\|_{L^2(\Gamma)}^2 \\ \text{subject to} & \begin{cases} -\Delta u(x) = f(x) & \text{in } \Omega \subset \mathbb{R}^2, \\ \gamma \frac{\partial u}{\partial \eta}(s) = -\lambda(s) & \text{on } \Gamma := \partial \Omega, \end{cases} \end{split} \tag{1}$$

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where  $u_*$  and f are given functions in  $L^2(\Omega) \setminus \mathbb{R}$ ,  $\gamma$  is the trace operator on  $\Gamma$ , and  $\alpha$ and  $\beta$  are nonnegative given stabilization parameters. The minimization is taken on 31  $u \in H^1(\Omega) \setminus \mathbb{R}$  and  $\lambda \in L^2(\Gamma) \setminus \mathbb{R}$ . Here, "\\R" stands for functions with zero average 32 on  $\Omega$  or  $\Gamma$ . We assume that the domain  $\Omega$  is a convex polygonal domain, hence, 33  $H^2$ -regularity of u is assumed. The norm  $H^{-1/2}(\Gamma)$  is defined as

$$\|\lambda\|_{H^{-1/2}(\Gamma)}^2 := |\nu_{\lambda}|_{H^1(\Omega)}^2,$$
 (2)

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where  $v_{\lambda} \in H^1(\Omega) \backslash \mathbb{R}$  is the harmonic extension of  $\lambda$  in  $\Omega$ . We remark that the 35 assumption  $\alpha + \beta > 0$  is necessary for the well-posedness of the problem (1), see 36 [7, 9, 11] and references therein. The case  $\alpha = \beta = 0$  can also be treated by enlarging the minimizing space for  $\lambda$  from  $H^{-1/2}(\Gamma)\backslash\mathbb{R}$  to  $H^{-3/2}_{t,00}(\Gamma)\backslash\mathbb{R}$ ; see [6] for 38 details. To make the notation less cumbersome, we sometimes drop " $\mathbb{R}$ " below.

We consider the following discretization for the LOECP (1). We consider the 41 space of piecewise linear and continuous functions  $V_h(\Omega) \subset H^1(\Omega)$  to approximate 42 u and p, and  $\Lambda_h(\Gamma) \subset H^{1/2}(\Gamma)$  (the restriction of  $V_h(\Omega)$  to  $\Gamma$ ) to approximate  $\lambda$ . The 43 underlying triangulation  $\mathcal{T}_h(\Omega)$  is assumed to be quasi-uniform with mesh size O(h). 44 Let  $\{\phi_1(x),\ldots,\phi_n(x)\}$  and  $\{\varphi_1(x),\ldots,\varphi_m(x)\}$  denote the standard hat nodal basis 45 functions for  $V_h(\Omega)$  and  $\Lambda_h(\Gamma)$ , respectively. The corresponding discrete problem 46 associated to (1) results in

$$\begin{bmatrix} M & 0 & A^T \\ 0 & G & Q^T E^T \\ A & E Q & 0 \end{bmatrix} \begin{bmatrix} \mathbf{u} \\ \lambda \\ \mathbf{p} \end{bmatrix} = \begin{bmatrix} \mathbf{f}_1 \\ \mathbf{f}_2 \\ \mathbf{f}_3 \end{bmatrix}, \tag{3}$$

where the matrices M and A are the mass and stiffness matrices on  $\Omega$ , and Q is the 48 mass matrix on  $\Gamma$ . We define  $Q_{ext_{ij}}=(\phi_i, \varphi_j)_{L^2(\Gamma)}; \phi_i \in V_h(\Omega)$  and  $\varphi_j \in \Lambda_h(\Gamma)$ . It is easy to see that  $Q_{ext} = EQ$ , where  $E \in \mathbb{R}^{n \times m}$  is the trivial zero discrete extension op-50 erator defined from  $\Lambda_h(\Gamma)$  to  $V_h(\Omega)$ . We define  $G \in \mathbb{R}^{m \times m}$  as be the matrix associated 51 to the norm  $\frac{\alpha}{2} \|\cdot\|_{H_h^{-1/2}(\Gamma)}^2 + \frac{\beta}{2} \|\cdot\|_{L^2(\Gamma)}^2$  on  $\Lambda_h(\Gamma)$ , where  $\|\lambda\|_{H_h^{-1/2}(\Gamma)} := |\nu_\lambda^h|_{H^1(\Omega)}$  52 with  $\nu_{\lambda}^h:=A^{\dagger}Q_{ext}\lambda$ , i.e.,  $\nu_{\lambda}^h$  is the discrete harmonic extension version of (2) with  $\lambda\in$  53  $\Lambda_h(\Gamma)$ . Hence, we have  $G=\alpha(Q_{ext}^TA^{\dagger})A(A^{\dagger}Q_{ext})+\beta Q=Q^T(\alpha E^TA^{\dagger}E+\beta Q^{-1})Q$ . 54 Here and the following  $A^{\dagger}$  is the pseudo inverse of A. The discrete forcing terms are 55 defined by  $(\mathbf{f}_1)_i = \int_{\Omega} u_*(x) \phi_i(x) dx$ , for  $1 \le i \le n$ ,  $\mathbf{f}_2 = 0$  and  $(\mathbf{f}_3)_i = \int_{\Omega} f(x) \phi_i(x) dx$ . 56

## 3 The Reduced Hessian $\mathcal{H}$

In this paper we propose and analyze preconditioners for the reduced Hessian 58 associated to (3). Eliminating the variables u and p from Eq.(3), and denoting 59  $S_1^{\dagger} := E^T A^{\dagger} E$  and  $S_2^{\dagger} := E^T A^{\dagger} M A^{\dagger} E$ , we obtain

$$\mathcal{H} \lambda := Q(\alpha S_1^{\dagger} + \beta Q^{-1} + S_3^{\dagger}) Q \lambda = b := Q_{ext}^T A^{\dagger} M A^{\dagger} f_3 - Q_{ext}^T A^{\dagger} f_1. \tag{4}$$

The matrix  $\mathcal{H}$  is known as the Schur complement (reduced Hessian) with respect 61 to the discrete control variable  $\lambda$ . We observe that the state variable u can be obtained 62

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by solving (4) and using the third equation of (3). We note that the Reduced matrix 63  $\mathcal{H}$  is a symmetric positive definite matrix on 64

$$\Lambda_h(\Gamma)\backslash_{Q}\mathbb{R}:=\{\lambda\in\Lambda_h(\Gamma); (\lambda,1)_{L^2(\Gamma)}=(Q\lambda,1_m)_{\ell^2}=0\},$$
 65

hence, we consider the Preconditioned Conjugate Gradient (PCG) with a preconditioner acting on  $\Lambda_h(\Gamma)\setminus_o\mathbb{R}$ . Note also that  $A^{\dagger}$  is also symmetric positive definite 67 matrix on

$$V_h(\Omega)\setminus_M \mathbb{R} := \{u \in V_h(\Omega); (u,1)_{L^2(\Omega)} = (Mu,1_n)_{\ell^2} = 0\}.$$
 69

The main goal of this paper is to develop robust preconditioned multilevel methods 70 for the matrix  $\mathcal{H}$  such that the condition number estimates that do not depend on  $\alpha$ and  $\beta$ , and depend on  $\log^2(h)$ .

We point out that several block preconditioners for solving systems like (3) were 74 proposed in the past; see [1, 8, 11, 14] and references therein. These preconditioners 75 depend heavily on the availability of a good preconditioner for the Schur complement 76 matrix. To the best of our knowledge, no robust and mathematically sounded pre- 77 conditioner was systematically carried out for the reduced Hessian (4). Most of the 78 existing work is toward problems where the control variable is f rather than  $\lambda$ , and 79 even for these cases, condition number estimates typically deteriorate when all the 80 stabilization parameters go to zero. Related work to ours is developed in [13] where 81 it is proposed a preconditioner for the first biharmonic problem discretized by the 82 mixed finite element method introduced by Ciarlet and Raviart [4]. Using techniques 83 developed in [5], Peisker transforms the discrete problem to an interface problem and 84 a preconditioner based on FFT is proposed and analyzed. This approach can also be 85 interpreted as a control problem like (1), however, replacing the Neumann control 86 by a Dirichlet control. We note that Dirichlet control problems are much easier to 87 handle and to study since in (4) the operator  $S_3^{\dagger}$  is replaced by  $S_1^{\dagger}$ , and therefore, a 88 multilevel method such as in [2], can be applied. An attempt to precondition the Neu-89 mann control problem via FFT was considered in [7], however, such as in Peisker's 90 work, it holds only for special meshes where the Schur complement matrix and the 91 mass matrix on  $\Gamma$  share the same set of eigenvectors.

### 4 Theoretical Remarks on the Reduced Hessian ${\mathcal H}$

In this section we associate the Reduced Hessian  ${\mathscr H}$  to a linear combination of 94 Sobolev norms. Here and below we use the notation  $a \prec (\succ) b$  to indicate that 95 a < (>)Cb, where the positive constant C depends only on the shape of  $\Omega$  and 96  $\mathcal{T}_h(\Omega)$ . When  $a \leq b \leq a$ , we say  $a \approx b$ .

98 First we observe that G is associated to the norm  $\frac{\alpha}{2} \|\cdot\|_{H_h^{-1/2}(\Gamma)}^2 + \frac{\beta}{2} \|\cdot\|_{L^2(\Gamma)}^2$  in 99  $\Lambda_h(\Gamma)$ . It is well known that for  $\lambda \in \Lambda_h(\Gamma) \setminus_{\Omega} \mathbb{R}$  we have 100

$$\lambda^{T} Q S_{1}^{\dagger} Q \lambda = \|\lambda\|_{H_{b}^{-1/2}(\Gamma)}^{2} \asymp \|\lambda\|_{H^{-1/2}(\Gamma)}^{2}. \tag{5}$$

What is not obvious is how to associate the matrix  $QS_3^{\dagger}Q$  to a Sobolev norm, and this is given in the following result (see [6]):

**Theorem 1.** Let  $\Omega \subset \mathbb{R}^2$  be a convex polygonal domain. Let  $v_{\lambda}^h := A^{\dagger}Q_{ext} \lambda \in \mathbb{R}^2$  be  $V_h(\Omega) \setminus_M \mathbb{R}$  be the discrete harmonic function with Neumann data  $\lambda \in \Lambda_h(\Gamma) \setminus_Q \mathbb{R}$ . 10. Then,  $\lambda^T Q S_3^{\dagger} Q \lambda = \|v_{\lambda}^h\|_{L^2(\Omega)}^2 \asymp \|\lambda\|_{H^{-3/2}(\Gamma)}^2 + h^2 \|\lambda\|_{H^{-1/2}(\Gamma)}^2. \tag{6}$ 

Using these results we conclude that  ${\mathscr H}$  is associated to the following linear 106 combination of Sobolev norms

$$\lambda^{T} \mathcal{H} \lambda \simeq (\alpha + h^{2}) \|\lambda\|_{H^{-1/2}(\Gamma)}^{2} + \beta \|\lambda\|_{L^{2}(\Gamma)}^{2} + \|\lambda\|_{H_{1,00}^{-3/2}(\Gamma)}^{2}. \tag{7}$$

*Remark 1.* We next hint why the norm  $\|\cdot\|_{H_{t,00}^{-3/2}(\Gamma)}^2$  is fundamental for this problem. 109

Let  $\{\Gamma_k\}_{1\leq k\leq K}$  and  $\{\delta_k\}_{1\leq k\leq K}$  be the edges and the vertices of the polygonal  $\Gamma$ , 110 respectively. Let  $C^{\infty}_{t,00}(\Gamma_k):=\{\lambda\in C^{\infty}(\Gamma_k);\partial\lambda/\partial\tau_k\in C^{\infty}_0(\Gamma_k)\}$ , where  $\tau_k$  stands for 111 the tangential unit vector on  $\Gamma_k$ . Define  $H^2_{t,00}(\Gamma_k)$  by the closure of  $C^{\infty}_{t,00}(\Gamma_k)$  in the 112  $H^2(\Gamma_k)$ -norm, that is,

$$H_{t,00}^{2}(\Gamma_{k}) := \{ \lambda \in H^{2}(\Gamma_{k}); \frac{\partial \lambda}{\partial \tau_{k}}(\delta_{k-1}) = \frac{\partial \lambda}{\partial \tau_{k}}(\delta_{k}) = 0 \}. \tag{8}$$

Using interpolation theory of operators and a characterization of  $H_{t,00}^{3/2}(\Gamma_k)$ , see [10], 114 it is possible to show that

$$H_{t,00}^{3/2}(\Gamma_k) := \left[ H_{t,00}^2(\Gamma_k), H^1(\Gamma_k) \right]_{1/2} = \left\{ \lambda \in H^{3/2}(\Gamma_k); \partial \lambda / \partial \tau_k \in H_{00}^{1/2}(\Gamma_k) \right\}.$$

We define  $H_{t,00}^{3/2}(\Gamma) = H^{1/2}(\Gamma) \cap \prod_{k=1}^K H_{t,00}^{3/2}(\Gamma_k)$  endowed with the norm

$$\|\lambda\|_{H^{3/2}_{t,00}(\Gamma)} := \|\lambda\|_{H^{1/2}(\Gamma)}^2 + \sum_{k=1}^K \|\frac{\partial \lambda}{\partial \tau_k}\|_{H^{1/2}_{00}(\Gamma_k)}^2, \tag{9}$$

and define  $H_{t,00}^{-3/2}(\Gamma)=(H_{t,00}^{3/2}(\Gamma))'$ . The fundamental property of this space is that

$$\|\lambda\|_{H_{r,00}^{-3/2}(\Gamma)} \asymp \|\nu_{\lambda}\|_{L^{2}(\Omega)},$$
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where  $v_{\lambda}$  is defined by (2); see [6].

# 5 Preconditioning Sobolev Norms Using Multilevel Methods

In this section, using multilevel based preconditioners, we develop spectral approximations for matrices associated to several Sobolev norms; see [2, 3, 12, 15], and 122 references therein.

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#### 5.1 Notation and Technical Tools

From now on, we assume that the triangulation  $\mathcal{T}_h$  of  $\Gamma$  has a multilevel structure. 125 More precisely, denoting  $\mathcal{T}_h$  as the restriction of  $\mathcal{T}_h(\Omega)$  to  $\Gamma$ , we assume that the triangulation  $\mathcal{T}_h$  is obtained from (L-1) successive refinements of an initial coarse triangulation  $\tau_0$  with initial grid size  $h_0$ . We assume also that  $h_\ell = h_{\ell-1}/2$  is the grid 128 size on the  $\ell$ -th triangulation  $\mathcal{T}_{\ell}$  and associate the standard  $P_1$  finite element space 129  $V_\ell(\Gamma)$  generated by continuous and piecewise linear basis functions  $\{\phi_i^\ell\}_{i=1}^{m_\ell}$ . Hence, 130 we have

$$V_0(\Gamma) \subset V_1(\Gamma) \subset \cdots \subset V_L(\Gamma) := V_h(\Gamma) \subset L^2(\Gamma).$$
 132

Let  $P_{\ell}$  denote the  $L^2(\Gamma)$ -orthogonal projection onto  $V_{\ell}(\Gamma)$ , and let  $\Delta P_{\ell} := (P_{\ell} - P_{\ell})$  $P_{\ell-1}$ ), that is, the  $L^2(\Gamma)$ -orthogonal projection onto  $V_{\ell}(\Gamma) \cap V_{\ell-1}(\Gamma)^{\perp}$ . We have that  $P_0, (P_1 - P_0), \dots, (P_L - P_{L-1})$  restricted to  $V_L(\Gamma)$  are mutually  $L^2$ -orthogonal projections which satisfy: 136

$$I = P_0 + (P_1 - P_0) + \dots + (P_L - P_{L-1}). \tag{10}$$

Note that  $P_L = I$ . The matrix form of  $P_\ell$  restricted to  $V_L(\Gamma)$  is given by

$$P_{\ell} = R_{\ell}^T Q_{\ell}^{-1} R_{\ell} Q, \tag{11}$$

where  $R_{\ell}$  is the  $m_{\ell} \times m_L$  restriction matrix, that is, the i-th row of  $R_{\ell}$  is obtained by interpolating the basis function  $\varphi_i^\ell \in V_\ell := V_\ell(\Gamma)$  at the nodes of the finest triangulation  $au_L := au_h$ .

It follows from [2, 12], that for -3/2 < s < 3/2

$$\|\mathbf{v}\|_{H^{s}(\Gamma)}^{2} \approx \sum_{\ell=0}^{L} h_{\ell}^{-2s} \|(P_{\ell} - P_{\ell-1})\mathbf{v}\|_{L^{2}(\Gamma)}^{2}, \text{ for all } \mathbf{v} \in V_{L}.$$
 (12)

This constraint for s comes from the fact that for  $s \ge 3/2$  we have  $V_h(\Gamma) \not\subset H^s(\Gamma)$ , 143 therefore, the equivalence deteriorates when s tends to 3/2. Results for negative 144 norms are obtained by duality. 145

146 We now describe how to represent the splitting  $\sum_{\ell=0}^L \mu_\ell \|(P_\ell - P_{\ell-1})\mathbf{v}\|_{L^2(\Gamma)}^2$  into a 147 matrix form. Let  $\Delta_{\ell} := (P_{\ell} - P_{\ell-1})Q^{-1} = R_{\ell}^T Q_{\ell}^{-1} R_{\ell} - R_{\ell-1}^T Q_{\ell-1}^{-1} R_{\ell-1}$ . Then we have

$$\Delta_k Q \Delta_\ell = \delta_{k\ell} \Delta_\ell \text{ and } \sum_{\ell=0}^L \mu_\ell \| (P_\ell - P_{\ell-1}) \mathbf{v} \|_{L^2(\Gamma)}^2 = \sum_{\ell=0}^L \mu_\ell \mathbf{v}^T Q (P_\ell - P_{\ell-1}) \mathbf{v}, \quad (13)$$

where  $P_{-1}=0$ . We observe that  $Q(P_{\ell}-P_{\ell-1})=Q\Delta_{\ell}Q$  is symmetric semi-positive 149 definite. By (12) and (13), for all  $v \in V_L$  we have 150

$$\|\mathbf{v}\|_{H^{-1/2}(\Gamma)}^2 \asymp \left(\sum_{\ell=0}^L h_\ell \Delta_\ell Q \mathbf{v}, Q \mathbf{v}\right). \tag{14}$$

To invert a matrix of the form  $\sum_{k=0}^{L} \mu_k^{-1} \Delta_k Q$ , we first assume that  $\mu_k > 0$ ,  $0 \le 1$  $k \le L$ . Then, from (10) and (13) we obtain 152

$$(\sum_{k=0}^{L} \mu_k^{-1} \Delta_k Q) (\sum_{\ell=0}^{L} \mu_\ell \Delta_\ell Q) = I.$$
 (15)

#### 5.2 Multilevel Preconditioner for the Reduced Hessian $\mathcal{H}$

In this subsection we analyze a multilevel preconditioner for Reduced Hessian  $\mathcal{H}$ . 154 We first present a preconditioner for G as follows. Using (2), (14) and (15) we obtain 155

$$\begin{cases}
S_1 & \approx Q \sum_{\ell=0}^L h_\ell^{-1} \Delta_\ell Q, \\
Q S_1^{\dagger} Q & \approx Q \sum_{\ell=0}^L h_\ell \Delta_\ell Q.
\end{cases}$$
(16)

The above equivalences yield simultaneous approximation for the spectral representations of  $G := \beta Q + \alpha Q S_1^{\dagger} Q$  in terms of the  $\Delta_{\ell}$  and Q. More precisely, 158

$$G \simeq Q \sum_{\ell=1}^{L} (\beta + \alpha h_{\ell}) \Delta_{\ell} Q, \tag{17}$$

and using (15) and (17), the following spectral equivalency holds

$$G^{-1} \simeq \sum_{\ell=0}^{L} (\beta + \alpha h_{\ell})^{-1} \Delta_{\ell}. \tag{18}$$

We next establish that  $\sum_{\ell=0}^L (h_\ell^{-3}) \Delta_\ell$  is a quasi-optimal preconditioner for  $QS_3^\dagger Q$ . 160 More precisely, we have the following result (see [6]): 161

**Theorem 2.** For all  $v_L \in V_L$ , the following inequalities hold:

$$\|\mathbf{v}_{L}\|_{H_{t,00}^{-3/2}(\Gamma)}^{2} \leq \sum_{\ell=1}^{L} h_{\ell}^{3} \|\Delta P_{\ell} \mathbf{v}_{L}\|_{L^{2}}^{2} \leq (L+1)^{2} \|\mathbf{v}_{L}\|_{H_{t,00}^{-3/2}(\Gamma)}^{2}.$$
(19)

From Theorems 1 and 2 and (15), we establish the main result, the quasioptimality for a preconditioner for  $\mathcal{H}$ . 164

**Theorem 3.** Let 
$$\mathscr{PC}:=\sum_{\ell=0}^L (\alpha h_\ell + \beta + h_\ell^3)^{-1} \Delta_\ell$$
. Then

$$(L+1)^{-2}\mathscr{P}\mathscr{C} \ \preceq \ \mathscr{H}^{-1} \ \preceq \ \mathscr{P}\mathscr{C}. \tag{20}$$

# 6 Numerical Results

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In this section we show numerical results conforming the theory developed. For all 167 tests presented,  $\Omega$  is the square domain  $[0,1] \times [0,1]$ . The triangulation of  $\Omega$  is constructed as follows. We divide each edge of  $\partial \Omega$  into  $2^N$  parts of equal length, where N is an integer denoting the number of refinements. In all tests (cond) means condition number, (it) indicates the number of iterations of the PCG, (eig min) means 171 the lowest eigenvalue for preconditioned system. To calculate the eigenvalues we 172 build the preconditioned system and use the function eig of MATLAB. We can see 173 from tables below the asymptotic  $\log^2(h)$  behavior for the case  $\alpha = \beta = 0$ , i.e., 174 cond(N+1) - cond(N) grows linearly with N. As expected, larger is  $\alpha$  or  $\beta$ , better 175 conditioned are the preconditioned systems (Tables 1–4).

Remark 2. Numerical experiments show (not reported here) that the largest eigenvalue of  $(\sum_{\ell=0}^{L} \Delta_{\ell}) * Q$  divided by the largest eigenvalue of  $(\sum_{\ell=0}^{L} h_{\ell}^{-3} \Delta_{\ell}) * QS_{3}^{\dagger}Q$  178 converges to 36 when h decreases to zero. In tables above, we considered the rescaled 179 preconditioner 180

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	$\mathcal{PC}_r * \mathcal{H}$	with $\beta =$	1	$\mathcal{PC}_r * \mathcal{H}$	with $\beta = (0.1)$	3
$\overline{N\downarrow}$	cond	eig min	it	cond	eig min	it
4	1.04237	0.02756	2	4.94294	0.01622	7
5	1.04222	0.02757	2	4.87258	0.01655	7
6	1.04218	0.02757	2	4.85515	0.01663	7
7	1.04217	0.02757	2	4.85084	0.01665	7

**Table 1.** Equivalence between  $\mathscr{H}$  and  $\mathscr{PC}_r$  with r=36 and  $\alpha=0$ .

	$\mathscr{PC}_r * \mathscr{H}$	with $\beta = (0.1)$	6	$\mathscr{PC}_r * \mathfrak{I}$	$\mathscr{H}$ with $\beta = 0$	)
N	↓ cond	eig min	it	cond	eig min	it
4	28.1662	0.004747	15	33.5522	0.004016	16
5	24.3303	0.005739	20	41.9737	0.003407	25
6	20.3042	0.006984	22	50.5193	0.002930	35
7	18.9576	0.007514	20	59.2085	0.002550	44

**Table 2.** Equivalence between  $\mathcal{H}$  and  $\mathcal{PC}_r$  with r=36 and  $\alpha=0$ .

	$\mathscr{PC}_r * \mathscr{H}$	with $\alpha = 1$		$\mathscr{PC}_r * \mathscr{H}$	with $\alpha = (0.1)$	3
$\overline{\mathrm{N}}\downarrow$	cond	eig min	it	cond	eig min	it
4	4.62312	0.11893	10	13.7601	0.010698	14
5	5.12018	0.11826	10	18.3917	0.012503	19
6	5.33402	0.11798	11	26.2878	0.013139	22
7	5.45327	0.11788	12	35.6393	0.013312	26

**Table 3.** Equivalence between  $\mathcal{H}$  and  $\mathcal{PC}_r$  with r = 36 and  $\beta = 0$ .

$\mathscr{PC}_r * \mathscr{H}$	with $\alpha = (0.1)^6$	Ó	$\mathscr{PC}_r * \mathcal{F}$	$\mathscr{C}$ with $\alpha = 0$	)
4 33.4363	0.004031	16	33.5522	0.0040164	16
5 41.4318	0.003452	25	41.9737	0.0034074	25
6 48.1852	0.003073	33	50.5193	0.0029301	35
7 50.8326	0.002973	43	59.2085	0.0025501	44

**Table 4.** Equivalence between  $\mathcal{H}$  and  $\mathcal{PC}_r$  with r = 36 and  $\beta = 0$ .

$$\mathscr{P}\mathscr{C}_r:=\sum_{\ell=0}^L(lpha h_\ell+roldsymbol{eta}+h_\ell^3)^{-1}\Delta_\ell,$$
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with r=36, instead of  $\mathscr{PC}:=\sum_{\ell=0}^L(\alpha h_\ell+\beta+h_\ell^3)^{-1}\Delta_\ell$ . This change improves 182 considerably the condition number of preconditioners and improve slightly the number of iterations.

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# An Overlapping Domain Decomposition Method for a 3D PEMFC Model

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**Summary.** In this paper, an overlapping domain decomposition method is developed to sim- 11 ulate the water management of the polymer exchange membrane fuel cell on the local struc- 12 tured grids. Numerical experiments demonstrate that our methods are effective to deal with 13 the simulation on the non-matching grids with low mass balance error. 14

1 Introduction 15

Polymer exchange membrane fuel cells (PEMFCs) have been used in a large number 16 of industries worldwide because of their advantages such as low environmental impact, rapid start-up and high power density [15, 16]. The performance of fuel cell is 18 affected by many factors, such as material parameters, operating conditions, different 19 channel structures and so on [2, 9, 10].

For better performance, different structures for the anode and cathode gas channels are used in the PEMFC practical design. This asymmetrical structure can keep 22
the balance of pressures on both sides of the membrane. Thus the water management in cathode can be improved and the duration of fuel cell can be prolonged. An 24
unstructured grid partitioned by tetrahedra or triangles can be used for this asymmetrical fuel cell in single domain approach, but structured grids, such as hexahedron 26
and quadrilateral, are easily implemented and have super convergence [1, 4, 14]. 27
However, non-matching grids would be generated when partitioning with structured 28
grids in numerical simulations. Besides, since oxygen reduction reaction occurs in 29
cathode, the variation of physical quantities such as water concentration are more 30
significant in cathode than in anode. So it is necessary for cathode to simulate these 31
phenomena accurately by a refined grid. The objective of this paper is to provide an 32
overlapping domain decomposition method for the simulation of a 3D single-phase 33
PEMFC model with local structured grid in anode and cathode respectively. 34

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#### 1.1 Governing Equations

Based on [5, 16], a fundamental fuel cell model consists of five principles of conservation: mass, momentum, species, charge, and thermal energy. Typically the fuel 37 cell is divided into seven subregions: the anode gas channel, anode gas diffusion 38 layer (GDL), anode catalyst layer (CL), membrane, cathode gas channel, cathode 39 GDL, and cathode CL. In the following we specifically focus our interests on mass. 40 momentum conservation and water concentration arising in all seven subregions.

**Flow equations.** For flow field with velocity **u** and pressure P as unknowns, we 42 have the following modified Navier-Stokes equations

$$\nabla \cdot (\rho \mathbf{u}) = 0, \tag{1}$$

$$\frac{1}{\varepsilon^2} \nabla \cdot (\rho \mathbf{u} \mathbf{u}) = -\nabla P + \nabla \cdot (\mu \nabla \mathbf{u}) + S_u, \tag{2}$$

where  $\varepsilon$  is porosity,  $\rho$  is density, and  $\mu$  is effective viscosity. In (2) we indicate that 44 the additional source term  $S_u$  in GDL and CL is named as Darcy's drag and defined 45 by  $S_{\mu} = -\frac{\mu}{K} \mathbf{u}$ , where K is hydraulic permeability.

Species concentration equation. Water management is critical to achieve high 47 performance for PEMFC. Therefore, without loss of generality, in order to focus on 48 water management topics, we typically consider water as the only component in the 49 following simplified species concentration equation. Water concentration equation in 50 single gaseous phase is defined as follows with respect to concentration C

$$\nabla \cdot (\mathbf{u}C) = \nabla \cdot (D_g^{eff} \nabla C) + S_{H_2O}, \tag{3}$$

equation where  $D_g^{eff} = \varepsilon^{1.5} D_{gas}$  is the effective water vapor diffusivity. The source 52 term  $S_{H_2O}$  is given as follows. 53

$$S_{H_2O} = \begin{cases} -\nabla \cdot \left(\frac{n_d}{F} \mathbf{i}_e\right) - \frac{j}{2F} & \text{in cathode CL} \\ -\nabla \cdot \left(\frac{n_d}{F} \mathbf{i}_e\right) & \text{in anode CL} \\ 0 & \text{otherwise,} \end{cases}$$
(4)

where  $n_d$ , the electro-osmotic drag coefficient, is a constant value in our simulation. 54  $\nabla \cdot \mathbf{i}_e = -j$  which is derived from the continuity equation of proton potential.  $\mathbf{i}_e$  is 55 the current density vector and j is the volumetric transfer current of the reaction (or 56 transfer current density) defined by  $j = j_1 - (j_1 - j_2)z/l_{cell}$ . This is an approximation 57 of transfer current density for our simplified single-phase PEMFC model due to the 58 absence of proton and electron potentials [12]. 59

#### 1.2 Computational Domain and Boundary Conditions

The computational domain and its geometric sizes are schematically shown in Fig. 1 61 and Table 1.

For flow field (1), (2) and water concentration equation (3), the following boundary conditions are imposed: 64

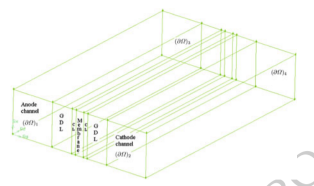


Fig. 1. Geometry of a single straight-channel PEMFC

Table 1. Physical coefficients and parameters

Parameter	Symbol	Parameter	Symbol	
Anode/cathode channel width $\delta_{CH}$	6.180mm	Anode/cathode GDL width $\delta_{GDL}$	0.235mm	
Anode/cathode CL width $\delta_{CL}$	0.010mm	Membrane width $\delta_{mem}$	0.018mm	
Cell length $l_{cell}$	70mm	Cell depth $h_{cell}$	6.360mm	
Porosity of membrane $\varepsilon$	0.26	Effective viscosity $\mu$	$3.166 \times 10^{-5} kg/(m \cdot s)$	
Porosity of GDL and CL $\varepsilon$	0.6	Water vapor diffusivity $D_{gas}$	$2.6 \times 10^{-5} m^2/s$	
Vapor density $\rho$	$0.882 \ kg/m^3$	Permeability of GDL and CL K	$2 \times 10^{-12} m^2$	
Electro-osmotic drag coefficient $n_d$	1.5	Transfer current density $j_1/j_2$	$20000/10000A/m^2$	

$$u_1 = u_2 = 0, u_3 = u_3|_{inlet}, C = C_{in} \text{ on inlet } (\partial \Omega)_1, (\partial \Omega)_2, \tag{5}$$

$$(PI - \mu \nabla \mathbf{u}) \cdot \mathbf{n} = 0$$
 on outlet  $(\partial \Omega)_3, (\partial \Omega)_4,$  (6)

$$u_1 = u_2 = u_3 = 0, \frac{\partial C}{\partial n} = 0$$
 on other boundaries. (7)

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# 2 Numerical Algorithm

#### 2.1 Domain Decomposition Method and Weak Forms

First, we split the domain  $(\Omega)$ , shown in Fig. 1, to two overlapping subdomains: 67 one is the anode and membrane  $(\Omega_a)$ , the other is the cathode and membrane  $(\Omega_c)$ . 68 The interface between anode CL and membrane is denoted as  $\mathscr{S}_a$ , and the interface between cathode CL and membrane is denoted as  $\mathscr{S}_c$ . The classical overlapping 70 Schwarz alternating method [13] is used in these two subdomains. Thus we are able 71 to reformulate Eqs. (1)–(3) to two Dirichlet-type interfacial boundary value subproblems.

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$$(\text{Problem A}) \left\{ \begin{array}{ll} \nabla \cdot (\rho \mathbf{u}_a) = 0 & \text{in } \Omega_a \\ \frac{1}{\varepsilon^2} \nabla \cdot (\rho \mathbf{u}_a \mathbf{u_a}) = -\nabla P_a + \nabla \cdot (\mu \nabla \mathbf{u}_a) - \frac{\mu}{K} \mathbf{u}_a & \text{in } \Omega_a \\ \nabla \cdot (\mathbf{u}_a C_a) = \nabla \cdot (D_g^{eff} \nabla C_a) + S_{H_2O} & \text{in } \Omega_a \\ u_{1,a} = u_{2,a} = 0, u_{3,a} = u_3|_{inlet}, C_a = C_{a,in} & \text{on } (\partial \Omega)_1 \\ (P_a I - \mu \nabla \mathbf{u_a}) \cdot \mathbf{n} = 0 & \text{on } (\partial \Omega)_3 \\ C_a = C_c & \text{on } \mathscr{S}_c \\ u_{1,a} = u_{2,a} = u_{3,a} = 0, \frac{\partial C}{\partial n} = 0 & \text{on other boundaries.} \end{array} \right.$$

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$$(\text{Problem C}) \left\{ \begin{array}{ll} \nabla \cdot \left( \rho \, \mathbf{u}_c \right) = 0 & \text{in } \Omega_c \\ \frac{1}{\varepsilon^2} \nabla \cdot \left( \rho \, \mathbf{u}_c \, \mathbf{u}_c \right) = - \nabla P_c + \nabla \cdot \left( \mu \nabla \mathbf{u}_c \right) - \frac{\mu}{K} \, \mathbf{u}_c & \text{in } \Omega_c \\ \nabla \cdot \left( \mathbf{u}_c C_c \right) = \nabla \cdot \left( D_g^{eff} \nabla C_c \right) + S_{H_2O} & \text{in } \Omega_c \\ u_{1,c} = u_{2,c} = 0, u_{3,c} = u_3|_{inlet}, C_c = C_{c,in} & \text{on } (\partial \Omega)_2 \\ \left( P_c I - \mu \nabla \mathbf{u}_c \right) \cdot \mathbf{n} = 0 & \text{on } (\partial \Omega)_4 \\ C_c = C_a & \text{on } \mathscr{S}_a \\ u_{1,c} = u_{2,c} = u_{3,c} = 0, \frac{\partial C}{\partial n} = 0 & \text{on other boundaries.} \end{array} \right.$$

Considering various nonlinearities of equations, we particularly employ Picard's 75 scheme to linearize the nonlinear source term. Define 76

$$\begin{split} &V_a := \{\mathbf{v}_a = (v_{1,a}, v_{2,a}, v_{3,a})^\top \in [H^1]^3 \mid v_{1,a}|_{(\partial\Omega)_1} = v_{2,a}|_{(\partial\Omega)_1} = 0, v_{3,a}|_{(\partial\Omega)_1} = u_{3,a}|_{inlet}\}, \\ &\widetilde{V}_a := \{\mathbf{v}_a = (v_{1,a}, v_{2,a}, v_{3,a})^\top \in [H^1]^3 \mid v_{1,a}|_{(\partial\Omega)_1} = v_{2,a}|_{(\partial\Omega)_1} = v_{3,a}|_{(\partial\Omega)_1} = 0\}, \\ &Q_a := \{w \in H^1 \mid w|_{(\partial\Omega)_1} = C_{in,a} \text{ and } w|_{\mathscr{S}_c} = C_c\}, \quad \widetilde{Q}_a := \{w \in H^1 \mid w|_{(\partial\Omega)_1} = 0 \text{ and } w|_{\mathscr{S}_c} = 0\}, \\ &P_a := L^2(\Omega_a). \end{split}$$

Then for any  $(\mathbf{v}_a,q_a,w_a)\in \widetilde{V}_a\times P_a\times \widetilde{Q}_a$ , find  $(\mathbf{u}_a^{k+1},P_a^{k+1},C_a^{k+1})\in V_a\times P_a\times Q_a$ , 77 such that

$$\begin{cases}
(\mu \nabla \mathbf{u}_{a}^{k+1}, \nabla \mathbf{v}_{a})_{\Omega_{a}} + (\frac{\rho}{\varepsilon^{2}} \nabla \mathbf{u}_{a}^{k} \mathbf{u}_{a}^{k+1}, \mathbf{v}_{a})_{\Omega_{a}} - (P_{a}^{k+1}, \nabla \mathbf{v}_{a})_{\Omega_{a}} + (\frac{\mu}{K} \mathbf{u}_{a}^{k+1}, \mathbf{v}_{a})_{\Omega_{a}} = 0 \\
(\nabla \mathbf{u}_{g}^{k+1}, q_{a})_{\Omega_{a}} = 0 \\
(D_{g}^{eff} \nabla C_{a}^{k+1}, \nabla w_{a})_{\Omega_{a}} + (\nabla \cdot (\mathbf{u}_{a}^{k} C_{a}), w_{a})_{\Omega_{a}} = (S_{H_{2}O}, w_{a})_{\Omega_{a}},
\end{cases}$$
(8)

which  $(\cdot, \cdot)_{\Omega_i}$  stands for the  $L^2$  inner product in  $\Omega_i$ . And in subdomain  $\Omega_c$ , we have 79 the same weak form with (8).

### 2.2 An Overlapping Domain Decomposition Algorithm

Firstly, the subdomains  $\Omega_a$  and  $\Omega_c$  are partitioned into cuboids independently, which 82 implies that the grids are local structured in anode and cathode. Define a partition 83  $\mathscr{T}_{h_i}$  in  $\Omega_i$  (i,j) represent a or c), and  $\Sigma_{i,j}$  is the set of mesh points of  $\mathscr{T}_{h_i}$  on  $\mathscr{S}_j$ . 84

To discretize weak form (8), we introduce the finite element space  $V_{h_i} \times P_{h_i} \subseteq 85$   $V_i \times P_i$  on  $\mathcal{T}_{h_i}$ , where  $V_{h_i} \times P_{h_i}$  denotes the Q2Q1 (triquadratic velocity and trilinear 86 pressure) finite element spaces.  $Q_{h_a}$  denotes the triquadratic finite element space for 87 water concentration whose members equal  $f_a$  on  $\mathcal{S}_c$ , where  $f_a$  represents the values 88 of points in the sets of  $\Sigma_{a,c}$ , which are obtained from the previous alternating step  $C^k$  89

by lagrange interpolation. Moreover, let  $\widetilde{Q}_{h_a}\subseteq\widetilde{Q}_a$  be the triquadratic finite element 90 space and  $\widetilde{V}_{h_a}\subseteq \widetilde{V}_a$  be the triquadratic finite element space. In subdomain  $\Omega_c,\,Q_{h_c}$ and  $V_{h_a}$  are defined in the same ways.

For flow and water concentration equations, we introduce the following combined finite element-upwind finite volume schemes [11].

For any given  $(\mathbf{u}_{h_i}^k, P_{h_i}^k, C_{h_i}^k) \in V_{h_i} \times P_{h_i} \times Q_{h_i}$  (k = 0, 1, 2, ...), find  $(\mathbf{u}_{h_i}^{k+1}, P_{h_i}^{k+1}, ...)$  $C_{h_i}^{k+1}$ )  $\in V_{h_i} \times P_{h_i} \times Q_{h_i}$  (k = 0, 1, 2, ...), such that

$$(\mu \nabla \mathbf{u}_{h_{i}}^{k+1}, \nabla \mathbf{v}_{h_{i}})_{\Omega_{i}} + (\frac{\rho}{\varepsilon^{2}} \nabla \mathbf{u}_{h_{i}}^{k} \mathbf{u}_{h_{i}}^{k+1}, \mathbf{v}_{h_{i}})_{\Omega_{i}} - (P_{h_{i}}^{k+1}, \nabla \mathbf{v}_{h_{i}})_{\Omega_{i}} + (\frac{\mu}{K} \mathbf{u}_{h_{i}}^{k+1}, \mathbf{v}_{h_{i}})_{\Omega_{i}} = 0$$

$$(\nabla \mathbf{u}_{h_{i}}^{k+1}, q_{h_{i}})_{\Omega_{i}} = 0 \quad \forall (\mathbf{v}_{h_{i}}, q_{h_{i}}) \in \widetilde{V}_{h_{i}} \times P_{h_{i}}, \quad (9)$$

$$(D_{g}^{eff} \nabla C_{h_{i}}^{k+1}, \nabla w_{h_{i}})_{\Omega_{i}} + (\nabla \cdot (\mathbf{u}_{h_{i}}^{k+1} C_{h_{i}}^{k+1}), w_{h_{i}})_{\Omega_{i}} + \delta(h_{i}) \mathbf{u}_{h_{i}}^{k+1} \cdot (\nabla C_{h_{i}}^{k+1}, \nabla w_{h_{i}})_{\Omega_{i}}$$

$$= (S_{H_{2}O}, w_{h_{i}})_{\Omega_{i}} \quad \forall w_{h_{i}} \in \widetilde{Q}_{h_{i}}, \quad (10)$$

where the last term in the left hand side of (10) is a stabilizing term, derived from 97 streamline-diffusion scheme [3, 6–8]. Basically we hold  $\delta(h) = Ch$ , C is a certain 98 constant parameter, which is chosen artificially with least possible on the premise of 99 optimal stability. Usually starting with small ones, we gradually increase the value of 100 C and compute the corresponding finite element equation (10) until gained numerical 101 solutions are not oscillating any more in convection-dominated gas channel.

Now, we are in position to describe the overlapping domain decomposition algo- 103 rithm with the finite element discretizations.

**Algorithm**: Given  $\mathbf{u}_h^0, C_h^0$ , the following procedures are successively executed 105 106

Step 1. Solve (9) in 
$$\Omega_a$$
 and  $\Omega_c$  for  $(\mathbf{u}_{h_i}^{k+1}, P_{h_i}^{k+1})$ , respectively, until

$$\|\mathbf{u}_{h_i}^{k+1} - \mathbf{u}_{h_i}^k\|_{L^2(\Omega_i)} + \|P_{h_i}^{k+1} - P_{h_i}^k\|_{L^2(\Omega_i)} < \text{tolerance}.$$
 (11)

Step 2. Solve (10) for  $C_{h_a}^{k+1}$ , and construct the finite element space  $\widetilde{Q}_{h_c}$  for  $\Omega_c$ . Step 3. Solve (10) for  $C_{h_c}^{k+1}$ , and construct the finite element space  $\widetilde{Q}_{h_a}$  for  $\Omega_a$ . Step 4. Compute the following stopping criteria: 108 109 110

$$\|C_{h_a}^{k+1} - C_{h_a}^k\|_{L^2(\Omega_a)} < \text{tolerance.}$$
 (12)

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If yes, then numerical computation is complete. Otherwise, go back to the step 2 111 and continue. 112

#### 3 Numerical Results

In this section, we will carry out the following numerical experiments which indicate that our methods are effective to deal with the non-matching grids, see Fig. 2 115 for example, in the simulation of the PEMFC. The velocity  $u_3|_{inlet}$  is defined as a 116 paraboloidal-like function given in (13). 117

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$$u_{3}|_{inlet} = \begin{cases} 0.2\sin\frac{x\pi}{\delta_{CH}}\sin\frac{y\pi}{\delta_{CH}} & \text{on anode inlet } (\partial\Omega)_{1} \\ 0.3\sin\frac{x\pi}{\delta_{CH}}\sin\frac{(y-l_{add})\pi}{\delta_{CH}} & \text{on cathode inlet } (\partial\Omega)_{2} \end{cases},$$
(13)

where  $l_{add} = \delta_{CH} + \delta_{GDL} + \delta_{CL} + \delta_{mem}$ .

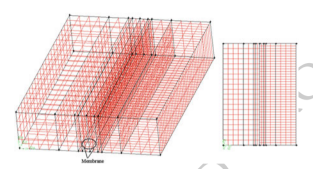


Fig. 2. An example of non-matching grids

Figures 3 and 4 show the velocity field in anode and cathode of fuel cell at the face of x=3.18 mm with this two method. As expected, there is a large difference in the velocity scale between the porous media and the open channel. The velocity in porous GDL is at least two orders of magnitude smaller than that in the open gas channel, indicating that gas diffusion is the dominant transport mechanism in porous GDL. Porous CL has a smaller velocity than GDL due to the inferior diffusion ability.

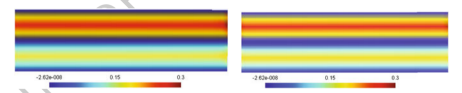


Fig. 3. Velocity with DDM

Fig. 4. Velocity with single domain

Figure 5 displays the water concentration distribution, presenting in the phase of water vapor, in anode and cathode. As shown in the figure, significant variations are displayed in both anode and cathode; in the porous media there is an increased water vapor concentration along the channel.

In order to verify the correctness of our numerical solutions, we compute the 129 relative error of mass balance in terms of the numerical fluxes at the inlet and outlet. 130

$$\text{mass balance error} = \frac{\left| \int_{(\partial \Omega)_{outlet}} C u_3 dS - \int_{(\partial \Omega)_{inlet}} C_{in} u_3 \right|_{inlet} dS - \int_{\Omega} S_{H_2O} dV |}{\int_{(\partial \Omega)_{inlet}} C_{in} u_3 |_{inlet} dS}. \tag{14}$$

#### An Overlapping Domain Decomposition Method for a 3D PEMFC Model

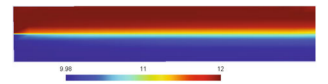


Fig. 5. Distributions of water concentration with DDM

The tolerance of our stopping criteria (12) for Schwarz alternating iteration is 131  $10^{-20}$ . By plugging the assigned and the computed concentration C as well as horizontal velocity  $u_3$  in Eq. (14), we attain a convergent mass balance error for our 133 numerical solutions along with the continuously refining grids, shown in Table 2. A 134 more accurate mass balance error is attained for the numerical solutions with DDM. 135

Table 2. Convergent mass balance error for with different grids

	Grids	Unknowns		Error with single domain
Mesh1	720	36260	$9.731 \times 10^{-3}$	$8.112 \times 10^{-3}$
Mesh2	1440	58660	$8.338 \times 10^{-3}$	$6.909 \times 10^{-3}$
Mesh3	2880	115884	$3.774 \times 10^{-3}$	$2.233 \times 10^{-3}$
Mesh4	3600	139840	$1.528 \times 10^{-3}$	Overflow

# 4 Conclusions and Future Work

In this paper, a simplified single-phase 3D steady PEMFC model is introduced 137 by a modified Navier-Stokes equations for mass and momentum, and a conser- 138 vation equation for water concentration. Based on the combined finite element- 139 upwind finite volume methods and the overlapping domain decomposition method, 140 a new discretization scheme is designed and implemented for the PEMFC model. 141 Numerical experiments demonstrate that our methods are effective to deal with the 142 non-matching grids and obtain a relatively accurate numerical solution with low mass 143 balance error. The derived discretization scheme will be also studied for two-phase 144 unsteady and/or fuel cell stack model in our further work.

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t2 1 t2.2

t2.5

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# Multigrid Methods for the Biharmonic Problem with Cahn-Hilliard Boundary Conditions

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1 Introduction 11

Let  $\Omega \subset \mathbb{R}^2$  be a bounded polygonal domain,  $V = \{v \in H^2(\Omega) : \partial v/\partial n = 0 \text{ on } 12 \partial \Omega\}$  and  $f \in L_2(\Omega)$ . In this paper we consider multigrid methods for the following 13 biharmonic problem: Find  $u \in V$  such that

$$\int_{\Omega} \nabla^2 u : \nabla^2 v \, dx = \int_{\Omega} f v \, dx \qquad \forall v \in V, \tag{1}$$

where  $\nabla^2 w$ :  $\nabla^2 v = \sum_{i,j=1}^2 w_{x_i x_j} v_{x_i x_j}$  is the inner product of the Hessian matrices of w 15 and v. Under the (assumed) compatibility condition,

$$\int_{\Omega} f \, dx = 0,\tag{2}$$

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the biharmonic problem (1) is solvable and the solution is unique up to an additive 17 constant. Furthermore we have an elliptic regularity estimate 18

$$\|\hat{u}\|_{H^{2+\alpha}(\Omega)} \le C\|f\|_{L_2(\Omega)} \tag{3}$$

for the solution  $\hat{u}$  of (1) that satisfies  $\int_{\Omega} \hat{u} dx = 0$ . Note that, unlike the biharmonic 19 problem with the boundary conditions of clamped plates, the index of elliptic regularity  $\alpha$  in (3), which is determined by the angles of  $\Omega$ , can be close to 0 even if  $\Omega$  21 is convex (cf. [2]).

The essential boundary condition  $\partial u/\partial n = 0$  and the natural boundary condition 23  $\partial (\Delta u)/\partial n = 0$  satisfied by the solution u of (1) appear in the Cahn-Hilliard model 24 for phase separation phenomena (cf. [8]). In particular, the boundary value problem 25 (1) appears when the Cahn-Hilliard equation is discretized in time by an implicit 26 method and the resulting nonlinear fourth order elliptic boundary value problem is 27 solved by an Newton iteration.

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We will describe a  $C^0$  interior penalty method for (1) in Sect. 2 and introduce in 29 Sect. 3 multigrid methods that are based on a new smoother. The convergence properties of the multigrid methods are briefly discussed in Sect. 4, followed by numerical 31 results in Sect. 5.

# 2 A Quadratic C<sup>0</sup> Interior Penalty Method

 $C^0$  interior penalty methods (cf. [6, 9]) are discontinuous Galerkin methods for fourth 34 order problems. Let  $\mathscr{T}_h$  be a simplicial triangulation of  $\Omega$ ,  $V_h \subset H^1(\Omega)$  be the associated  $P_2$  Lagrange finite element space (cf. [5]), and  $\hat{V}_h$  be the subspace of  $V_h$  consisting of functions with zero mean, i.e.,  $v \in V_h$  belongs to  $\hat{V}_h$  if and only if  $\int_{\Omega} v \, dx = 0$ . 37 The quadratic  $C^0$  interior penalty method for (1) is to find  $\hat{u}_h \in \hat{V}_h$  such that

$$a_h(\hat{u}_h, v) = \int_{\Omega} f v \, dx \qquad \forall v \in \hat{V}_h, \tag{4}$$

where 39

$$a_{h}(\hat{u}_{h}, v) = \int_{\Omega} f v dx \qquad \forall v \in \hat{V}_{h},$$

$$a_{h}(w, v) = \sum_{T \in \mathcal{T}_{h}} \int_{T} \nabla^{2} w : \nabla^{2} v dx + \sum_{e \in \mathcal{E}_{h}} \int_{e} \left\{ \left\{ \frac{\partial^{2} w}{\partial n^{2}} \right\} \right\} \left[ \left[ \frac{\partial v}{\partial n} \right] \right] ds$$

$$+ \sum_{e \in \mathcal{E}_{h}} \int_{e} \left\{ \left\{ \frac{\partial^{2} v}{\partial n^{2}} \right\} \right\} \left[ \left[ \frac{\partial w}{\partial n} \right] \right] ds + \sum_{e \in \mathcal{E}_{h}} \int_{e} \left[ \left[ \frac{\partial w}{\partial n} \right] \right] \left[ \left[ \frac{\partial v}{\partial n} \right] \right] ds.$$

$$(5)$$

Here  $\mathcal{E}_h$  is the set of the edges in  $\mathcal{T}_h$ ,  $\{\{\partial^2 v/\partial n^2\}\}$  (resp.  $[\![\partial v/\partial n]\!]$ ) is the average of 40 the second normal derivative of v (resp. the jump of the first normal derivative of v) 41 across an edge, |e| is the length of the edge e, and  $\sigma > 0$  is a penalty parameter.

The quadratic  $C^0$  interior penalty method is consistent. It is also stable if  $\sigma$  is 43 sufficiently large, which is assumed to be the case. (The magnitude of  $\sigma$  is related to certain inverse estimates. It can be taken to be 5 in practice.) It can be shown (cf. [3]) 45 that the solution  $\hat{u}_h$  of (4) satisfies the following error estimate: 46

$$\|\hat{u} - \hat{u}_h\|_h \le Ch^\alpha \|f\|_{L_2(\Omega)},\tag{6}$$

where  $\hat{u}$  is the zero mean solution of (1),  $\alpha$  is the index of elliptic regularity in (3), 47 and the norm  $\|\cdot\|_h$  is given by

$$\|v\|_h^2 = \sum_{T \in \mathcal{T}_h} |v|_{H^2(T)}^2 + \sum_{e \in \mathcal{E}_h} |e|^{-1} \| [\![ \partial v / \partial n ]\!] \|_{L_2(e)}^2.$$

C<sup>0</sup> interior penalty methods have certain advantages over other finite element 49 methods for fourth order problems. They are simpler than conforming methods 50 which require  $C^1$  elements. They come in a natural hierarchy that can capture smooth 51 solutions efficiently, which is not the case for classical nonconforming methods. Un- 52 like mixed methods they preserve the positive definiteness of the continuous problem 53 and are easier to develop for more complicated problems (cf. [9]). 54

Another significant advantage of  $C^0$  interior penalty methods comes from the 55 fact that the underlying finite element spaces are standard spaces for second order 56 problems. (Note that the essential boundary condition for (1) is only enforced weakly 57 in (4) and the finite element space  $V_h$  does not involve any boundary condition.) 58 Therefore multigrid solves for second order problems can be easily implemented as 59 a preconditioner. By using such a preconditioner in the smoothing steps of multigrid 60 algorithms for fourth order problems, the performance of the smoother and hence 61 the overall performance of the multigrid algorithms can be significantly improved. 62 This approach was carried out in [7] for the biharmonic problem with the boundary 63 conditions of clamped plates. Below we will use this approach to develop multigrid 64 methods for (4).

# 3 Multigrid Methods

Let  $\mathscr{T}_k$   $(k=0,1,\cdots)$  be a sequence of simplicial triangulations obtained from the 67 initial triangulation  $\mathscr{T}_0$  by uniform refinement. We will use  $V_k$  (resp.  $a_k(\cdot,\cdot)$ ) to denote 68 the finite element space (resp. the bilinear form for the  $C^0$  interior penalty method) 69 associated with  $\mathscr{T}_k$ .

Let  $V_k'$  be the dual space of  $V_k$  and  $\hat{V}_k = \{v \in V_k : \int_{\Omega} v \, dx = 0\}$  be the zero-mean 71 subspace of  $V_k$ . We can identify  $\hat{V}_k'$  with the subspace of  $V_k'$  whose members annihilate 72 the constant functions, i.e.,  $\hat{V}_k' = \{\gamma \in V_k' : \langle \gamma, 1 \rangle = 0\}$ , where  $\langle \cdot, \cdot \rangle$  is the canonical 73 bilinear form between a vector space and its dual.

Let the operator  $A_k: V_k \longrightarrow \hat{V}'_k$  be defined by  $\langle A_k v, w \rangle = a_k(v, w)$  for all  $v, w \in V_k$ . 75 We can then rewrite the discrete problem (4) as  $A_k \hat{u}_k = \phi_k$ , where  $\hat{u}_k \in \hat{V}_k$  and  $\phi_k \in \hat{V}'_k$  76 satisfies  $\langle \phi_k, v \rangle = \int_{\Omega} f v \, dx$  for all  $v \in V_k$ . Below we will develop multigrid algorithms 77 for equations of the form

$$A_k z = \psi \tag{7}$$

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where  $z \in \hat{V}_k$  and  $\psi \in \hat{V}'_k$ .

There are two ingredients in the design of multigrid algorithms. First of all, we 80 need intergrid transfer operators to move data between consecutive levels. Since 81 the finite element spaces are nested, we can take the coarse-to-fine operator  $I_{k-1}^k$ : 82  $V_{k-1} \longrightarrow V_k$  to be the natural injection and the fine-to-coarse operator  $I_k^{k-1}: V_k' \longrightarrow$  83  $V_{k-1}'$  to be the transpose of  $I_{k-1}^k$  with respect to the canonical bilinear forms, i.e., 84  $\langle I_k^{k-1} \gamma, \nu \rangle = \langle \gamma, I_{k-1}^k \nu \rangle$  for all  $\gamma \in V_k'$ ,  $\nu \in V_{k-1}$ . Note that  $I_{k-1}^k$  maps  $\hat{V}_{k-1}$  into  $\hat{V}_k$  and 85 consequently  $I_k^{k-1}$  maps  $\hat{V}_k'$  into  $\hat{V}_{k-1}'$ .

The second ingredient is a good smoother that can damp out the highly oscillatory part of the error of an approximate solution so that the remaining part of the error 88 can be captured accurately on a coarser grid. Here we take advantage of the fact that 89 the  $P_2$  Lagrange finite element space is a standard space for second order problems 90 to incorporate a multigrid Poisson solve in the smoother. Let  $L_k: \hat{V}_k \longrightarrow \hat{V}'_k$  be the 91 discrete Laplace operator defined by 92

$$\langle L_k v, w \rangle = \int_{\Omega} \nabla v \cdot \nabla w \, dx \qquad \forall v, w \in \hat{V}_k.$$

We take  $S_k^{-1}: \hat{V}_k' \longrightarrow \hat{V}_k$  to be an approximate inverse of  $L_k$  obtained from a multigrid 94 Poisson solve such that

$$\langle S_k v, v \rangle \approx |v|_{H^1(\Omega)}^2 \qquad \forall v \in \hat{V}_k.$$
 (8)

The smoothing step in our multigrid algorithms for (7) is then given by

$$z_{\text{new}} = z_{\text{old}} + \lambda_k S_k^{-1} (\psi - A_k z_{\text{old}}), \tag{9}$$

where  $\lambda_k$  is a damping factor chosen so that the spectral radius  $\rho(\lambda_k S_k^{-1} A_k)$  is <2. It follows from (8) and standard inverse estimates (cf. [5]) that we can take  $\lambda_k = Ch_k^2$ . 98 Note that the computational cost of (9) is proportional to the dimension of  $\hat{V}_k$ , which 99 implies that the overall computational costs of the multigrid algorithms in Sects. 3.1 and 3.2 are also proportional to the dimension of  $\hat{V}_k$ .

We can now describe the V-cycle and W-cycle algorithms (cf. [10]) in terms of 102 the integrid transfer operators and the smoothing scheme. 103

#### 3.1 V-Cycle Algorithm

The V-cycle algorithm computes an approximate solution  $MG_V(k, \psi, z_0, m)$  of (7) 105 with initial guess  $z_0 \in \hat{V}_k$  and m pre-smoothing and m post-smoothing steps. For 106 k=0, we take  $MG_V(0,\psi,z_0,m)$  to be the output of a direct solve. For  $k\geq 1$ , we compute  $MG_V(k, \psi, z_0, m)$  recursively in three steps. 108

*Pre-smoothing* For  $1 \le \ell \le m$ , compute  $z_{\ell}$  recursively by

$$z_{\ell} = z_{\ell-1} + \lambda_k S_k^{-1} (\psi - A_k z_{\ell-1}).$$

Coarse Grid Correction Compute

$$z_{m+1} = z_m + I_{k-1}^k MG_V(k-1, \rho_{k-1}, 0, m),$$

where  $\rho_{k-1} = I_k^{k-1}(\psi - A_k z_m) \in \hat{V}_{k-1}'$  is the transferred residual of  $z_m$ . 111

*Post-smoothing* For  $m+2 \le \ell \le 2m+1$ , compute  $z_{\ell}$  recursively by

$$z_{\ell} = z_{\ell-1} + \lambda_k S_k^{-1} (\psi - A_k z_{\ell-1}).$$

The final output is  $MG_V(k, \psi, z_0, m) = z_{2m+1}$ .

#### 3.2 W-Cycle Algorithm

The W-cycle algorithm computes an approximate solution  $MG_W(k, \psi, z_0, m)$  of (7) 115 with initial guess  $z_0 \in \hat{V}_k$  and m pre-smoothing and m post-smoothing steps. The only difference between the V-cycle algorithm and the W-cycle algorithm is in the coarse grid correction step, where the coarse grid algorithm is applied twice to the coarse 118 grid residual equation. More precisely, we have 119

$$\begin{split} z_{m+\frac{1}{2}} &= MG_W(k-1,\rho_{k-1},0,m), \\ z_{m+1} &= z_m + MG_W(k-1,\rho_{k-1},z_{m+\frac{1}{2}},m). \end{split}$$

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Remark 1. For simplicity we have described the multigrid algorithms in terms of the 121 space  $\hat{V}_k$  where the bilinear form  $a_k(\cdot,\cdot)$  is nonsingular. But the multigrid Poisson 122 solve  $S_k^{-1}$  (and hence the V-cycle and W-cycle algorithms) can be implemented on 123  $V_k$  for  $k \ge 1$ . The implementation of multigrid algorithms for the singular Neumann 124 problem is discussed for example in [1].

# **4 Convergence Properties**

Let  $z_0 \in \hat{V}_k$  be the initial guess and  $z_{\dagger} \in \hat{V}_k$  be the output of the *V*-cycle or *W*-cycle 127 algorithm for (7). Numerical results indicate that

$$||z - z_{\dagger}||_{a_h} \le Cm^{-\alpha} ||z - z_0||_{a_h}, \tag{10}$$

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where  $\alpha$  is the index of elliptic regularity in (3) and  $\|\cdot\|_{a_h} = \sqrt{a_h(\cdot,\cdot)}$  is the energy 129 norm, provided that the number of smoothing steps  $m \ge m_*$ . Here  $m_*$  is a sufficiently large positive integer independent of k. In particular the multigrid algorithms are 131 contractions for sufficiently large m and the contraction numbers are bounded away 132 from 1 uniformly. A similar estimate was obtained in [7] for the boundary conditions 133 of clamped plates. The derivation of (10) for the Cahn-Hilliard boundary conditions 134 will be carried out in [4] where general fourth order problems are considered.

A significant benefit of including a multigrid Poisson solve in the smoothing step 136 (9) is that the resulting smoothing property is similar to that for second order problems (cf. [7]) so that the contraction number estimate (10) is also similar to that for 138 second order problems. Indeed, because of the estimate (8), we can derive a smoothing property for (9) with respect to a family of mesh dependent norms  $\|\cdot\|_{s,k}$  such 140 that  $\|\|\cdot\|\|_{0,k} \approx |\cdot|_{H^1(\Omega)}$  and  $\|\|\cdot\|\|_{1,k} \approx |\cdot|_{H^2(\Omega)}$  on the space  $\hat{V}_k$ . Note that the smoothing properties of standard smoothers for second order problems are described in terms 142 of mesh dependent norms  $\|\cdot\|_{s,k}$  such that  $\|\cdot\|_{0,k} \approx \|\cdot\|_{L_2(\Omega)}$  and  $\|\cdot\|_{1,k} \approx |\cdot|_{H^1(\Omega)}$ on the finite element spaces. The good performance of the smoothing step (9) is due 144 to the similarity between the Hilbert scales  $[H^1(\Omega), H^2(\Omega)]$  and  $[L_2(\Omega), H^1(\Omega)]$ .

If we use a standard smoother such as the Richardson relaxation in a multigrid 146 algorithm for (7), then the smoothing property will be determined by the Hilbert 147 scale  $[L_2(\Omega), H^2(\Omega)]$ . In this case the estimate (10) will be replaced by the estimate

$$||z - z_{\dagger}||_{a_h} \le Cm^{-\alpha/2}||z - z_0||_{a_h},$$
 (11)

which means that the effect of 100 smoothing steps without the preconditioner is 149 roughly equivalent to the effect of 10 smoothing steps with the preconditioner. As 150 far as we know, all existing multigrid methods for fourth order problems (except 151 those in [6]) use standard smoothers and their convergence is governed by (11).

#### 5 Numerical Results

The numerical experiments were performed on sienna@IMA (Intel P4, 3.4 GHz 154 CPU, 2 G memory) at the Institute for Mathematics and its Applications. In the numerical experiments we take  $\sigma = 5$  and the preconditioner to be a V-cycle Poisson 156 solve with one pre-smoothing step and one post-smoothing step. (Other multigrid 157 Poisson solves can also be used, but the V(1,1) solve appears to be the most efficient.) The contraction numbers for the V-cycle and W-cycle algorithms on the unit 159 square (with two elements in the initial mesh) are reported in Tables 1 and 2. It is 160 observed that the V-cycle (resp. W-cycle) algorithm is a contraction for m > 4 (resp. 161 m > 2).

Table 1. Contraction numbers for the V-cycle algorithm on the unit square.

m k	4	5	6	7	8	9	10	11	12	13
1	0.212	0.126	0.0813	0.0594	0.0442	0.0332	0.0252	0.0192	0.0147	0.0114
2	0.329	0.223	0.190	0.164	0.142	0.124	0.109	0.0967	0.0861	0.0771
3	0.412	0.342	0.308	0.279	0.255	0.234	0.217	0.203	0.190	0.179
4	0.479	0.420	0.386	0.357	0.334	0.314	0.296	0.282	0.266	0.257
5	0.537	0.467	0.434	0.408	0.386	0.367	0.351	0.336	0.324	0.312
6	0.578	0.494	0.462	0.436	0.415	0.396	0.380	0.366	0.353	0.341
7	0.619	0.503	0.472	0.446	0.425	0.406	0.391	0.376	0.364	0.351

t1.1 t1 2 t1.3 t1 4 t1.5 t1 6 t1.7 t1 8

> t2.1 t2 2 t2.3 t2.4 t2.5 t2.6 t2 7 t2.8

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**Table 2.** Contraction numbers for the W-cycle algorithm on the unit square.

k m	2	3	4	5	6	7	8	9	10	11
1	0.661	0.368	0.212	0.126	0.0813	0.0594	0.0442	0.0332	0.0252	0.0192
2	0.483	0.360	0.291	0.241	0.203	0.172	0.148	0.128	0.112	0.0983
3	0.475	0.375	0.335	0.282	0.263	0.229	0.215	0.195	0.182	0.171
4	0.455	0.383	0.335	0.308	0.287	0.270	0.256	0.244	0.233	0.223
5	0.456	0.384	0.344	0.315	0.297	0.279	0.267	0.255	0.245	0.237
6	0.455	0.384	0.344	0.316	0.297	0.280	0.268	0.256	0.248	0.239
7	0.455	0.384	0.344	0.317	0.297	0.281	0.269	0.258	0.248	0.240

For comparison we report in Table 3 the contraction numbers for the V-cycle 163 algorithm that does not use a preconditioner in the smoothing steps. The smoothing step in this algorithm is the standard Richardson relaxation scheme.

We have also carried out numerical experiments for the L-shaped domain with 166 vertices (0,0), (1,0), (1,1), (-1,1), (-1,-1) and (0,-1). The initial mesh consists of six isosceles triangles sharing (0,0) as a common vertex. The contraction numbers 168 for the W-cycle algorithm with/without the preconditioner are presented in Tables 4 169 and 5.

We note that the contraction numbers in Table 1 (resp. Table 4) for m smoothing 171 steps are comparable to the contraction numbers in Table 3 (resp. Tables 5) for  $m^2$ smoothing steps.

**Page 134** 

t3.1 t3 2 t3.3 t3 4 t3.5 t3.6 t3.7 t3.8

t5.1 t5.2 t5.3 t5.4

t5.6 t5.7 t5.8

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Table 3. Contraction numbers for the V-cycle algorithm without a preconditioner on the unit square.

m k	21	22	23	24	25	26	27	28	29	30
1	0.428	0.410	0.392	0.376	0.361	0.346	0.332	0.320	0.307	0.296
2	0.646	0.614	0.583	0.555	0.529	0.504	0.481	0.459	0.439	0.420
3	0.770	0.728	0.690	0.654	0.621	0.591	0.562	0.535	0.510	0.487
4	0.844	0.797	0.753	0.713	0.676	0.641	0.609	0.579	0.551	0.525
5	0.895	0.843	0.795	0.752	0.711	0.674	0.639	0.607	0.577	0.548
6	0.931	0.876	0.826	0.780	0.737	0.697	0.661	0.627	0.595	0.565
7	0.960	0.902	0.849	0.801	0.757	0.715	0.677	0.642	0.609	0.578

Table 4. Contraction numbers for the W-cycle algorithm with a preconditioner on the L-shaped domain.

k m	3	5	7	9	11	13	15	17	19	21	23	t4.1
1	0.319	0.187	0.125	0.105	0.0913	0.0798	0.0699	0.0614	0.0540	0.0476	0.0420	t4.2
2	0.383	0.273	0.206	0.161	0.139	0.132	0.125	0.119	0.113	0.108	0.103	t4.3
3	0.390	0.302	0.238	0.208	0.182	0.163	0.152	0.148	0.144	0.141	0.137	t4.4
4	0.386	0.309	0.271	0.245	0.224	0.208	0.193	0.181	0.170	0.161	0.153	t4.5
5	0.384	0.315	0.279	0.255	0.237	0.222	0.209	0.198	0.189	0.180	0.172	t4.6
6	0.384	0.316	0.281	0.257	0.240	0.226	0.213	0.203	0.193	0.185	0.177	t4.7
7	0.387	0.317	0.281	0.258	0.240	0.226	0.214	0.203	0.194	0.186	0.178	t4.8

Table 5. Contraction numbers for the W-cycle algorithm without a preconditioner on the L-shaped domain.

k	m	5	7	9	11	13	15	17	19	21	23
	1	0.943	0.788	0.680	0.600	0.537	0.486	0.443	0.407	0.375	0.347
	2	0.790	0.585	0.505	0.459	0.426	0.394	0.375	0.358	0.342	0.328
١	3	0.666	0.512	0.469	0.456	0.434	0.416	0.400	0.386	0.373	0.362
	4	0.580	0.519	0.484	0.454	0.434	0.418	0.405	0.394	0.385	0.376
	5	0.581	0.527	0.491	0.465	0.444	0.427	0.414	0.402	0.392	0.384
	6	0.587	0.531	0.494	0.467	0.446	0.429	0.415	0.404	0.394	0.386
	7	0.587	0.530	0.493	0.467	0.446	0.429	0.415	0.404	0.394	0.386

Finally we compare the computational cost between the preconditioned schemes 174 and the un-preconditioned schemes. On the unit square, the contraction numbers for 175 the preconditioned V-cycle algorithm with m = 4 (cf. Table 1) are about the same as 176 the contraction numbers for the un-preconditioned V-cycle algorithm with  $m=29\,$  177 (cf. Table 3). For k = 7, the former takes  $1.4 \times 10^8$  floating point operations and 178 0.55 s while the latter takes  $3.2 \times 10^8$  floating point operations and 1.2 s.

On the L-shaped domain, the contraction numbers for the preconditioned W-cycle algorithm with $m=3$ (cf. Table 4) are about the same as the contraction numbers for the un-preconditioned W-cycle algorithm with $m=23$ (cf. Table 5). For $k=7$ , the former takes $4.7 \times 10^8$ floating point operations and $2.1$ s while the latter takes $1.1 \times 10^9$ floating point operations and $4.7$ s.	180 181 182 183 184
1.1 × 10 Houting point operations and 4.7 s.	104
<b>Acknowledgments</b> This work was supported in part by the National Science Foundation under Grant No. DMS-10-16332 and by the Institute for Mathematics and its Applications with funds provided by the National Science Foundation.	185 186 187
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# A Two-Level Additive Schwarz Preconditioner for $\mathbb{C}^0$ Interior Penalty Methods for Cahn-Hilliard Equations

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**Summary.** We study a two-level additive Schwarz preconditioner for  $C^0$  interior penalty 7 methods for a biharmonic problem with essential and natural boundary conditions with Cahn-8 Hilliard type. We show that the condition number of the preconditioned system is bounded 9 by  $C(1+(H^3/\delta^3))$ , where H is the typical diameter of a subdomain,  $\delta$  measures the overlap 10 among the subdomains, and the positive constant C is independent of the mesh sizes and the 11 number of subdomains.

1 Introduction

Let  $\Omega$  be a bounded polygonal domain in  $\mathbb{R}^2$ , and  $\mathbb{V} = \{v \in H^2(\Omega) : \partial v/\partial n = 0 \text{ on } 14 \partial \Omega\}$ , where  $\partial/\partial n$  denotes the outward normal derivative. Consider the following 15 model problem which is the weak form of the biharmonic problem with boundary 16 conditions of Cahn-Hilliard type:

Find  $u \in H^2(\Omega)$  such that

$$a(u,v) = (f,v) \quad \forall v \in \mathbb{V},$$
 (1)

$$\frac{\partial u}{\partial n} = 0 \qquad \text{on } \partial\Omega, \tag{2}$$

where  $f \in L_2(\Omega)$ ,  $(\cdot,\cdot)$  is the  $L_2(\Omega)$  inner product, and

$$a(w,v) = \sum_{i,j=1}^{2} \int_{\Omega} \frac{\partial^{2} w}{\partial x_{i} \partial x_{j}} \frac{\partial^{2} v}{\partial x_{i} \partial x_{j}} dx$$

is the inner product of the Hessian matrices of w and v.

Let  $p_*$  be a corner of  $\Omega$ , and

$$\mathbb{V}^* = \{ v \in \mathbb{V} : v(p_*) = 0 \}.$$
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Then by elliptic regularity [1], the unique solution  $u \in V^*$  of our model problem 24 belongs to  $H^{2+\alpha}(\Omega)$ , where  $0 < \alpha \le 2$  is the index of elliptic regularity. 25

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 $C^0$  interior penalty methods are discontinuous Galerkin methods for fourth order 26 problems. These approaches for our model problem have recently been analyzed in 27 [5]. Let  $\mathcal{T}_h$  be a simplicial or convex quadrilateral triangulation of  $\Omega$ , and  $V_h$  be a 28 Lagrange (triangular or tensor product) finite element space associated with  $\mathcal{T}_h$ . Let 29

$$V_h^* = \{ v \in V_h : v(p_*) = 0 \}.$$
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Then the  $C^0$  interior penalty method for (1) and (2) is to find  $u_h \in V_h^*$  such that

$$\mathscr{A}_h(u_h, v) = (f, v) \qquad \forall v \in V_h^*, \tag{3}$$

where for  $w, v \in V_h^*$ ,

$$\mathcal{A}_{h}(w,v) = \sum_{D \in \mathcal{T}_{h}} \sum_{i,j=1}^{2} \int_{D} \frac{\partial^{2} w}{\partial x_{i} \partial x_{j}} \frac{\partial^{2} v}{\partial x_{i} \partial x_{j}} dx + \sum_{e \in \mathcal{E}_{h}} \frac{\eta}{|e|} \int_{e} \left[ \left[ \frac{\partial w}{\partial n} \right] \right] \left[ \left[ \frac{\partial v}{\partial n} \right] \right] ds + \sum_{e \in \mathcal{E}_{h}} \int_{e} \left( \left\{ \left[ \frac{\partial^{2} w}{\partial n^{2}} \right] \right\} \left[ \left[ \frac{\partial v}{\partial n} \right] \right] + \left\{ \left[ \frac{\partial^{2} v}{\partial n^{2}} \right] \right\} \left[ \left[ \frac{\partial w}{\partial n} \right] \right] \right) ds,$$

$$(4)$$

 $\mathcal{E}_h$  denotes the set of edges of the triangulation  $\mathcal{T}_h$ , and  $\eta$  is a penalty parameter. The jumps and averages are defined as follows.

For interior edges  $e \in \mathcal{E}_h$  shared by two elements  $D_{\pm} \in \mathcal{T}_h$ , we take  $n_e$  to be the 35 unit normal of e pointing from  $D_{-}$  into  $D_{+}$ , and define 36

$$\left[ \left[ \frac{\partial v}{\partial n} \right] \right] = \frac{\partial v_+}{\partial n_e} - \frac{\partial v_-}{\partial n_e} \quad \text{and} \quad \left\{ \left\{ \frac{\partial^2 v}{\partial n^2} \right\} \right\} = \frac{1}{2} \left( \frac{\partial^2 v_+}{\partial n_e^2} + \frac{\partial^2 v_-}{\partial n_e^2} \right),$$
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where  $v_{\pm} = v |_{D_{+}}$ . Note that the definitions of  $[\![\partial v/\partial n]\!]$  and  $\{\![\partial^{2}v/\partial n^{2}]\!\}$  are independent of the choice of e.

For  $e \in \mathcal{E}_h$  which is on the boundary of  $\Omega$ , we take  $n_e$  to be the unit normal of  $e^{-40}$ pointing outside  $\Omega$  and define 41

$$\left[ \left[ \frac{\partial v}{\partial n} \right] \right] = -\frac{\partial v}{\partial n_e} \quad \text{and} \quad \left\{ \left\{ \frac{\partial^2 v}{\partial n^2} \right\} \right\} = \frac{\partial^2 v}{\partial n_e^2}.$$

Remark 1. The discrete problem (3) resulting from the  $C^0$  interior penalty method is 43 consistent, and for the penalty parameter  $\eta$  large enough, it is also stable [3].

For fourth order problems,  $C^0$  interior penalty methods have certain advantages 45 over classical finite element methods. However, due to the nature of fourth order 46 problems, the discrete system resulting from the  $C^0$  interior penalty method is very 47 ill-conditioned. Therefore, it is necessary to develop modern fast solvers to overcome 48 this drawback. In this paper, we construct a two-level additive Schwarz preconditioner and extend the results in [4] for biharmonic problems with essential Dirichlet 50 boundary conditions to the ones with the essential and natural boundary conditions. 51

The rest of this paper is organized as follows. We first introduce the framework 52 of a two-level additive Schwarz preconditioner in Sect. 2, followed by the condition 53 number estimates of the preconditioned system in Sect. 3. Section 4 demonstrates 54 some numerical results.

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### 2 A Two-Level Additive Schwarz Preconditioner

For simplicity, we will focus on the case where  $\mathcal{T}_h$  is a rectangular mesh. The results obtained in this paper are also true for triangular and general convex quadrilateral meshes.

Let  $V_h^* = \{v : v \in C(\bar{\Omega}), v(p_*) = 0, v_D = v \big|_D = \mathbb{Q}_2(D) \ \forall D \in \mathscr{T}_h \}$  be the standard 60 quadratic Lagrange finite element space associated with  $\mathscr{T}_h$ , and the operator  $A_h$ : 61  $V_h^* \longrightarrow V_h^{*'}$  can then be defined by

$$\langle A_h v, w \rangle = \mathscr{A}_h(v, w) \qquad \forall v, w \in V_h^*,$$

where  $\langle \cdot, \cdot \rangle$  is the canonical bilinear form between a vector space and its dual.

Note that for  $\eta$  sufficiently large, the following relation [3] is true.

$$C_1|v|^2_{H^2(\Omega,\mathscr{T}_h)} \le \langle A_h v, v \rangle \le C_2|v|^2_{H^2(\Omega,\mathscr{T}_h)} \qquad \forall v \in V_h^*,$$

where

$$|v|_{H^2(\Omega,\mathcal{T}_h)}^2 = \sum_{D \in \mathcal{T}_h} |v|_{H^2(D)}^2 + \sum_{e \in \mathcal{E}_h} \frac{1}{|e|} \| [\![\partial v/\partial n]\!] \|_{L_2(e)}^2,$$

and the constants  $C_1$  and  $C_2$  depend only on the shape regularity of  $\mathcal{T}_h$ .

We now construct a two-level additive Schwarz preconditioner for the operator  $A_h$  which involves a coarse grid solve and subdomain solves.

First of all, let  $\mathscr{T}_H$  be a coarse rectangular mesh for  $\Omega$ , and  $V_0 \subset H^1(\Omega)$  be the 69  $\mathbb{Q}_1$  finite element space associated with  $\mathscr{T}_H$ . We define  $A_0: V_0^* \longrightarrow V_0^{*'}$  by 70

$$\langle A_0 v, w \rangle = \mathscr{A}_H(v, w) \qquad \forall v, w \in V_0^*,$$

where  $\mathscr{A}_H$  is the analog of  $\mathscr{A}_h$  for the coarse grid  $\mathscr{T}_H$ , and  $V_0^* = \{v : v \in V_0, v(p_*) = 710\}$ .

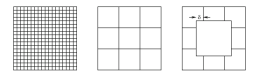
Let  $\Omega_j, 1 \leq j \leq J$ , be overlapping subdomains of  $\Omega$  such that  $\Omega = \bigcup_{j=1}^J \Omega_j$ , and 73 the boundaries of  $\Omega_j$  are aligned with the edges of  $\mathscr{T}_h$ . We assume that there exist 74 nonnegative  $\theta_j \in C^{\infty}(\bar{\Omega})$  for  $1 \leq j \leq J$  such that

$$\begin{split} \theta_j &= 0 \qquad & \text{on} \quad \Omega \backslash \Omega_j, \\ \sum_{j=1}^J \theta_j &= 1 \qquad & \text{on} \quad \bar{\Omega}, \\ \| \nabla \theta_j \|_{L_{\infty}(\Omega)} &\leq \frac{C}{\delta}, \qquad & \| \nabla^2 \theta_j \|_{L_{\infty}(\Omega)} \leq \frac{C}{\delta^2}, \end{split}$$

where  $\nabla^2 \theta_j$  is the Hessian of  $\theta_j$ ,  $\delta > 0$  measures the overlap among the subdomains, 76 and C is a positive constant independent of h, H and J.

Remark 2. Suppose  $\mathcal{T}_h$  is a refinement of  $\mathcal{T}_H$ . We can construct  $\Omega_j$  by enlarging the 78 elements of  $\mathcal{T}_H$  by the amount of  $\delta$  so that the boundaries of  $\Omega_j, 1 \leq j \leq J$ , are 79 aligned with the edges of  $\mathcal{T}_h$  (cf. Fig. 1). The construction of  $\theta_j, 1 \leq j \leq J$ , is then 80 standard.

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**Fig. 1.**  $\mathcal{T}_h$ ,  $\mathcal{T}_H$  and  $\Omega_i$ 

Moreover, we assume that the maximum number of subdomains  $\Omega_i$  that share a common point is bounded by a constant  $N_c$ . 83

Let  $V_j = \{v : v \in V_h^*, v = 0 \text{ on } \bar{\Omega}_\ell \text{ if } \ell \neq j\}$  be the  $\mathbb{Q}_2$  finite element space asso-84 ciated with  $\mathscr{T}_h$  on  $\bar{\Omega}_j$ . Then we define the operator  $A_j: V_j \longrightarrow V_j'$  by  $\langle A_j v, w \rangle = \mathscr{A}_j(v, w) \qquad \forall v, w \in V_j,$ 85

$$\langle A_j v, w \rangle = \mathscr{A}_j(v, w) \qquad \forall v, w \in V_j,$$

where  $\mathscr{A}_i, 1 \leq j \leq J$ , are the analogs of  $\mathscr{A}_h$  restricted on  $\bar{\Omega}_j$ . Similarly, we obtain 86 that

$$C_3|v|^2_{H^2(\Omega_j,\mathscr{T}_h)} \le \langle A_j v, v \rangle \le C_4|v|^2_{H^2(\Omega_j,\mathscr{T}_h)} \qquad \forall v \in V_j,$$

where

$$|v|^2_{H^2(\Omega_j,\mathscr{T}_h)} = \sum_{\substack{D \in \mathscr{T}_h \\ D \subset \Omega_j}} |v|^2_{H^2(D)} + \sum_{\substack{e \in \mathscr{E}_h \\ e \subset \Omega_j}} \| [\![\partial v/\partial n]\!] \|^2_{L_2(e)},$$

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and  $C_3$ ,  $C_4$  are constants independent of h, H, J,  $N_c$  and  $\delta$ .

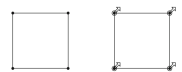
For simplicity, from now on, we will use C to denote a generic positive constant 90 independent of  $h, H, \delta$ , and J that will take different values in different occurrences. 91

The subdomain finite element space  $V_j$ ,  $1 \le j \le J$ , is connected to  $V_h^*$  by the 92 natural injection operator  $I_i$  which satisfies the following inequality. 93

$$|I_j v|_{H^2(\Omega, \mathscr{T}_h)} \le C|v|_{H^2(\Omega_i, \mathscr{T}_h)} \qquad \forall v \in V_j.$$

Furthermore, the coarse space  $V_0^st$  and the fine space  $V_h^st$  are connected by the 94 operator  $I_0$  which is defined as follows.

Let  $\tilde{V}_0 \subset H^2(\Omega)$  be the  $\mathbb{Q}_3$  Bogner-Fox-Schmit finite element space associated 96 with  $\mathcal{T}_H$ , and  $\tilde{V}_0^* = \{v : v \in \tilde{V}_0, v(p_*) = 0\}$ . The  $\mathbb{Q}_1$  Lagrange element and the  $\mathbb{Q}_3$  97 Bogner-Fox-Schmit element are depicted in Fig. 2, where we use the solid dot ● to 98 denote pointwise evaluation of the shape functions, the circle  $\circ$  and the arrow  $\nearrow$  to 99 denote pointwise evaluation of all the first order derivatives and the mixed second 100 order derivative of the shape functions, respectively.



**Fig. 2.**  $\mathbb{Q}_1$  element and  $\mathbb{Q}_3$  Bogner-Fox-Schmit element

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We define 
$$E_H: V_0^* \longrightarrow \tilde{V}_0^*$$
 to be the operator that for all  $p \in \mathcal{T}_H$ ,

$$(E_{H}v)(p) = v(p),$$

$$\nabla(E_{H}v)(p) = \begin{cases} \frac{1}{|\mathcal{T}_{p}|} \sum_{D \in \mathcal{T}_{p}} \nabla v_{D}(p), & \text{if } p \in \Omega, \\ 0, & \text{if } p \in \partial \Omega, \end{cases}$$

$$\frac{\partial^{2}(E_{H}v)}{\partial x_{1}\partial x_{2}}(p) = \begin{cases} \frac{1}{|\mathcal{T}_{p}|} \sum_{D \in \mathcal{T}_{p}} \frac{\partial^{2}v_{D}}{\partial x_{1}\partial x_{2}}(p), & \text{if } p \in \Omega, \\ 0, & \text{if } p \in \partial \Omega, \end{cases}$$

where  $\mathscr{T}_p$  is the set of rectangles in  $\mathscr{T}_H$  sharing p as a vertex,  $|\mathscr{T}_p|$  is the number of elements in  $\mathscr{T}_p$ , and  $v_D = v|_D$ .

Then for all  $v \in V_0^*$ , we take  $I_0 v \in V_h^*$  to be the one whose nodal values are 105 identical with the corresponding nodal values of  $E_H v$ . 106

Remark 3. Instead of using the operator  $E_H$ , if we define the operator  $I_0$  as the natural injection operator from  $V_0^*$  to  $V_h^*$ , then the performance of the preconditioner will be 108 affected by the different scalings that appear in the penalty terms for  $\mathcal{A}_h$  and  $\mathcal{A}_H$ . 109 However, this problems can be avoided by defining  $I_0$  as above since  $E_H v \in H^2(\Omega)$ . 110

We can now define the two-level additive Schwarz preconditioner  $B: V_h^{*'} \longrightarrow V_h^*$  111 by

$$B = \sum_{j=0}^{J} I_{j} A_{j}^{-1} I_{j}^{t},$$

where  $I_j^t: V_h^{*'} \longrightarrow V_j'$  is the transpose of  $I_j$ , i.e.,  $\langle I_j^t \Psi, v \rangle = \langle \Psi, I_j v \rangle \qquad \forall \Psi \in V_h^{*'}, v \in V_j.$ 

$$\langle I_j^t \Psi, v \rangle = \langle \Psi, I_j v \rangle \qquad \forall \Psi \in V_h^{*'}, v \in V_j.$$

From the additive Schwarz theory [2, 6], the preconditioner B is symmetric positive definite and therefore the eigenvalues of  $BA_h$  are positive. Moreover, the maximum and minimum eigenvalues of  $BA_h$  are given by the following formulas, which will be used to estimate the condition number of the preconditioned system. 117

$$\lambda_{\max}(BA_h) = \max_{\substack{v \in V_h \\ v \neq 0}} \frac{\langle A_h v, v \rangle}{\min_{\substack{v = \sum_{j=0}^J I_j v_j \\ v_j \in V_j}} \sum_{j=0}^J \langle A_j v_j, v_j \rangle},$$

$$\lambda_{\min}(BA_h) = \min_{\substack{v \in V_h \\ v \neq 0}} \frac{\langle A_h v, v \rangle}{\min_{\substack{v = \sum_{j=0}^{J} I_j v_j \\ v_i \in V_i}} \sum_{j=0}^{J} \langle A_j v_j, v_j \rangle}.$$

#### 3 Condition Number Estimates

From the construction of our two-level additive Schwarz preconditioner, by the similar arguments as we did in [4], it is not difficult to derive the following results on the 120 estimates of the eigenvalues of the preconditioned system.

**Theorem 1.** The following upper bound for the eigenvalues of  $BA_h$  holds:

$$\lambda_{\max}(BA_h) \leq C,$$

where the positive constant C depends on the shape regularity of  $\mathcal{T}_h$  and  $\mathcal{T}_H$  but not 123  $h, H, \delta \text{ nor } J.$ 124

**Theorem 2.** The following lower bound for the eigenvalues of  $BA_h$  holds:

$$\lambda_{\min}(BA_h) \geq C\left(1 + \frac{H^4}{\delta^4}\right),$$

where the positive constant C depends on the shape regularity of  $\mathcal{T}_h$  and  $\mathcal{T}_H$  but not 126  $h, H, \delta$  nor J. 127

Finally, from Theorems 1 and 2, the following estimate on the condition number of 128 the preconditioned system can be obtained immediately.

**Theorem 3.** It holds that

 $\kappa(BA_h) = \frac{\lambda_{\max}(BA_h)}{\lambda_{\min}(BA_h)} \le C\left(1 + \frac{H^4}{\delta^4}\right),\,$ 

where the positive constant C depends on the shape regularity of  $\mathscr{T}_h$  and  $\mathscr{T}_H$  but not 131  $h, H, \delta$  nor J. 132

Remark 4. In the case of a small overlap, i.e.  $\delta \ll H$ , the estimate on the condition 133 number of the preconditioned system can be improved to  $(1 + (H/\delta)^3)$ , provided 134 with more assumptions on the subdomains  $\Omega_i$  [4]. 135

#### 4 Numerical Results

In this section, we present some numerical results for the biharmonic problem with Cahn-Hilliard type of boundary conditions on the unit square. We choose the penalty 138 parameter in  $\mathcal{A}_h$ ,  $\mathcal{A}_H$  and  $\mathcal{A}_i$  to be 5, which guarantees the coerciveness of the variational form (4) on  $V_h^*$ .

First of all, for different choices of H and h, we generate a vector  $v_h \in V_h^*$ , compute the right-hand side vector  $g = A_h v_h$ , and apply the preconditioned conjugate 142 gradient algorithm to the system  $A_h z = g$  using our two-level additive Schwarz pre- 143 conditioner. We compute the iteration numbers needed for reducing the energy norm 144 error by a factor of  $10^{-6}$  for five random choices of  $v_h$  and then average them. The

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numbers are collected in Tables 1 and 2. Also, to illustrate the practical performance 146 of our preconditioner, such iteration numbers needed for reducing the energy norm 147 error by a factor of  $10^{-2}$  with 16 subdomains are reported in Table 3. They show that 148 the bound for the condition number of  $BA_h$  is independent of h.

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We also compute, in the case of 4 and 16 subdomains, the maximum eigenvalue, 150 the minimum eigenvalue, and the condition number of the preconditioned system for 151 the fine mesh  $h = 2^{-6}$  and various overlaps among subdomains by using Lanczos 152 methods. The results are tabulated in Tables 4 and 5. They show that the maximum 153 eigenvalue is bounded and the minimum eigenvalue increases as the overlap among 154 subdomains decreases.

**Table 1.** Average number of iterations for reducing the energy norm error by a factor of  $10^{-6}$ with H = 1/2 and J = 4

	$h = 2^{-2}$	$h = 2^{-3}$	$h = 2^{-4}$	$h = 2^{-5}$	$h = 2^{-6}$	t1.1
$\delta = 2^{-2}$	17	17	17	15	15	t1.2
$\delta = 2^{-3}$	-	20	20	19	17	t1.3
$\delta = 2^{-4}$	-	-	26	25	24	t1.4
$\delta = 2^{-5}$	-	- /	-	47	45	t1.5
$\delta = 2^{-6}$	-	- (	-	-	93	t1.6

**Table 2.** Average number of iterations for reducing the energy norm error by a factor of  $10^{-6}$ with H = 1/4 and J = 16

	$h = 2^{-3}$	$h = 2^{-4}$	$h = 2^{-5}$	$h = 2^{-6}$
$\delta = 2^{-3}$	27	29	27	24
$\delta = 2^{-4}$ $\delta = 2^{-5}$	-	28	26	24
	-	-	42	39
$\delta = 2^{-6}$	-	-	-	83

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**Table 3.** Average number of iterations for reducing the energy norm error by a factor of  $10^{-2}$  with H = 1/4 and J = 16

t3

t3

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	$h = 2^{-3}$	$h = 2^{-4}$	$h = 2^{-5}$	$h = 2^{-6}$
$\delta = 2^{-3}$	6	6	5	5
$\delta = 2^{-4}$	-	5	5	4
$\delta = 2^{-5}$	-	-	5	4
$\delta = 2^{-6}$	-	-	-	5

**Table 4.**  $\lambda_{\max}(BA_h), \lambda_{\min}(BA_h)$  and  $\kappa(BA_h)$  with  $H = 1/2, h = 2^{-6}$  and J = 4

$H/\delta$	$\lambda_{\max}(BA_h)$	$\lambda_{\min}(BA_h)$	$\kappa(BA_h)$
2	4.8394	0.4259	$1.1363 \times 10^{1}$
4	4.8029	0.3045	$1.5775 \times 10^{1}$
8	4.7526	0.1279	$3.7149 \times 10^{1}$
16	4.6600	0.0247	$1.8850 \times 10^{2}$
32	4.5849	0.0036	$1.2895 \times 10^3$

**Table 5.**  $\lambda_{\max}(BA_h)$ ,  $\lambda_{\min}(BA_h)$  and  $\kappa(BA_h)$  with  $H=1/4, h=2^{-6}$  and J=16

$H/\delta$	$\lambda_{\max}(BA_h)$	$\lambda_{\min}(BA_h)$	$\kappa(BA_h)$
2	6.5195	0.1811	$3.5992 \times 10^{1}$
4	4.8740	0.1633	$2.9852 \times 10^{1}$
8	4.6968	0.0631	$7.4402 \times 10^{1}$
16	4.5865	0.0103	$4.4698 \times 10^{2}$

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# An Algebraic Multigrid Method Based on Matching in Graphs

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1 Introduction 10

We present an Algebraic Multigrid (AMG) method for graph Laplacian problems. 11 The coarse graphs are constructed recursively by pair-wise aggregation, or matching 12 as in [3] and we use an Algebraic Multilevel Iterations (AMLI) [1, 6] for the solution 13 phase.

The two-level method constructs a splitting of the underlying vector space into 15 two subspaces  $V_S$  and  $V_P$  and then corrects the error successively on  $V_S$  and  $V_P$ . The 16 coarse space  $V_P$  is obtained using matching on the underlying graph. Such a two-level 17 method is shown to be uniformly convergent. In the AMLI method (multilevel), m 18 coarse level corrections are applied on each level. For large m, while the convergence rate of the method is comparable to that of the two-level method and, hence, 20 uniformly convergent, it is clear that the overall complexity of such method could 21 be too high for large values of m. In our approach, the AMLI convergence rate is 22 estimated solely based on the underlying two-level method, which allows us to show 23 that m=2 gives a balance between the complexity and the desired convergence rate, 24 thus, resulting in an efficient algorithm.

The paper is organized as follows. In Sect. 2 the graph Laplacian problem is described. In Sect. 3, the graph matching algorithm is introduced and it is indicated that the  $\ell_2$  projection on the coarse space is the key quantity for obtaining the multilevel estimates of the AMLI method. In Sect. 4, an analysis of a specific two-level method is presented and in Sect. 5 its convergence and complexity are estimated. In the following section, numerical results are reported.

# 2 Graph Laplacian Problems

Graph Laplacian solvers can be used as preconditioners for various discrete numeri- 33 cal models, e.g., ones arising from discretizations of partial differential equations, 34

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machine learning algorithms, and spectral clustering of images. Consider a con- 35 nected unweighted graph  $\mathscr{G} = (\mathscr{V}, \mathscr{E})$  where  $\mathscr{V}$  and  $\mathscr{E}$  are the sets of vertices and 36 edges. The graph Laplacian  $A \in \mathbb{R}^{n \times n}$ , where  $n = |\mathcal{V}|$  (cardinality of  $\mathcal{V}$ ), corre- 37 sponding to the graph  $\mathcal{G}$ , can be defined as follows:

$$(Au,v) = \sum_{k=(i,j)\in\mathscr{E}} (u_i - u_j)(v_i - v_j).$$

The matrix A is symmetric and positive semi-definite. The null space of A is one 39 dimensional, and its basis is given by  $\{1\}$ , where 1 is a vector whose components are 40 all equal to 1. Our aim here is to solve graph Laplacian problems, or to find u, such 41 that  $(u, \mathbf{1}) = 0$  and

$$Au = f$$
,

for a given f satisfying (f, 1) = 0.

We want to find an AMG method to solve graph Laplacians with simple settings, 44 so that we can estimate the performance of the AMG method, with as few assump- 45 tions introduced as possible. The construction of this AMG method can also help 46 us to derive similar methods for weighted graph Laplacian problems, which come 47 from finite element or finite difference discretizations of elliptic partial differential 48 equations, circuit simulations, and in general, network flow simulations.

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# 3 Graph Matching

Given a graph  $\mathcal{G}$ , assume that we can find a set of aggregates  $\mathcal{M}$  called a *matching*, 51 where each aggregate contains exactly two vertices, and every vertex of  $\mathscr G$  is contained in exactly one aggregate. For a certain aggregate that contains vertices i and 53j, we merge the two vertices, and the newly formed vertex, named k, is considered 54 connected to the vertex l if and only if l is connected to i or j on graph  $\mathscr{G}$ . By merging vertices in each aggregate, a reduced graph of the graph  $\mathscr{G}$  is formed. Applying 56 such a matching algorithm recursively will result in a sequence of graphs. We then 57 construct a solver for the graph Laplacian of  $\mathscr{G}$  based on the sequence of reduced 58 graphs.

In the matching  $\mathcal{M}$ , we consider the k-th aggregate as a graph  $\mathcal{G}_k = (\mathcal{V}_k, \mathcal{E}_k)$ . Let 60 Q be the  $\ell_2$ -orthogonal projection on the coarse space, which consists of vectors that 61 are piecewise constant on each set  $\mathcal{V}_k$ . An alternative definition of Q is as follows.

$$(Qu)_i = \frac{1}{|\mathcal{V}_k|} \sum_{j \in \mathcal{V}_k} u_j, \quad i \in \mathcal{V}_k.$$

Classical AMG theory suggests that the coarse space should cover, or approx- 63 imate algebraically smooth error components. Detailed explanations can be found, 64 e.g., in the appendix of [5]. In the following section, we will compute how well piecewise constant vectors can approximate smooth vectors and will discuss the properties 66 of two-level and multilevel methods using the subspace(s) associated with the pro- 67 jection Q. 68

#### 4 A Two-Level Method

Define matrices P and S for a given matching  $\mathcal{M}$ , such that

$$P \cdot e_k = e_i + e_j$$
,  $S \cdot e_k = e_i - e_j$ ,  $(i, j) \in \mathcal{V}_k$ ,

where  $e_i$  and  $e_j$  are Euclidean basis vectors. Since a prerequisite for designing an 71 efficient AMLI method is an efficient two-level method, in this section we focus on 72 two-level methods and their convergence rates. Given an initial guess  $u_0$ , a typical 73 two-level algorithm which takes as input  $u_k$  and returns the next iterate  $u_{k+1}$  is as 74 follows:

1. 
$$v = u_k + SR^{-1}S^T(f - Au_k)$$
,  
2.  $w = v + PA_c^{-1}P^T(f - Av)$ ,  
3.  $u_{k+1} = w + SR^{-T}S^T(f - Aw)$ .

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Here the matrix R is a preconditioner of  $S^TAS$ , which is the restriction of A on the 77 space  $\operatorname{range}(S) = [\operatorname{range}(P)]^{\perp}$ . The matrix  $A_c$  is an approximation of the restriction 78 of A on the coarse space  $V_c = \operatorname{range}(P)$ . In our algorithm,  $A_c$  is first defined as the 79 graph Laplacian of the unweighted coarse graph and thus  $A_c \neq P^TAP$ . We then scale 80  $A_c$  such that  $(v^TA_cv)/(v^TP^TAPv) \in [1,c_c]$ . A proper scaling results in  $c_c = 2$  for 81 P that corresponds to an aligned matching and A that is a structured grid of any 82 dimension. The matrix representation of this two-level method, denoted by G, can 83 be deduced via the error propagation matrix given as follows.

$$E = (I - SR^{-T}S^{T}A)(I - PA_{c}^{-1}P^{T}A)(I - SR^{-1}S^{T}A) = I - G^{-1}A.$$
 (1)

We now derive an estimate on the angle between the spaces range(S) and 85 range(P), which in our setting amounts to obtaining a bound on the energy norm 86 of Q, the  $\ell_2$ -orthogonal projection onto range(P). Let  $\gamma$  be the C.B.S. constant such 87 that it is the smallest number satisfying  $(Sw, Pv)_A \leq \gamma |Sw|_A |Pv|_A$ , then (cf. [6, Corol-88 lary 3.7]):

$$|Q|_A^2 = 1/(1-\gamma^2).$$

Using [2, Theorem 4.2] we can show that, if the symmetrized smoother  $\widetilde{R} = R + 90$   $R^T - S^T A S$  is positive definite, and  $(w^T \widetilde{R} w) / (w^T S^T A S w) \in [1, \kappa_s]$ , then

$$\frac{v^T G v}{v^T A v} \in [1, |Q|_A^2 (\kappa_s + c_c - 1)].$$

If a two-level method using a certain matching is already given, then both  $|Q|_A$  92 and  $\kappa_s$  can be estimated using the properties of the underlying graph. The norm  $|Q|_A$  93 is estimated as follows: 94

$$u^{T}QAQu = \sum_{(i,j) \in \mathscr{E}} ((Qu)_{i} - (Qu)_{j})^{2} \le 2d \sum_{(i,j) \in E} (u_{i} - u_{j})^{2} \le (2d)u^{T}Au$$

where d is the maximum degree of the graph. This implies that  $|Q|_A^2 \le 2d$ . Assuming 95 that the matching  $\mathcal{M}$  is perfect, we show that the smallest eigenvalue of  $S^TAS$  is 96 larger or equal to 4, by computing 97

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$$w^T S^T A S w \ge \sum_{(i,j) \in \mathcal{M}} ((Sw)_i - (Sw)_j)^2 = \sum_{(i,j) \in \mathcal{M}} 4(Sw)_i^2 = 4 \|w\|_{\ell_2}^2.$$

According to the Gershgorin theorem, the largest eigenvalue of  $S^TAS$  is bounded 98 by a function of d and for a simple smoother R, such as Richardson iteration,  $\kappa_s$  is 99 also bounded by a function of d. From the above results (i.e, the stability estimate 100 of Q in the A-seminorm and the lower bound on the smallest eigenvalue of  $S^{T}AS$ ) it 101 follows that the two-level method is uniformly convergent with respect to the size of 102 the matrix A. Based on the two-level convergence estimate, AMLI cycles with low 103 complexity and predictable convergence is then constructed.

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# 5 Algebraic Multilevel Iterations

An estimate of the two-level convergence rate does not automatically carry over to an 106 estimate of the convergence of a multilevel V-cycle, and in general, for piece-wise 107 constant coarse spaces, it can be shown that the convergence rate degrades expo- 108 nentially with respect to the number of levels. A remedy for this issue is to use more 109 complicated cycles such as AMLI, and keep a balance between complexity of a cycle 110 and its convergence rate so that the resulting algorithm is optimal or nearly optimal.

We describe an AMLI method by first rewriting the two-level preconditioner G, 112 as well as G which is G under the hierarchical basis (S, P), in block form: 113

$$\widehat{G}^{-1} = \widehat{L}^{-T} \begin{pmatrix} (R + R^T - S^T A S)^{-1} & 0 \\ 0 & A_c^{-1} \end{pmatrix} \widehat{L}^{-1},$$

$$G = (S, P)^{-1} \widehat{G}(S, P)^{-T},$$

where

$$\widehat{L} = \begin{pmatrix} I & 0 \\ P^T A S R^{-1} & I \end{pmatrix}.$$

Then define an AMLI preconditioner B as follows

$$\widehat{B}^{-1} = \widehat{L}^{-T} \begin{pmatrix} (R + R^T - S^T A S)^{-1} & 0 \\ 0 & B_c^{-1} q (A_c B_c^{-1}) \end{pmatrix} \widehat{L}^{-1},$$

$$B^{-1} = (S, P)^T \widehat{B}^{-1} (S, P).$$

Here  $A_c$  is the scaled unweighted graph Laplacian of the coarse graph and  $B_c$  is a 116 preconditioner of  $A_c$ , and q(t) is a polynomial. When q(t) = 1, the action  $\widehat{B}^{-1}$  stands for a V-cycle with an inexact solver  $B_c^{-1}$  on the coarse level. In the case of a W-cycle, we have q(t) = 2 - t. 119

The following lemma shows how well the AMLI preconditioner B approximates 120 the two-level preconditioner G.

**Lemma 1.** If 
$$\lambda_1 \leq \lambda(B_c^{-1}A_c) \leq \lambda_2$$
 and  $tq(t) > 0$  for  $t \in [\lambda_1, \lambda_2]$ , then

$$\min(1, \min_{\lambda_1 \leq t \leq \lambda_2} \frac{1}{tq(t)}) \leq \frac{v^T G^{-1} v}{v^T B^{-1} v} \leq \max(1, \max_{\lambda_1 < t < \lambda_2} \frac{1}{tq(t)}).$$

This lemma suggests that, the AMLI method is spectrally equivalent to a two- 123 level method, given that the coarse-level preconditioner is spectrally equivalent to 124 the coarser-level matrix. The upper and lower bounds in the lemma above are related 125 to estimates on |tq(t)| for t in a given interval. As shown in [1, 6], using higher 126 order polynomials q(t), the matrix  $B^{-1}$  can approximate  $G^{-1}$  arbitrarily well and 127 thus we will have a method with excellent convergence rate. However, a higher order 128 polynomial q(t) leads to a much more expensive computation of the coarser level 129 correction, and the resulting multilevel methods can have a very high complexity 130 and one should be careful in the choice of the polynomial degree.

Assume that a multilevel hierarchy is formed by a recursive application of the 132 matching algorithm. Denote the graph Laplacians on each level, and the correspond- 133 ing two-level preconditioners by  $A_k$  and  $G_k$ . Following the ordering of levels in [1, 6] 134 we set  $A = A_0$  and denote by  $A_J$  the coarsest matrix. Define a sequence of solvers as

$$\begin{split} \widehat{B}_J^{-1} &= \widehat{A}_J^{\dagger} = (S_J, P_J)^{-T} A_J^{\dagger} (S_J, P_J)^{-1}, \\ B_k^{-1} &= (S_k, P_k)^T \widehat{B}_k^{-1} (S_k, P_k), \quad k = 0, \dots, J, \\ \widehat{B}_k^{-1} &= \widehat{L}_k^{-T} \begin{pmatrix} (R_k + R_k^T - S_k^T A_k S_k)^{-1} & 0 \\ 0 & B_{k+1}^{-1} q(A_{k+1} B_{k+1}^{-1}) \end{pmatrix} L_k^{-1}, \quad k = 0 \dots J - 1. \end{split}$$

Then, a multilevel proof of convergence follows.

**Lemma 2.** Assume that there is a constant  $c_g$ ,  $1 \le c_g < 4$ , such that the following relation holds. 138

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$$v^T \widehat{A}_k v \le v^T \widehat{G}_k v \le c_g v^T \widehat{A}_k v, \quad \forall v \text{ and } k = 0, \dots, J.$$

Then there exists a linear function q(t), such that

$$\frac{2}{\sqrt{c_g}} - 1 \le \frac{v^T B_k^{-1} v}{v^T A_k^{-1} v} \le 1, \quad \forall v \text{ and } k = 0, \dots, J.$$

Here q(t) is a scaled and shifted Chebyshev type polynomial (see [1]).

This lemma shows that, if  $c_g$  is strictly less than 4, then the action  $B_0^{-1}$  is an 141 uniformly convergent AMLI cycle with  $O(n \log n)$  complexity. Even if  $c_g = 4$  on all levels, one may prove that the condition number of  $B_I^{-1}A_I$  for the case of second 143 order q(t) (similar to a W-cycle) grows linearly with respect to the number of levels  $J = \log n$ . This results in a convergence factor  $1 - 1/\log n$  at a complexity of 145  $O(n \log n)$  for each cycle.

The two-level method we suggest is based on graph matching, thus  $c_g \le |Q|_A^2 (\kappa_s + 147)$  $c_c-1$ ). In a simple case where the graph  ${\mathscr G}$  is a two-dimensional uniform grid, an 148 aligned regular matching yields  $|Q|_A^2 \leq 2$ ,  $\kappa_s = 1 + \varepsilon$  for arbitrary small  $\varepsilon$ , and  $c_c \leq 2$ . 149

This yields  $c_g \leq 4$  and thus the W-cycle AMLI preconditioner will result in 150 a nearly optimal order method (cf. Lemma 2 and the discussion below). For un- 151 structured or higher dimensional grids, numerical experiments indicate that random 152 matching may still result in two-level methods for which  $c_g \leq 4$ .

#### 6 Numerical Results

We use the matching based AMLI method to solve a family of unweighted graph 155 Laplacians, corresponding to graphs that represent structured grids or unstructured 156 triangulations.

Structured grids. In the structured grid case on a rectangular domain, we match in a fixed direction. After several levels of matching the graph corresponding to the 159 coarsest grid is a line. For the test on L-shaped domain, we still use matching in a 160 fixed direction until a part of the coarsest graph becomes a tree. In such case, the 161 unknowns can be ordered so that the fill-in during LU factorization on the coarsest 162 grid is small.

A similar strategy can be used for graph Laplacians corresponding to three- 164 dimensional structured grids. The matching procedure is applied only in two fixed 165 directions.

Convergence analysis indicates that, choosing as a smoother  $R^{-1} = (S^T A S)^{-1}$ guarantees the bound  $c_g \le 4$ , for a matching based two-level method on structured 168 grids. In the numerical experiments, we instead use a Gauss-Seidel smoother for all 169 structured grid problems. Using such a smoother retains a convergence rate  $\sim (1 - 170)$  $1/\log n$ ) and  $O(n\log n)$  computational complexity.

Unstructured grids. Each of the unstructured grids in our tests are constructed 172 by first perturbing the coordinates of vertices of a structured grid, followed by De- 173 launay triangulation of the resulting set of vertices. For unstructured grids, we use 174 a random matching algorithm. Numerical results show that the maximum degree of 175 the coarser graphs grow only during the first few coarsening steps. Hence, smoothers 176 such as Gauss-Seidel can approximate well  $(S_k^T A_k S_k)^{-1}$  on all levels and the application of such a smoother has a complexity proportional to the number of de- 178 grees of freedom (DOF) on level k. We use the CG method to perform the action 179 of  $(S_k^T A_k S_k)^{-1}$  on a vector. Such approach is practical since  $S^T A S$  is equally well 180 conditioned on all levels.

Instead of using the same AMLI polynomial q(t) on all levels, we determine the 182 polynomials  $q_k(t)$  on each level recursively, starting from the second coarsest level. 183 After constructing a multilevel hierarchy, we use 6 AMLI two level cycles (level 184 (J-1) and level J) and a Lanczos algorithm to estimate the condition number of 185  $B_{I-1}^{-1}A_{J-1}$ . We apply this procedure recursively (and with 6 AMLI multilevel cycles 186 from level (k+1) to J) to estimate the condition number of  $B_k^{-1}A_k$  on level k, for  $k = 1, \dots, J - 2$ . When all polynomials are determined, they are used in the AMLI 188 cycle during the solving phase.

Numerical tests. We use the AMLI cycle as a preconditioner of Conjugate Gra- 190 dient (CG) method. We stop the iterations when the relative residual becomes smaller 191 than  $10^{-10}$ . The results are summarized in Table 1. The number of CG iterations is 192 denoted by M, and the average convergence rate of the last five iterations is denoted by  $r_a$ . The CG coefficients are also used to estimate the condition number  $\kappa(B_0^{-1}A_0)$ , 194 as suggested in [4]. The operator and grid complexities are less than 2 in all the 195 examples presented below.

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(a)	$^{2}$ D	unit	SQ	uare

DOF	К	$r_a$	М
$256^{2}$			
512 <sup>2</sup>			
$1024^{2}$	32.9	0.69	40

#### (b) 3D unit cube

DOF	К	$r_a$	M
		0.36	
$64^{3}$	11.4	0.45	25
$128^{3}$	19.2	0.51	29

(c) 2D L-Shaped

DOF	κ	$r_a$	M
$(3/4) \cdot 256^2$	17.8	0.56	33
$(3/4) \cdot 512^2$			
$(3/4) \cdot 1024^2$	31.7	0.69	38

#### (d) 3D Fichera

DOF	К	$r_a$	M
$(7/8) \cdot 32^3$	7.5	0.40	22
$(7/8) \cdot 64^3$	11.1	0.48	25
$(7/8) \cdot 128^3$	15.8	0.55	29

(e) 2D unit square (ug)

DOF	к	$r_a$	M
$256^2$	31.4	0.58	35
512 <sup>2</sup>	36.7	0.63	39
1024 <sup>2</sup>	42.0	0.58	41

(f) 3D unit cube (ug)

DOF	κ	$r_a$	M
$32^{3}$	29.5	0.51	35
64 <sup>3</sup>	37.6	0.68	46
$128^{3}$	48.3	0.72	52

Table 1. Results for structured grids on square, cubic, L-shaped and Fichera domain, and for unstructured grids (ug) on square and cubic domain. Here,  $\kappa$  is an estimate (from CG) of  $\kappa(B_0^{-1}A_0).$ 

Note that for the 2D and 3D unstructured grid problems, the number of levels 197 for a given unstructured grid is the same as that of a structured grid with the same 198 degrees of freedom. We observe a logarithmic growth of the condition numbers with respect to the size of the grids, and fast convergence rates of the preconditioned CG 200 method in all cases.

7 Conclusions 202

We present an AMLI (AMG) method based on graph matching with a nearly optimal 203 convergence rate and computational complexity. We have also presented numerical 204 tests which confirming our estimates. Our ongoing research is on extending the es- 205 timates to general aggregation algorithms and aggregates configurations and we are 206 also investigating improvements of the AMLI method components.

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# Shifted Laplacian RAS Solvers for the Helmholtz Equation

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1 Introduction 10

We consider the Helmholtz equation:

$$-\Delta u^* - k^2 u^* = f \quad \text{in} \quad \Omega$$

$$u^* = g_D \text{ on } \partial \Omega_D, \quad \frac{\partial u^*}{\partial n} = g_N \text{ on } \partial \Omega_N, \quad \frac{\partial u^*}{\partial n} + iku^* = g_S \text{ on } \partial \Omega_S$$
(1)

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where  $\Omega$  is a bounded polygonal region in  $\Re^2$ , and the  $\partial\Omega_D$ ,  $\partial\Omega_N$  and  $\partial\Omega_S$  correspond to subsets of  $\partial\Omega$  where the Dirichlet, Neumann and Sommerfeld boundary conditions are imposed.

The main purpose of this paper is to introduce novel two-level overlapping 15 Schwarz methods for solving the Helmholtz equation. Among the most effective parallel two-level domain decomposition solvers for the Helmholtz equation on general 17 unstructured meshes, we mention the FETI-H method introduced by Farhat et al. [5], 18 and the WRAS-H-RC method introduced by Kimn and Sarkis [10]. FETI-H type pre- 19 conditioners belong to the class of nonoverlapping domain decomposition methods. 20 FETI-H methods can be viewed as a modification of the original FETI method in- 21 troduced by Farhat et al. [6]. The local solvers in FETI-H are based on Sommerfeld 22 boundary conditions, see [3], while the coarse problem is based on plane waves. 23 WRAS-H-RC type preconditioners belong to the class of overlapping Schwarz 24 methods. They can be viewed as a miscellaneous of several methods to enhance the 25 effectiveness of the solver for Helmholtz problems. The first ingredient of WRAS- 26 H-RC preconditioners is the use of Sommerfeld boundary conditions for the local 27 solvers on overlapping subdomains. This idea is similar to what was done in FETI- 28 H, however, now for the overlapping case. This idea can be found for instance in the 29 work of Cai et al. [2] and Kimn [8]. The second ingredient is the use of the Weighted 30 Restricted Additive Schwarz (WRAS) method introduced by Cai and Sarkis [1] in 31 order to average the local overlapping solutions. The third ingredient is the use of 32

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partition of unity coarse spaces, see [13]. Here we consider the multiplication of a 33 partition of unity times plane waves; see [12]. The fourth ingredient is how to define 34 the coarse problem. It was discovered in [10] that a dramatic gain in performance 35 can be obtained if WRAS techniques are applied to the fine-to-coarse restriction operator and the coarse-to-fine prolongation operator. The idea is to force the to act 37 more locally on the fine-to-coarse transference of information and globally on the 38 coarse-to-fine phase. The last ingredient is to put all these pieces together. The idea 39 is to extend the Balancing Domain Decomposition (BDD) methods of Mandel [11], 40 which were originally developed for the nonoverlapping case, to the overlapping 41 case. This extension was introduced in [9] and the methods there were denoted by 42 Overlapping Balancing Domain Decomposition (OBDD) methods. The WRAS-H- 43 RC methods in [10] stand for "WRAS" for the local solvers, "H" for the FETI-H 44 ingredients included in the methods, and "RC" for the restricted flavor of coarse 45 problem.

Here in this paper we investigate numerically new techniques to improve further 47 the performance of the WRAS-H-RC. More precisely, the shifted Laplacian tech- 48 niques introduced in [7] and [4], are used to construct novel local solvers. We investigate how the various kinds of shifts affect the performance of the algorithms. As 50 a result, we discover novel preconditioners that are more effective than the existing 51 ones.

# 2 Discrete Formulation of the Problem

From a Green's formula, (1) can be reduced to: Find  $u^* - u_D^* \in H_D^1(\Omega)$  such that,

$$a(u^*, v) = \int_{\Omega} (\nabla u^* \cdot \nabla \bar{v} - k^2 u^* \bar{v}) \, dx + ik \int_{\partial \Omega_S} u^* \bar{v} \, ds$$

$$= \int_{\Omega} f \bar{v} \, dx + \int_{\partial \Omega_N} g_N \bar{v} \, ds + \int_{\partial \Omega_S} g_S \bar{v} = F(v), \ \forall v \in H_D^1(\Omega),$$
(2)

where  $u_D^*$  is an extension of  $g_D$  to  $H^1(\Omega)$ , and  $H^1_D(\Omega)$  is the space of  $H^1(\Omega)$  functions vanishing on  $\partial \Omega_D$ .

Let  $\mathscr{T}_h(\Omega)$  be a quasi-uniform triangulation of  $\Omega$  and let  $V \subset H^1_D(\Omega)$  be the 58 finite element space of continuous piecewise linear functions vanishing on  $\partial \Omega_D$ . We 59 assume that  $g_D$  on  $\partial \Omega_D$  is a piecewise linear continuous function on  $\mathcal{T}^h(\partial \Omega_D)$  and 60 we have eliminated  $g_D$  by a discrete trivial zero extension inside  $\Omega$ . We then obtain 61 a discrete problem of the following form: Find  $u \in V$  such that 62

$$a(u,v) = f(v), \ \forall \ v \in V. \tag{3}$$

Using the standard hat basis functions, (3) can be rewritten as a linear system of 63 equations of the form

$$Au = f. (4)$$

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# 3 Description of the WRAS-H-RC Methods

## 3.1 Partitioning and Subdomains

Given the triangulation  $\mathcal{T}^h(\Omega)$ , we assume that a domain partition by elements has 67 been applied and resulted in N nonoverlapping subdomains  $\Omega_i, i = 1, ...N$ , such that 68

$$\overline{\Omega} = \bigcup_{i=1}^N \overline{\Omega}_i$$
 and  $\Omega_i \cap \Omega_i = \emptyset$ , for  $j \neq i$ .

Let  $\delta$  be a nonnegative integer. Define  $\Omega_i^0 = \Omega_i$ . For  $\delta \geq 1$ , define the overlapping 70 subdomains  $\Omega_i^\delta$  as follows: let  $\Omega_i^1$  be the one-overlap element extension of  $\Omega_i^0$  by 71 including all the immediate neighboring elements  $\tau_h \in \mathscr{T}^h(\Omega)$  such that  $\overline{\tau}_h \cap \overline{\Omega}_i^0 \neq \emptyset$ . 72 Using this idea recursively, we can define a  $\delta$ -extension overlapping subdomains  $\Omega_i^\delta$  73

$$\Omega_i = \Omega_i^0 \subset \Omega_i^1 \subset \dots \subset \Omega_i^\delta \dots$$

## 3.2 Partition of the Unity

Let w be a nonnegative integer. For nodes x on  $\partial \Omega_i^0$  define  $\hat{v}_i^w(x)=1$ , for nodes x on 76  $\partial \Omega_i^1 \backslash \overline{\Omega}_i^0$  define  $\hat{v}_i^w(x)=1-1/(w+1)$ , for nodes x on  $\partial \Omega_i^2 \backslash \overline{\Omega}_i^1$  define  $\hat{v}_i^w(x)=1-77$  2/(w+1), and recursively until  $\hat{v}_i^w(x)=0$ . For nodes x in  $\overline{\Omega} \backslash \overline{\Omega}_i^w$  define  $\hat{v}_i^w(x)=0$ . 78 The partition of unity  $v_i^w$  is defined as

$$\vartheta_i^w = I_h(\frac{\hat{\vartheta}_i^w}{\sum_{j=1}^N \hat{\vartheta}_j^w}) \quad i = 1, \cdots, N,$$

where  $I_h$  is the nodal piecewise linear interpolant on  $\mathscr{T}^h(\overline{\Omega})$ . Note that the support 81 of  $\vartheta_i^w$  is  $\Omega_i^{w+1}$  and  $|\nabla \vartheta_i^w| \leq O((w+1)/h)$ . We define the weighting diagonal matrix 82  $D_i^w$  as equal to  $\vartheta_i^w(x)$  at the nodes x of  $\overline{\Omega}$ .

# 3.3 Local Problems 8

Let us denote by  $V_i^{\delta}$ ,  $i=1,\cdots,N$ , the local space of functions in  $H^1(\Omega_i^{\delta})$  which are 85 continuous piecewise linear and vanishes only on  $\partial \Omega_i^{\delta} \cap \partial \Omega_D$ . For each subdomain 86  $\Omega_i^{\delta}$ , let  $R_i^{\delta}: V \to V_i^{\delta}$  be the regular restriction operator on  $V_i^{\delta}$ , that is,  $v_i(x) = v(x)$  87 for nodes  $x \in \overline{\Omega}_i^{\delta}$ .

For the local solvers, we respect the original boundary condition and impose 90 Sommerfeld boundary condition on the interior boundaries  $\partial \Omega_i^{\delta} \backslash \partial \Omega$ . The associated local projections in matrix form are defined by 92

$$T_{i,WRAS-H}^{\delta} = (R_i^{\delta} D_i^{\delta})^T (\tilde{A}_i^{\delta})^{-1} R_i^{\delta} A \quad i = 1, \dots, N$$
 (5)

where  $\tilde{A}_{i}^{\delta}$  are the matrix form of

$$\tilde{a}_{i}^{\delta}(u_{i}, v_{i}) = \int_{\Omega^{\delta}} (\nabla u_{i} \cdot \nabla \overline{v}_{i} - k^{2} u_{i} \overline{v}_{i}) dx + ik \int_{\partial \Omega^{\delta} \setminus (\partial \Omega_{D} \cup \partial \Omega_{N})} u_{i} \overline{v}_{i} ds. \tag{6}$$

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3.4 Coarse Problem 94

Let c be a nonnegative integer. The coarse space  $V_0^{c,p} \in V$  is defined as the space 95 spanned by  $D_i^c Q_j^D$  for  $i=1,\ldots,N$  and  $j=1,\cdots,p$ . Here,  $Q_j:=e^{ik\eta_j^Tx}$ , where 96  $\eta_j=(\cos(\theta_j),\sin(\theta_j))$ , with  $\theta_j=(j-1)\times\frac{\pi}{p},j=1,\cdots,p$ , while  $Q_j^D(x):=Q_j(x)$  for 97 nodes  $x\in\overline{\Omega}\setminus\partial\Omega_D$  and  $Q_j^D(x):=0$  for nodes x on  $\partial\Omega_D$ . The coarse-to-fine prolongation matrix  $(E_0^{c,p})$  consists of columns  $D_i^\delta Q_j^D$ , while the fine-to-coarse restriction 99 matrix  $R_0^{\delta,p}$  consists of rows  $(R_i^\delta)^T R_i^\delta Q_j^D$ . The first coarse problem we consider in 100 this paper is given by

 $P_{0,RC}^{\delta,c,p} = E_0^{c,p} [R_0^{\delta,p} A E_0^{c,p}]^{-1} R_0^{\delta,p}.$  (7)

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## 3.5 Hybrid Preconditioners

The first preconditioner we consider is given by

$$T_{WRAS-H-RC}^{\delta,c,p} := P_{0,RC}^{\delta,c,p} + (I - P_{0,RC}^{\delta,c,p})(\sum_{i=1}^{N} T_{i,WRAS-H}^{\delta})(I - P_{0,RC}^{\delta,c,p}). \tag{8}$$

Because  $P_{0,RC}^{\delta,c,p}$  is a projection, only one coarse problem solver is necessary per iteration of the iterative method.

Other hybrid preconditioners can also be designed. For instance, we can replace the local problem  $T_{i,WRAS}^{\delta}$  by

$$P_{i,OBDD-H}^{\delta} := (R_i^{\delta} D_i^{\delta})^T (\tilde{A}_i^{\delta})^{-1} R_i^{\delta} D_i^{\delta} A$$
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or/and replace the coarse problem  $P_{0,RC}^{\delta,c,p}$  by something more classical such as

$$P_0^{c,p} = E_0^{c,p} [(E_0^{c,p})^T A E_0^{c,p}]^{-1} (E_0^{c,p})^T.$$
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Inserting these operators properly into (7) we obtain preconditioners which we denote by  $T_{WRAS-H}^{\delta,c,p}$ ,  $T_{OBDD-H}^{\delta,c,p}$  or  $T_{OBDD-H-RC}^{\delta,c,p}$ . An interesting structure that 113  $T_{WRAS-H-RC}^{\delta,e,p}$  has, and the others do not, is that the same restriction operators  $R_i^{\delta}$  are 114 used to compute the right-hand side for both the local and coarse problems, therefore, 115 computational efficiency can be explored.

# **4 Shifted Local Operators**

The matrix  $\tilde{A}_i^{\delta}$  obtained from the bilinear form (6) can be written as

$$\tilde{A}_i^{\delta} = A_i^{\delta} - k^2 M_i^{\delta} + ik B_i^{\delta},$$
 119

where  $A_i^{\delta}$ ,  $M_i^{\delta}$ , and  $B_i^{\delta}$  are the corresponding matrices associated to

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$$\int_{\Omega_{i}^{\delta}} \nabla u_{i} \cdot \nabla \overline{v}_{i} dx + ik \int_{\partial \Omega_{i}^{\delta} \cap \partial \Omega_{S}} u_{i} \overline{v}_{i} ds, \quad \int_{\Omega_{i}^{\delta}} u_{i} \overline{v}_{i} dx \quad \text{and} \quad \int_{\partial \Omega_{i}^{\delta} \setminus \partial \Omega} u_{i} \overline{v}_{i} ds,$$

respectively. We note that the local matrix  $A_i^{\delta} - k^2 M_i^{\delta}$  is singular if  $k^2$  is a generalized 122 eigenvalue of  $A_i^{\delta}$ . Alternatively, if we enforce zero Dirichlet boundary condition on 123 the interior boundaries  $\partial \Omega_i \cap \Omega_i^{\delta}$ , singularities also might occurs, specially when the subdomains are not small enough. The Sommerfeld term plays the rule of shifting 125 the real spectrum of  $A_i^{\delta} - k^2 M_i^{\delta}$  to the upper part of the complex plane, therefore, 126 elliminating possible zero eigenvalues. More general shifts were introduced recently 127 by Gijzen et al. [7] and Erlangga et al. [4] to move the spectrum to a disk on the first 128 quadrant. Inspired by this work, we now consider shifts to define the local solvers as 129

$$\tilde{A}_{i}^{\delta}(\alpha_{r}, \alpha_{i}, \beta_{r}, \beta_{i}) = A_{i}^{\delta} + (\alpha_{r} + i\alpha_{i})k^{2}M_{i}^{\delta} + (\beta_{r} + i\beta_{i})kB_{i}^{\delta}, \tag{9}$$

that is, the local Laplacians  $A_i^{\delta}$  are shifted by a complex combination of  $M_i^{\delta}$  and  $B_i^{\delta}$ . 130 Note that  $\tilde{A}_i^{\delta}(-1,0,0,1)$  reduces to the original local solver (6), while  $\tilde{A}_i^{\delta}(-1,0,0,0)$ to  $A_i^{\delta} - k^2 M_i^{\delta}$ .

## **5 Numerical Results**

As a numerical test, we consider a wave guided problem for solving the Helmholtz 134 equation on the unit square. We consider homogeneous Neumann boundary condi- 135 tion on the horizontal sides, homogeneous Sommerfeld on the right vertical side, and 136 a constant identical to one Dirichlet on the left vertical side. The stopping criteria for 137 the PGMRES is to reduce the initial residual by a factor of  $10^{-6}$ . In all tests the right 138 preconditioner is applied.

The triangulation is composed of Courant elements of mesh size h = 1/256. The 141 nonoverlapping subdomains  $\Omega_i^0$  are squares of size 1/M, and the number of subdomains is denoted by  $nsub = M \times M$ . The pair  $(\delta, c)$  refers to how many layers of 143 elements are used to define the extension of the overlapping subdomains  $\Omega_i^{\delta}$  and the 144 extension of the support of the coarse basis functions, respectively. The constant k 145 refers to the wave number and p denotes the number of local plane waves used in 146 the coarse space. Table 1 shows that the method  $P_{WRAS-H-RC}$  is the most effective 147 method among those introduced in Sect. 3.5. Table 2 shows that we should select 148 the support for the coarse basis functions larger enough, larger than the size of the 149 extended subdomains. Tables 1 and 2 show that the number of iterations decreases 150 when we increase the size of the overlap.

We now test the effectiveness of  $P_{WRAS-H-RC}$  for several combinations of local 153 solvers  $\tilde{A}_i^{\delta}(\alpha_r,\alpha_i,\beta_r,\beta_i)$ . Table 3 shows results for  $\delta=2$  and Table 4 for  $\delta=0$ . 154 We can see from Tables 3 and 4 that the number of iterations using the original 155 local problem are 13 and 34, respectively. It is very surprising and interesting to observe that the number of iterations are 9 and 18 for the combination (0,1,1,0), a 157 respectable gain in efficiency. Tables 3 and 4 reveal that there exist more effective 158 choices for local solvers rather than the common choice approach of adding a Som- 159 merfeld term on the interior boundary of the subdomains. These preliminary results 160 are very inspiring and encouraging for further numerical and theoretical investiga- 161 tions.

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Table 1. The Guided Wave Problem, Sommerfeld boundary condition on interior subdomain boundaries, n = 257,  $nsub = 64(8 \times 8)$ , Tol= $10^{-6}$ , k = 20

$(\delta,c,p)$	(0,7,4)	(1,7,4)	(2,7,4)
OBDD-H	158	85	43
WRAS-H	150	74	36
OBDD - H - RC	40	23	16
WRAS - H - RC	34	19	13

Table 2. WRAS-H-RC The Guided Wave Problem, Sommerfeld boundary condition on interior subdomain boundaries, n = 257,  $nsub = 64(8 \times 8)$ , p = 4, Tol= $10^{-6}$ , k = 20

		WRAS-H-RC									
c=	1	2	3	4	5	6	7	8			
$\delta = 0$	78	67	54	46	40	37	34	32			
$\delta = 1$	190	36	31	25	22	21	19	18			
$\delta = 2$	181	181	19	18	16	14	13	12			

Table 3. The Guided Wave Problem, WRAS-H-RC algorithm with Shifted Laplacian local problems, n = 257, nsub = 64, Tol= $10^{-6}$ , p = 4, k = 20, c = 7,  $\delta = 2$ 

	$\alpha_r =$	-1	-1	-1	0	0	0	1	1	1
	$\alpha_i =$	-1	0	1	-1	0	1	-1	0	1
$\beta_r = -1$	$\beta_i = -1$	37	53	116	22	28	210	17	22	48
$\beta_r = -1$	$\beta_i = 0$	236	123	199	154	275	139	105	300*	138
$\beta_r = -1$	$\beta_i = 1$	66	34	28	227	24	16	55	22	17
$\beta_r = 0$	$\beta_i = -1$	20	23	62	14	14	20	12	11	12
$\beta_r = 0$	$\beta_i = 0$	19	16	13	17	300*	12	14	13	10
$\beta_r = 0$	$\beta_i = 1$	55	13	13	23	13	11	15	12	11
$\beta_r = 1$	$\beta_i = -1$	15	12	12	13	10	10	12	10	9
$\beta_r = 1$	$\beta_i = 0$	13	17	11	12	10	9	12	10	8
$\beta_r = 1$	$\beta_i = 1$	17	10	11	12	10	9	11	10	9

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t4.1 t4.2 t4 3 t4.4 t4 5

t4 7 t4.8 t4 9 t4.10 t4 11

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Table 4. The Guided Wave Problem, WRAS-H-RC algorithm with Shifted Laplacian local problems, n = 257, nsub = 64, Tol= $10^{-6}$ , p = 4, k = 20, c = 7,  $\delta = 0$ 

	$\alpha_r =$	-1	-1	-1	0	0	0	1	1	1
	$\alpha_i =$	-1	0	1	-1	0	1	-1	0	1
$\beta_r = -1$	$\beta_i = -1$	168	213	300*	99	168	300*	69	106	300*
$\beta_r = -1$	$\beta_i = 0$	291	207	243	238	300*	209	221	300*	300*
$\beta_r = -1$	$\beta_i = 1$	300*	137	101	300*	130	63	300*	107	67
$\beta_r = 0$	$\beta_i = -1$	55	69	289	38	42	80	34	30	32
$\beta_r = 0$	$\beta_i = 0$	45	31	30	38	300*	27	34	24	24
$\beta_r = 0$	$\beta_i = 1$	279	34	33	94	39	30	40	35	31
$\beta_r = 1$	$\beta_i = -1$	34	31	39	29	25	22	27	24	21
$\beta_r = 1$	$\beta_i = 0$	27	22	21	24	20	18	24	21	20
$\beta_r = 1$	$\beta_i = 1$	51	23	21	25	21	20	23	21	21

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# A Subspace Correction Method for Nearly Singular **Linear Elasticity Problems**

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1 Introduction 9

The focus of this work is on constructing a robust (uniform in the problem param- 10 eters) iterative solution method for the system of linear algebraic equations arising 11 from a nonconforming finite element discretization based on reduced integration. We 12 introduce a specific space decomposition into two overlapping subspaces that serves 13 as a basis for devising a uniformly convergent subspace correction algorithm. We 14 consider the equations of linear elasticity in primal variables. For nearly incompressible materials, i.e., when the Poisson ratio  $\nu$  approaches 1/2, this problem becomes 16 ill-posed and the resulting discrete problem is nearly singular.

Subspace correction methods for nearly singular systems have been studied 18 in [10] leading to robust multigrid methods for planar linear elasticity problems 19 (see [11]). In [13] a multigrid method has been presented for a finite element dis- 20 cretization with  $P_2 - P_0$  elements. This approach relies on a local basis for the weakly 21 divergence-free functions.

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In this setting, presently known (multilevel) iterative solution methods are opti- 23 mal or nearly optimal for the pure displacement problem only, i.e., when Dirichlet 24 boundary conditions are imposed on the entire boundary, see, e.g., [1, 4]. For pure 25 traction or mixed boundary conditions the problem gets more involved. It is known, 26 that standard (conforming and nonconforming) finite element methods then require 27 certain stabilization techniques, see, e.g., [3, 6]. We employ a discretization scheme 28 introduced in [3] which achieves the stabilization via reduced integration. Note that 29 based on an appropriate discrete version of Korn's second inequality optimal error 30 estimates have been shown for this method (see [3]).

The remainder of this paper is organized as follows: The formulation of the linear elasticity problem with pure traction boundary conditions and its finite element 33 discretization are given in Sect. 2. We briefly recall some convergence results for the 34 Method of Successive Subspace Correction (MSSC) in Sect. 3. In Sect. 4 we present 35 a specific space decomposition which defines an MSSC preconditioner. Finally, we 36

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present a numerical test illustrating the optimal performance of the preconditioner in 37 Sect. 5.

#### 2 Problem Formulation

For the sake of simplicity we consider only two-dimensional problems in this paper. 40 Let  $\Omega$  be a bounded, connected and open subset of  $\mathbb{R}^2$ , denoting the reference con-41 figuration of an elastic body. The boundary of  $\Omega$  is denoted by  $\partial \Omega$ . Following [3] 42 we consider the pure traction problem of linear elasticity which reads 43

$$\boldsymbol{\sigma} = \mu \left[ \boldsymbol{\varepsilon}(\mathbf{u}) + \frac{v}{1 - 2v} \operatorname{div} \mathbf{u} I \right] \qquad \text{in } \Omega, \tag{1a}$$
$$-\operatorname{div} \boldsymbol{\sigma} = \mathbf{f} \qquad \text{in } \Omega, \tag{1b}$$
$$\boldsymbol{\sigma} \cdot \mathbf{n} = \mathbf{g} \qquad \text{on } \partial \Omega \tag{1c}$$

$$-\operatorname{div}\boldsymbol{\sigma} = \mathbf{f} \qquad \text{in } \Omega, \tag{1b}$$

$$\mathbf{\sigma} \cdot \mathbf{n} = \mathbf{g} \qquad \text{on } \partial \Omega. \tag{1c}$$

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where  $\sigma$  denotes the stress tensor and  $\boldsymbol{\varepsilon}(\mathbf{u}) := \nabla^{(s)} \mathbf{u}$  is the symmetric gradient, i.e., 44  $\varepsilon_{ij}(\mathbf{u})$ : =  $\frac{1}{2} \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right)$ . Further  $\mathbf{u}$  denotes the vector of displacements,  $\mathbf{f}$  denotes 45 the body forces, **n** is the outwards pointing unit normal vector on  $\Gamma = \partial \Omega$  and **g** is 46 the applied load on  $\Gamma$ . The properties of the material depend on the Poisson ratio 47  $v \in [0, 1/2)$ , and the shear modulus  $\mu := \frac{E}{1+v}$  where E is the modulus of elasticity.

We consider the space  $\mathbf{V}^{\text{RBM}}$ :=  $\{\mathbf{v}: \mathbf{v} = (a_1 + by, a_2 - bx)^t, a_1, a_2, b \in \mathbb{R}\}$  of 49 rigid body motions and define the subspace  $\hat{\mathbf{V}}$  of  $H^1$ -functions orthogonal to  $\mathbf{V}^{\text{RBM}}$ , 50 i.e.,

$$\hat{\mathbf{V}} := \{ \mathbf{v} \in [H^1(\Omega)]^2 : \int_{\Omega} \mathbf{v} \, d\mathbf{x} = \mathbf{0} \quad \text{and} \quad \int_{\Omega} v_1 y - v_2 x \, d\mathbf{x} = 0 \}.$$
 (2)

Let  $\mathcal{T}_H$  be a quasi-uniform triangulation of  $\Omega$ . Moreover, we subdivide each triangle 52  $T \in \mathcal{T}_H$  into four congruent triangles by adding the midpoints of the edges to the set 53 of vertices. The obtained refined triangulation  $\mathcal{T}_h$  of  $\Omega$  has a mesh size h=H/2. 54 We introduce the vector space  $\mathbf{V} := [V]^2 := [H^1(\Omega)]^2$  and the subspace  $\mathbf{V}_h := [V_h]^2$ , 55 which consists of the vector-valued continuous piecewise linear functions on the fine 56 mesh  $\mathcal{J}_h$ . Next we define  $\hat{\mathbf{V}}_h := \mathbf{V}_h \cap \hat{\mathbf{V}}$  and denote the space of piecewise constant 57 functions on  $\mathcal{T}_H$  by  $S_H$ . Then we consider the problem: Find  $\mathbf{u}_h \in \hat{\mathbf{V}}_h$  such that

$$a(\mathbf{u}_h, \mathbf{v}_h) = L(\mathbf{v}_h) := (\mathbf{f}, \mathbf{v}_h)_0 + \int_{\partial \Omega} \mathbf{g} \cdot \mathbf{v}_h \, ds \qquad \forall \mathbf{v}_h \in \hat{\mathbf{V}}_h,$$
 (3)

$$a(\mathbf{u}_h, \mathbf{v}_h) := \mu \left( (\boldsymbol{\varepsilon}(\mathbf{u}_h), \boldsymbol{\varepsilon}(\mathbf{v}_h))_0 + \frac{v}{1 - 2v} (P_0 \operatorname{div} \mathbf{u}_h, P_0 \operatorname{div} \mathbf{v}_h)_0 \right), \tag{4}$$

where  $\mathbf{f} \in [L_2(\Omega)]^2$  and  $\mathbf{g} \in [L_2(\partial \Omega)]^2$ .  $P_0$  is the  $L^2$ -projection onto  $S_H$ , that is,

$$P_0(v)|_{T_H} = \frac{1}{|T_H|} \int_{T_H} v \, \mathrm{d}\mathbf{x} \qquad \forall T_H \in \mathscr{T}_H, \tag{5}$$

for any scalar function  $v \in L^2(\Omega)$ . It is known that under the compatibility condition 60  $L(\mathbf{v}) = 0$  for all  $\mathbf{v} \in \mathbf{V}^{RBM}$  problem (3) has a unique solution  $\mathbf{u}_h \in \hat{\mathbf{V}}_h$ , see, e.g., [1]. 61 In [3] optimal order error estimates have been shown for this approximation, which 62 are robust with respect to the Poisson ratio  $\nu$ . 63

# 3 Subspace Correction Framework

The general framework of subspace correction methods is closely related to the abstract Schwarz theory, see, e.g., [5, 14].

Let us consider the variational problem: Find  $u \in V$  such that

$$a(u, v) = f(v) \quad \forall v \in V,$$
 (6)

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with  $V \subset H$  being a closed subset of the Hilbert space H. Moreover, we assume that 68 the bilinear form  $a(.,.): H \times H \to \mathbb{R}$  is continuous, symmetric, and H-elliptic. If f 69 is a continuous linear functional on H, then this problem is well-posed. 70

Now, let us split V into a-not necessarily direct-sum of closed subspaces  $V_i \subset 71$   $V, i = 1, \ldots, J$ , i.e.,  $V = \sum_{i=1}^J V_i$ . With each subspace  $V_i$  we associate a symmetric, 72 bounded, and elliptic bilinear form  $a_i(.,.)$  approximating a(.,.) on  $V_i$ . The MSSC 73 (see [16, Algorighm 2.1]) solves the residual equation for  $i = 1, \ldots, J$  with  $u_l = u^l$ : 74 Find  $e_i \in V_i$  such that for all  $v_i \in V_i$ , there holds:

$$a(e_i, v_i) = f(v_i) - a(u_{l+i-1}, v_i),$$
 and set  $u_{l+i} = u_{l+i-1} + e_i,$  (7)

Finally, the next iterate is  $u^{l+1} = u_{l+J}$ . Let  $T_i : V \to V_i$  be defined as

$$a_i(T_i v, v_i) = a(v, v_i)$$
, for all  $v_i \in V_i$ .

The assumptions on  $a_i(.,.)$  imply that  $T_i$  is well-defined,  $\mathcal{R}(T_i) = V_i$ , and  $T_i: V_i \to V_i$  77 is an isomorphism. The error after l iterations of the MSSC is given by  $u-u^l=E(u-78u^{l-1})=...=E^l(u-u^0)$ , where the error propagation operator E can be represented 79 in product form , i.e.,

$$E = (I - T_J)(I - T_{J-1}) \cdots (I - T_1). \tag{8}$$

In the following we consider the case of exact subspace solves, i.e.,  $a_i(.,.) = a(.,.)$  81 on  $V_i$ , in which  $T_i$  reduces to the idempotent, a-adjoint operator  $P_i$  defined by

$$a(P_i v, v_i) = a(v, v_i) \qquad \forall v_i \in V_i.$$
(9)

For a proof of the following identity for the energy norm of the error propagation 83 operator we refer the reader to [16].

**Theorem 1.** Under the assumptions (9) and  $V = \sum_{i=1}^{J} V_i$  we have

$$||E||_a^2 = ||(I - P_J)(I - P_{J-1})\cdots(I - P_1)||_a^2 = \frac{c_0}{1 + c_0}$$
(10)

where 
$$c_0 = \sup_{\|v\|_a = 1} \inf_{\sum_i v_i = v} \sum_{i=1}^J \|P_i \sum_{j=i+1}^J v_j\|_a^2 < \infty$$
.

Let  $\mathscr{E}_H$  be the set of edges of  $\mathscr{T}_H$  and  $\mathscr{V}_H$  be the set of (coarse) vertices of the mesh 87  $\mathscr{T}_H$ . Then for any vertex  $v_i \in \mathscr{V}_H$  we denote the set of edges sharing  $v_i$  by  $\mathscr{N}_i^{\mathscr{E}}$ . 88 For any edge  $E = (v_{E,1}, v_{E,2}) \in \mathscr{E}_H$  by  $\varphi_E$  we denote the scalar nodal basis function 89 corresponding to the midpoint of the edge E, and by  $\varphi_{E,1}$  and  $\varphi_{E,2}$  the nodal basis 90

functions corresponding to the vertices  $v_{E,1}$  and  $v_{E,2}$  of E. The corresponding vector-valued degrees of freedom (dof) of any function  $\mathbf{v}_h \in \mathbf{V}_h$  are denoted by  $\mathbf{v}_E$ ,  $\mathbf{v}_{E,1}$  and 92  $\mathbf{v}_{E,2}$ , respectively. We further use  $\varphi_i$  and  $\mathbf{v}_i$  to denote the basis functions and dof 93 associated with the vertices from  $\mathscr{V}_H$ .

For any edge  $E \in \mathcal{E}_H$  we assume that  $v_{E,1} < v_{E,2}$  and that the globally defined 95 tangential vector  $\tau_E$  points from  $v_{E,1}$  to  $v_{E,2}$ . The global edge normal vector  $\mathbf{n}_E$  is 96 orthogonal to  $\tau_E$  and is obtained from  $\tau_E$  by a clockwise rotation. By  $\mathbf{V}_H^{RT}$  we denote 97 the lowest order Raviart Thomas space (cf. [2]), i.e.,

$$\mathbf{V}_{H}^{RT} := \{ \mathbf{v} \in [L^{2}(\Omega)]^{2} : \mathbf{v} = \mathbf{a} + (bx, by)^{t} \text{ on each } T \in \mathcal{T}_{H}, \mathbf{a} \in \mathbb{R}^{2}, b \in \mathbb{R} \}$$
 (11)

where the degrees of freedom are the normal fluxes over the edges E, i.e.,  $F_E^{RT}(\mathbf{v})$ := 99  $\frac{1}{|E|}\int_E \mathbf{v}\cdot\mathbf{n}_E\,\mathrm{d}s$ . The basis functions  $\varphi_E^{RT}$  corresponding to an edge E of an element 100  $T\in\mathcal{T}_H$  are such that  $F_{E'}^{RT}(\varphi_E^{RT})$ :=  $\delta_{EE'}$ . We also use the projection  $\Pi^{RT}:\mathbf{V}\mapsto$  101  $\mathbf{V}_H^{RT}$  defined by  $\Pi^{RT}(\mathbf{v})=\sum_{E\in\mathscr{E}_H}F_E^{RT}(\mathbf{v})\varphi_E^{RT}$ , for which the commuting property 102  $P_0\,\mathrm{div}\,\mathbf{v}_h=\mathrm{div}\,\Pi^{RT}(\mathbf{v}_h)$  holds for any  $\mathbf{v}_h\in\mathbf{V}_h$  (cf. [2, p. 131]).

# **4 Space Decomposition**

Let us consider the following unique decomposition of any function  $\mathbf{v}_h \in \mathbf{V}_h$ :

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$$\mathbf{v}_{h} = \sum_{i \in \mathcal{V}_{H}} \boldsymbol{\varphi}_{i} \mathbf{v}_{i} + \sum_{E \in \mathcal{E}_{H}} \boldsymbol{\varphi}_{E} \mathbf{v}_{E}$$

$$= \underbrace{\sum_{i \in \mathcal{V}_{H}} \left[ \boldsymbol{\varphi}_{i} \mathbf{v}_{i} - \frac{1}{2} \sum_{E \in \mathcal{N}_{i}^{\mathcal{E}}} (\mathbf{v}_{i} \cdot \mathbf{n}_{E}) \boldsymbol{\varphi}_{E} \mathbf{n}_{E} \right]}_{=: \mathbf{v}_{\mathcal{V}}} + \underbrace{\sum_{E \in \mathcal{E}_{H}} \left( \left[ \mathbf{v}_{E} + \frac{1}{2} (\mathbf{v}_{E,1} + \mathbf{v}_{E,2}) \right] \cdot \mathbf{n}_{E} \right) \boldsymbol{\varphi}_{E} \mathbf{n}_{E}}_{=: \mathbf{v}_{\tau}}.$$

Next we define the splitting  $V_h = V_{\mathscr{V}} \oplus V_{\tau} \oplus V_n$ , where

$$\begin{aligned} \mathbf{V}_{\mathscr{V}} &:= \left\{ \mathbf{v}_h \in \mathbf{V}_h : \mathbf{v}_h = \sum_{i \in \mathscr{V}_H} \left[ \varphi_i \mathbf{v}_i - \frac{1}{2} \sum_{E \in \mathscr{N}_i^{\mathscr{E}}} (\mathbf{v}_i \cdot \mathbf{n}_E) \, \varphi_E \mathbf{n}_E \right] \right\}, \\ \mathbf{V}_{\tau} &:= \left\{ \mathbf{v}_h \in \mathbf{V}_h : \mathbf{v}_h = \sum_{E \in \mathscr{E}_H} \alpha_E \varphi_E \tau_E \right\}, \quad \mathbf{V}_n &:= \left\{ \mathbf{v}_h \in \mathbf{V}_h : \mathbf{v}_h = \sum_{E \in \mathscr{E}_H} \alpha_E \varphi_E \mathbf{n}_E \right\}. \end{aligned}$$

Note that  $\Pi^{RT}(\mathbf{V}_{\mathscr{V}}) = \Pi^{RT}(\mathbf{V}_{\tau}) = \{0\}$ . Next, we introduce the spaces

$$\begin{split} \boldsymbol{V}_{\text{curl}} &:= \left\{ \mathbf{v}_h \in \boldsymbol{V}_h : \mathbf{v}_h = \sum_{i \in \mathscr{V}_H} \beta_i \sum_{E \in \mathscr{N}_i^{\mathscr{E}}} \frac{\delta_{E,i}}{|E|} \varphi_E \mathbf{n}_E \right\}, \\ \boldsymbol{V}_{\nabla_h} &:= \left\{ \mathbf{v}_h \in \boldsymbol{V}_h : \mathbf{v}_h = \sum_{T \in \mathscr{T}_H} \gamma_T \sum_{E \subset T} \left( \mathbf{n}_E \cdot \mathbf{n}_{E,T} \right) \varphi_E \mathbf{n}_E \right\}. \end{split}$$

Here  $\delta_{E,i}$  is defined by

$$\delta_{E,i} = \begin{cases} -1 & \text{if } i = v_{E,1} \\ 1 & \text{if } i = v_{E,2} \end{cases}$$
 (12)

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Note that  $V_{\text{curl}} \subset V_n$ , and  $V_{\nabla_h} \subset V_n$ , and the following properties hold:

$$\begin{split} P_0 \operatorname{div}(\mathbf{v}_{\operatorname{curl}}) &= \operatorname{div} \Pi^{RT}(\mathbf{v}_{\operatorname{curl}}) = 0 \qquad \forall \mathbf{v}_{\operatorname{curl}} \in \boldsymbol{V}_{\operatorname{curl}}, \\ P_0 \operatorname{div}(\mathbf{v}_{\nabla_b}) &= \operatorname{div} \Pi^{RT}(\mathbf{v}_{\nabla_b}) \neq 0 \qquad \forall \mathbf{v}_{\nabla_b} \in \boldsymbol{V}_{\nabla_b}. \end{split}$$

Moreover,  $\dim(\mathbf{V}_{\text{curl}}) = n_{v,H} - 1$  and  $\dim(\mathbf{V}_{\nabla_h}) = n_{T,H}$ , and thus, using Euler's formula, i.e.,  $n_{v,H} - 1 + n_{T,H} = n_{E,H}$ , we find that  $\boldsymbol{V}_n = \boldsymbol{V}_{\text{curl}} \oplus \boldsymbol{V}_{\nabla_h}$ . Hence we obtain

$$\mathbf{V}_h = \mathbf{V}_{\mathcal{V}} \oplus \mathbf{V}_{\tau} \oplus \mathbf{V}_{\text{curl}} \oplus \mathbf{V}_{\nabla_h}. \tag{13}$$

Finally, we decompose  $V_h$  into two overlapping subspaces  $V_I$  and  $V_{II}$ :

$$\mathbf{V}_{I} = \mathbf{V}_{\mathscr{V}} \oplus \mathbf{V}_{\tau} \oplus \mathbf{V}_{\text{curl}} \tag{14}$$

$$V_{I} = V_{\mathscr{V}} \oplus V_{\tau} \oplus V_{\text{curl}}$$

$$V_{II} = V_{\tau} \oplus V_{\text{curl}} \oplus V_{\nabla_{h}}$$

$$\tag{14}$$

The overlap of  $m{V}_I$  and  $m{V}_{II}$  is given by  $m{V}_{ au} + m{V}_{ ext{curl}}$ , and any element  $m{v}_{II} \in m{V}_{II}$  can 113 be uniquely decomposed into  $\mathbf{v}_{II} = \mathbf{v}_{\tau} + \mathbf{v}_{\text{curl}} + \mathbf{v}_{\nabla_h}$ , with  $\mathbf{v}_{\tau} \in \mathbf{V}_{\tau}$ ,  $\mathbf{v}_{\text{curl}} \in \mathbf{V}_{\text{curl}}$  and  $\mathbf{v}_{\nabla_h} \in \mathbf{V}_{\nabla_h}$ . However, finding the components  $\mathbf{v}_{\mathrm{curl}} \stackrel{"}{\in} \mathbf{V}_{\mathrm{curl}}$  and  $\mathbf{v}_{\nabla_h} \in \mathbf{V}_{\nabla_h}$  for a given 115 function  $\mathbf{v}_n \in \mathbf{V}_n$  requires a solution of a system with an *M*-matrix corresponding to the lowest order mixed method for Laplace equation with lumped mass [2]. 117

Note that since  $P_0 \operatorname{div}(\mathbf{V}_I) = \operatorname{div} \Pi^{RT}(\mathbf{V}_I) = \{0\}$  the bilinear form a(.,.) satisfies

$$a(\mathbf{u}_I, \mathbf{v}_I) = \mu(\boldsymbol{\varepsilon}(\mathbf{u}_I), \boldsymbol{\varepsilon}(\mathbf{v}_I))_0 \qquad \forall \mathbf{u}_I, \mathbf{v}_I \in \boldsymbol{V}_I,$$
 (16)

and in the limit case v=0 we have  $a(\mathbf{u}_h,\mathbf{v}_h)=\mu(\boldsymbol{\varepsilon}(\mathbf{u}_h),\boldsymbol{\varepsilon}(\mathbf{v}_h))_0$  for all  $\mathbf{u}_h,\mathbf{v}_h\in \boldsymbol{V}_h$ . In the following, we use the operator representations  $A: V \to V$  and  $A_{\varepsilon}: V \to V$ for the bilinear forms a(.,.) and  $\mu(\boldsymbol{\varepsilon}(.),\boldsymbol{\varepsilon}(.))_0$ . If we symmetrize the MSSC, we 121 obtain the following error propagation  $\bar{E}_{MSSC}$ , compare with (8) in case of J=2 and exact subsolves, i.e., 123

$$\bar{E}_{MSSC} = (I - P_I)(I - P_{II})(I - P_I).$$

The error propagation operator can be rewritten as  $\bar{E}_{MSSC} = I - \bar{B}_{MSSC}A$ , with symmetric  $\bar{B}_{MSSC}$ . Further,  $\bar{B}_{MSSC}$  is positive definite, since  $\bar{E}_{MSSC}$  is non-expansive. Note that even though  $\bar{B}_{MSSC} = (I - \bar{E}_{MSSC})A^{-1}$  formally involves the inverse of A, we do not need  $A^{-1}$  in order to apply  $\bar{B}_{MSSC}$ . 127

If v is bounded away from the incompressible limit 1/2, we know that  $A_{\varepsilon}$  is 128 spectrally equivalent to A. Further, there are efficient preconditioners for  $A_{\mathcal{E}}$ . We now 129 define the additive preconditioner B by 130

$$B := \frac{1 - 2v}{1 - v} A_{\varepsilon}^{-1} + \frac{v}{1 - v} \bar{B}_{MSSC}. \tag{17}$$

Note that B is a convex combination of  $A_{\varepsilon}^{-1}$  and  $\bar{B}_{MSSC}$ .

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Remark 1. It has been shown in [14, 16] that an inexact solution of the subproblems (7) results in a uniform preconditioner under reasonable assumptions. The sub- 133 problems on the spaces  $V_I$  and  $V_h$  involve the bilinear form

$$\bar{a}(\mathbf{u}_i, \mathbf{v}_i) = \mu(\boldsymbol{\varepsilon}(\mathbf{u}_i), \boldsymbol{\varepsilon}(\mathbf{v}_i))_0 \qquad \forall \mathbf{u}_i, \mathbf{v}_i \in \boldsymbol{W} = \boldsymbol{V}_I, \boldsymbol{V}_h.$$
 (18)

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Any efficient preconditioning technique for the vector-Laplace equation can be employed in these steps, e.g., classical AMG (see [12]) or AMGm (see [8]). 136

The problem on  $V_H = V_E := \{ \mathbf{v}_h \in V_h : \mathbf{v}_h(\mathbf{x}_i) = \mathbf{0} \ v_i \in \mathcal{Y}_H \}$  is more involved. 137 First, by using Korn's inequality, Poincarè's inequality and the inverse inequality one 138 can show that 139

$$\|\boldsymbol{\varepsilon}(\mathbf{v}_E)\|_0^2 \approx \|\nabla \mathbf{v}_E\|_0^2 \approx H^{-2}\|\mathbf{v}_E\|_0^2$$
.

Second, any function  $\mathbf{v}_E \in \mathbf{V}_E$  can be uniquely decomposed into  $\mathbf{v}_E = \mathbf{v}_n + \mathbf{v}_{\tau}$  where  $\mathbf{v}_n \in \mathbf{V}_n$  and  $\mathbf{v}_\tau \in \mathbf{V}_\tau$ . Moreover, by locally estimating the angle between  $\mathbf{V}_n$  and  $\mathbf{V}_\tau$  141 in the  $a(\cdot, \cdot)$ -inner product, it can be shown that

$$\|\mathbf{v}_{E}\|_{0}^{2} = \|\mathbf{v}_{n} + \mathbf{v}_{\tau}\|_{0}^{2} \approx \|\mathbf{v}_{n}\|_{0}^{2} + \|\mathbf{v}_{\tau}\|_{0}^{2}$$
(19)

holds uniformly with respect to the mesh size h. Furthermore  $\Pi^{RT}(\mathbf{v}_{\tau}) = 0$  for all 143  $\mathbf{v}_{\tau} \in \mathbf{V}_{\tau}$ . Hence, the relation  $a(\mathbf{u}_{E}, \mathbf{v}_{E}) \approx \tilde{a}(\mathbf{u}_{E}, \mathbf{v}_{E})$  holds on  $\mathbf{V}_{II}$  where

$$\tilde{a}(\mathbf{u}_E, \mathbf{v}_E) := \mu \left\{ H^{-2}(\mathbf{u}_\tau, \mathbf{v}_\tau)_0 + H^{-2}(\mathbf{u}_n, \mathbf{v}_n)_0 + \frac{\nu}{1 - 2\nu} (P_0 \operatorname{div} \mathbf{u}_n, P_0 \operatorname{div} \mathbf{v}_n)_0 \right\}.$$
(20)

Now, using the interpolation operator  $I_{RT}^h: \boldsymbol{V}_H^{RT} \to \boldsymbol{V}_h$ , defined by  $I_{RT}^h(\varphi_E^{RT}) = 145$   $2\varphi_E \mathbf{n}_E \in \boldsymbol{V}_n$ , one can show that  $\boldsymbol{V}_n$  is isomorphic to  $\boldsymbol{V}_H^{RT}$ . Thus solving a variational 146 problem with  $\tilde{a}(.,.)$  on  $V_n$  is equivalent to solving a problem with the bilinear form 147

$$a_{RT}(\mathbf{u}_{RT}, \mathbf{v}_{RT}) := \mu \left\{ H^{-2}(\mathbf{u}_{RT}, \mathbf{v}_{RT})_0 + \frac{v}{1 - 2v} (\operatorname{div} \mathbf{u}_{RT}, \operatorname{div} \mathbf{v}_{RT})_0 \right\},$$
 (21)

on  $V_H^{RT}$  (see [7, 15]). An efficient solver for the latter problem can be designed by 148 using the auxiliary space preconditioner of [7], or by using the robust algebraic mul- 149 tilevel iteration method developed in [9]. 150

# 5 Numerical Experiment

We now perform a numerical test to show that the preconditioner (17) is an efficient and robust preconditioner. We consider the problem with homogenous Dirichlet 153 boundary conditions on the unit square  $\Omega = (0,1)^2$ . The number of PCG iterations 154 for a residual reduction by a factor  $10^8$  are shown in Table 1. The subproblems on 155  $V_I$  and  $V_{II}$  are solved exactly. Additionally, we list the estimated condition numbers 156  $\kappa(BA)$ , obtained from the Lanczos process. 157

t1 1 t1.2 t1 3 t1 4 t1.5 t1.6 t1.7 t1 8 t1 9 t1 10

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**Table 1.** Iteration numbers (#it.) and condition numbers ( $\kappa(BA)$ ) of the pcg-cycle.

#DOF	2	242	1	058	4	418	18	050	72	962	293	3378
	#it.	κ	#it.	κ	#it.	κ	#it.	κ	#it.	κ	#it.	κ
v = 0:	1	1.00	1	1.00	1	1.00	1	1.00	1	1.00	1	1.00
v = 0.25:	8	1.41	8	1.48	8	1.53	9	1.55	9	1.57	9	1.57
v = 0.4:	10	1.90	11	2.19	12	2.38	12	2.49	13	2.57	13	2.62
v = 0.45:	11	2.11	12	2.61	14	3.01	15	3.25	15	3.41	15	3.52
v = 0.49:	10	1.90	11	2.54	14	3.31	16	3.97	17	4.39	17	4.69
v = 0.499:	9	1.98	10	1.98	11	2.13	14	2.99	15	3.83	17	4.51
v = 0.4999:	9	1.99	9	1.99	9	1.99	10	1.99	12	2.43	13	3.34
v = 0.49999	:   9	1.99	9	1.99	9	2.00	9	2.00	9	2.00	10	2.00

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# Adaptive Finite Element Methods with Inexact Solvers 2 for the Nonlinear Poisson-Boltzmann Equation

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1 Introduction 13

In this article we study adaptive finite element methods (AFEM) with inexact solvers 14 for a class of semilinear elliptic interface problems. We are particularly interested in 15 nonlinear problems with discontinuous diffusion coefficients, such as the nonlinear 16 Poisson-Boltzmann equation and its regularizations. The algorithm we study consists of the standard SOLVE-ESTIMATE-MARK-REFINE procedure common to 18 many adaptive finite element algorithms, but where the SOLVE step involves only a 19 full solve on the coarsest level, and the remaining levels involve only single Newton 20 updates to the previous approximate solution. We summarize a recently developed 21 AFEM convergence theory for inexact solvers appearing in [3], and present a se- 22 quence of numerical experiments that give evidence that the theory does in fact pre- 23 dict the contraction properties of AFEM with inexact solvers. The various routines 24 used are all designed to maintain a linear-time computational complexity.

An outline of the paper is as follows. In Sect. 2, we give a brief overview of the 26 Poisson-Boltzmann equation. In Sect. 3, we describe AFEM algorithms, and intro- 27 duce a variation involving inexact solvers. In Sect. 4, we give a sequence of numerical 28 experiments that support the theoretical statements on convergence and optimality. 29 Finally, in Sect. 5 we make some final observations.

# 2 Regularized Poisson-Boltzmann Equation

We use standard notation for Sobolev spaces. In particular, we denote  $\|\cdot\|_{0,G}$  the  $L^2$ norm on any subset  $G \subset \mathbb{R}^3$ , and denote  $\|\cdot\|_{1,2,G}$  the  $H^1$  norm on G.

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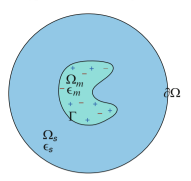


Fig. 1. Schematic of a molecular domain

Let  $\Omega := \Omega_m \cup \Gamma \cup \Omega_s$  be a bounded Lipschitz domain in  $\mathbb{R}^3$ , which consists of the molecular region  $\Omega_m$ , the solvent region  $\Omega_s$  and their interface  $\Gamma := \overline{\Omega}_m \cap \overline{\Omega}_s$  35 (see Fig. 1). Our interest in this paper is to solve the following regularized Poisson-Boltzmann equation in the weak form: find  $u \in H_g^1(\Omega) := \{u \in H^1(\Omega) : u|_{\partial\Omega} = g\}$  37 such that

$$a(u,v) + (b(u),v) = (f,v) \quad \forall v \in H_0^1(\Omega), \tag{1}$$

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where  $a(u,v) = \int_{\Omega} \varepsilon \nabla u \cdot \nabla v dx$ ,  $(b(u),v) = \int_{\Omega} \kappa^2 \sinh(u)v dx$ . Here we assume that 39 the diffusion coefficient  $\varepsilon$  is piecewise positive constant  $\varepsilon|_{\Omega_m} = \varepsilon_m$  and  $\varepsilon|_{\Omega_s} = \varepsilon_s$ . The 40 modified Debye-Hückel parameter  $\kappa^2$  is also piecewise constant with  $\kappa^2(x)|_{\Omega_m} = 0$  41 and  $\kappa^2(x)|_{\Omega_s} > 0$ . The equation (1) arises from several regularization schemes (cf. 42 [5, 6]) of the nonlinear Poisson-Boltzmann equation:

$$-\nabla \cdot (\varepsilon \nabla u) + \kappa^2 \sinh u = \sum_{i=1}^N z_i \delta(x_i),$$

where the right hand side represents N fixed points with charges  $z_i$  at positions  $x_i$ , 44 and  $\delta$  is the Dirac delta distribution.

It is easy to verify that the bilinear form in (1) satisfies:

$$c_0 \|u\|_{1,2}^2 \le a(u,u), \qquad a(u,v) \le c_1 \|u\|_{1,2} \|v\|_{1,2}, \qquad \forall u,v \in H_0^1(\Omega),$$

where  $0 < c_0 \le c_1 < \infty$  are constants depending only on  $\varepsilon$ . These properties imply 47 the norm on  $H_0^1(\Omega)$  is equivalent to the energy norm  $||\cdot||: H_0^1(\Omega) \to \mathbb{R}$ , 48

$$|||u|||^2 = a(u,u),$$
  $c_0||u||_{1,2}^2 \le |||u|||^2 \le c_1||u||_{1,2}^2.$ 

Let  $\mathscr{T}_h$  be a shape-regular conforming triangulation of  $\Omega$ , and let  $V_g(\mathscr{T}_h) := \{ v \in \mathcal{T}_h \}$  be the standard piecewise linear finite element space of defined on  $\mathscr{T}_h$ . For simplicity, we assume that the interface  $\Gamma$  is resolved by  $\mathscr{T}_h$ . Then the finite element approximation of (1) reads: find  $u_h \in V_g(\mathscr{T}_h)$  such that

$$a(u_h, v) + (b(u_h), v) = (f, v), \quad \forall v \in V_0(\mathscr{T}_h). \tag{2}$$

We close this section with a summary of a priori  $L^{\infty}$  bounds for the solution u 53 to (1) and the discrete solution  $u_h$  to (2), which play a key role in the finite element 54 error analysis of (2) and adaptive algorithms. For interested reader, we refer to [5, 9] 55 for details.

**Theorem 1.** There exist  $u_+, u_- \in L^{\infty}(\Omega)$  such that the solution u of (1) satisfies the 57 following a priori  $L^{\infty}$  bounds: 58

$$u_{-} \le u \le u_{+}, \qquad a.e. \text{ in } \Omega.$$
 (3)

Moreover, if the triangulation  $\mathcal{T}_h$  satisfies that

$$a(\phi_i, \phi_j) \le -\frac{\sigma}{h^2} \sum_{e_i : \subset \tau} |\tau|, \quad for some \quad \sigma > 0,$$
 (4)

for all the adjacent vertices  $i \neq j$  with the basis function  $\phi_i$  and  $\phi_j$ , then the discrete 60 solution  $u_h$  of (2) also has the a priori  $L^{\infty}$  bound

$$||u_h||_{L^{\infty}(\Omega)} \le C, \tag{5}$$

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where C is a constant independent of h.

We note that the mesh condition is generally not needed practically, and in fact can 63 also be avoided in analysis for certain nonlinearites [2].

# 3 Adaptive FEM with Inexact Solvers

Given a discrete solution  $u_h \in V_g(\mathcal{T}_h)$ , let us define the residual based error indicator 66  $\eta(u_h, \tau)$ :

$$\eta^2(u_h, \tau) = h_{\tau}^2 \|b(u_h) - f\|_{0, \tau}^2 + \sum_{e \subset \partial \tau} h_e \|[(\varepsilon \nabla u_h) \cdot n_e]\|_{0, e}^2,$$
 68

where  $[(\varepsilon \nabla u_h) \cdot n_e]$  denote the jump of the flux across a face e of  $\tau$ . For any subset  $\mathscr{S} \subset \mathscr{T}_h$ , we set  $\eta^2(u_h,\mathscr{S}) := \sum_{\tau \in \mathscr{S}} \eta^2(u_h,\tau)$ . By using the a priori  $L^{\infty}$  bounds 70 Theorem 1, we can show (cf. [9]) that the error indicator satisfies: 71

$$|||u - u_h||^2 < C_1 \eta^2(u_h, \hat{\mathcal{T}}_h);$$
 (6)

and 72

$$|\eta(v,\tau) - \eta(w,\tau)| \le C_2 ||v - w||_{\omega_{\tau}}, \quad \forall v, w \in V_g(\mathscr{T}_h)$$

$$\tag{7}$$

where  $\omega_{\tau} = \bigcup_{\tau' \in \mathscr{T}_h, \bar{\tau}' \cap \bar{\tau} \neq 0} \tau'$  and  $\|\|v\|\|_{\omega_{\tau}}^2 = \int_{\omega_{\tau}} \varepsilon |\nabla v|^2 dx$ . 73 Given an initial triangulation  $\mathscr{T}_0$ , the standard adaptive finite element method 74 (AFEM) generates a sequence  $|u_k, \mathcal{T}_k, \{\eta(u_k, \tau)\}_{\tau \in \mathcal{T}_k}|$  based on the iteration of the 75 form: 76

$$SOLVE \rightarrow ESTIMATE \rightarrow MARK \rightarrow REFINE.$$
 77

Here the SOLVE subroutine is usually assumed to be exact, namely  $u_k$  is the exact 78 solution to the nonlinear equation (2); the ESTIMATE routine computes the elementwise residual indicator  $\eta(u_k, \tau)$ ; the MARK routine uses standard Dörfler marking 80 (cf. [7]) where  $\mathcal{M}_k \subset \mathcal{T}_k$  is chosen so that

$$\eta(u_k, \mathscr{M}_k) \ge \theta \eta(u_k, \mathscr{T}_k)$$
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for some parameter  $\theta \in (0,1]$ ; finally, the routine REFINE subdivide the marked 83 elements and possibly some neighboring elements in certain way such that the new 84 triangulation preserves shape-regularity and conformity.

During last decade, a lot of theoretical work has been done to show the conver-86 gence of the AFEM with exact solver (see [11] and the references cited therein for 87 linear PDE case, and [10] for nonlinear PDE case). To the best of the authors knowledge, there are only a couple of convergence results of AFEM for symmetric linear 89 elliptic equations (cf. [1, 12]) which take the numerical error into account. To distinct 90 with the exact solver case, we use  $\hat{u}_k$  and  $\hat{\mathcal{T}}_k$  to denote the numerical approximation 91 to (2) and the triangulation obtained from the adaptive refinement using the inexact 92 solutions.

Due to the page limitation, we only state the main convergence result of the 94 AFEM with inexact solver for solving (1) below. More detailed analysis and extension are reported in [3].

**Theorem 2.** Let  $\{\hat{\mathcal{T}}_k, \hat{u}_k\}_{k\geq 0}$  be the sequence of meshes and approximate solutions 97 computed by the AFEM algorithm. Let u denote the exact solution and  $u_k$  denote 98 the exact discrete solutions on the meshes  $\hat{\mathcal{T}}_k$ . Then, there exist constants  $\mu > 0$ , 99  $v \in (0,1), \gamma > 0$ , and  $\alpha \in (0,1)$  such that if the inexact solutions satisfy 100

$$\mu \| u_k - \hat{u}_k \|^2 + \| u_{k+1} - \hat{u}_{k+1} \|^2 \le \nu \eta^2(\hat{u}_k, \hat{\mathcal{T}}_k)$$
(8)

then

$$\mu \| u_{k} - \hat{u}_{k} \|^{2} + \| u_{k+1} - \hat{u}_{k+1} \|^{2} \leq \nu \eta^{2} (\hat{u}_{k}, \hat{\mathcal{T}}_{k})$$

$$\| u - u_{k+1} \|^{2} + \gamma \eta^{2} (\hat{u}_{k+1}, \hat{\mathcal{T}}_{k+1}) \leq \alpha^{2} (\| u - u_{k} \|^{2} + \gamma \eta^{2} (\hat{u}_{k}, \hat{\mathcal{T}}_{k})).$$

$$(9)$$

Consequently,  $\lim_{k\to\infty} u_k = \lim_{k\to\infty} \hat{u}_k = u$ . 102

The proof of this theorem is based on the upper bound (6) of the exact solution, 103 the Lipschitz property (7) of the error indicator, Dörfler marking, and the following 104 quasi-orthogonality between the exact solutions: 105

$$|||u - u_{k+1}||^2 \le \Lambda |||u - u_k||^2 - ||u_{k+1} - u_k||^2$$
(10)

where  $\Lambda$  can be made close to 1 by refinement. For a proof of the inequality (10), see 106 for example [9].

To achieve the optimal computational complexity, we should avoid solving the 108 nonlinear system (2) as much as we could. The two-grid algorithm [13] shows that a 109 nonlinear solver on a coarse grid combined with a Newton update on the fine grid still 110 yield quasi-optimal approximation. Motivated by this idea, we propose the following AFEM algorithm with inexact solver, which contains only one nonlinear solver 112 on the coarsest grid, and Newton updates on each follow-up steps: In Algorithm 1, 113

121

126

```
Algorithm 1: \left[\hat{u}_k, \hat{\mathcal{T}}_k, \{\eta(\hat{u}_k, \tau)\}_{\tau \in \hat{\mathcal{T}}_k}\right] := \text{Inexact AFEM}(\mathcal{T}_0, \theta)
1 \hat{u}_0 = u_0 := \mathsf{NSOLVE}(\mathscr{T}_0)
                                                                  %Nonlinear solver on initial triangulation
2 for k := 0, 1, \cdots do
            \{\eta(\hat{u}_k, 	au)\}_{	au \in \hat{\mathscr{T}}_k} := \mathsf{ESTIMATE}(\hat{u}_k, \hat{\mathscr{T}}_k)
            \mathcal{M}_k := \mathsf{MARK}(\{\eta(\hat{u}_k, \tau)\}_{\tau \in \widehat{\mathscr{T}}_k}, \widehat{\mathscr{T}}_k, \theta)
            \hat{\mathscr{T}}_{k+1} := \mathsf{REFINE}(\hat{\mathscr{T}}_k, \mathscr{M}_k)
            \hat{u}_{k+1} := \mathsf{UPDATE}(\hat{u}_k, \hat{\mathcal{T}}_{k+1})
                                                                           %One-step Newton update
7 end
```

the NSOLVE routine is used only on the coarsest mesh and is implemented using 114 Newton's method run to certain convergence tolerance. For the rest of the solutions, 115 a single step of Newton's method is used to update the previous approximation. That 116 is, UPDATE computes  $\hat{u}_{k+1}$  such that

$$a(\hat{u}_{k+1} - \hat{u}_k, \phi) + (b'(\hat{u}_k)(\hat{u}_{k+1} - \hat{u}_k), \phi) = 0$$
(11)

for every  $\phi \in V(\hat{\mathcal{D}}_{k+1})$ . We remark that since (11) is only a linear problem, we could use the local multilevel method to solve it in (near) optimal complexity (cf. [4]). 119 Therefore, the overall computational complexity of the Algorithm 1 is nearly opti- 120 mal.

We should point out that it is not obvious how to enforce the required approximation property (8) that  $\hat{u}_k$  must satisfy for the theorem. This is examined in more detail 123 in [3]. However, numerical evidence in the following section shows Algorithm 1 is an 124 efficient algorithm, and the results matches the ones from AFEM with exact solver.

# **4 Numerical Experiments**

In this section we present some numerical experiments to illustrate the result in Theorem 2, implemented with FETK [8]. The software utilizes the standard piecewise- 128 linear finite element space for discretizing (1). Algorithm 1 is implemented with care 129 taken to guarantee that each of the steps runs in linear time relative to the number 130 of vertices in the mesh. The linear solver used is Multigrid preconditioned Conju- 131 gate Gradients. The estimator is computed using a high-order quadrature rule, and, 132 as mentioned above, the marking strategy is Dörfler marking where the estimated 133 errors have been binned to maintain linear complexity while still marking the ele- 134 ments with the largest error. Finally, the refinement is longest edge bisection, with 135 refinement outside of the marked set to maintain conformity of the mesh.

We present two sets of results in order to explore the effects of the inexact solver 137 in multiple contexts. For each problem, we present a convergence plot using both 138 inexact and exact solvers (including a reference line of order  $N^{-\frac{1}{3}}$ ) as well as a 139 representative cut-away of a mesh with around 30,000 vertices. The exact discrete 140 solution is computed using the standard AFEM algorithm where the solution on each 141 mesh is computed by allowing Newton's method to continue running to convergence 142 with the tolerance  $10^{-7}$ . For the exact solution, one could choose to start with an 143 arbitrary initial guess, such as the zero solution, or, as we've chosen, use the solution 144 computed on the previous mesh. Making this choice can drastically decrease the 145 number of Newton steps needed to achieve convergence. For each problem below, 146 we discuss the amount of time/computation saved using the inexact solver over this 147 exact solver.

Note that using the inexact solver modifies not only the solution on a given mesh, 149 but also the sequence of meshes generated, since the algorithm may mark different 150 simplices. However, as shown in the examples below, the inexact solutions still maintain optimal convergence rates.

The first result uses constant coefficients across the entire domain  $\Omega = [0, 1]^3$ , an 153 exponential nonlinearity, and a right hand side chosen so that the derivative of the 154 exact solution is large near the origin. The boundary conditions chosen for this prob- 155 lem are homogeneous Dirichlet boundary conditions. Specifically, the exact solution 156 is given by  $u = u_1 u_2$  where

$$u_1 = \sin(\pi x)\sin(\pi y)\sin(\pi z)$$

152

157

159

is chosen to satisfy the boundary condition and

$$u_2 = 3(x^2 + y^2 + x^2 + 10^{-4})^{-1.5}$$
.

The results can be seen in Fig. 2.

For this problem, the number of iterations in Newton's method by the exact solver 162 varied between 3 and 7, depending on the refinement level. Because all steps of 163 the algorithm are designed to be linear, this suggests that the inexact solver runs 164 at least three times faster for this problem, while still maintaining optimal order of 165 convergence.

In order to test the robustness to the addition of jump coefficients, the second 167 result uses the domain  $\Omega = [-1,1]^3$  and  $\Omega_m = \left[-\frac{1}{4},\frac{1}{4}\right]$  with constants  $\varepsilon_s = 80, \varepsilon_m = 168$ 2,  $\kappa_s = 1$ , and  $\kappa_m = 0$ . Homogeneous Neumann conditions are chosen for the boundary and the right hand side is simplified to a constant. Because an exact solution is 170 unavailable for this (and the following) problem, the error is computed by comparing to a discrete solution on a mesh with around ten times the number of vertices 172 as the finest mesh used in the adaptive algorithm. Figure 3 shows the results for this 173 problem. As can be seen the refinement favors the interface and the inexact and exact 174 solvers perform as expected.

Once again, for this problem, the exact solver required between 3 and 9 iterations 176 of Newton's method to reach convergence, depending on the refinement level. Since 177 the run time is linear is the number of iterations, this result gives a speedup of at least 178 three times using the inexact solver, without causing a loss in convergence rate.

5 Conclusion 180

In this article we have studied AFEM with inexact solvers for a class of semilinear 181 elliptic interface problems with discontinuous diffusion coefficients. The algorithm 182

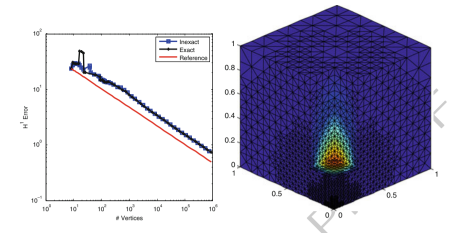


Fig. 2. Convergence plot and mesh cut-away for the corner singularity problem

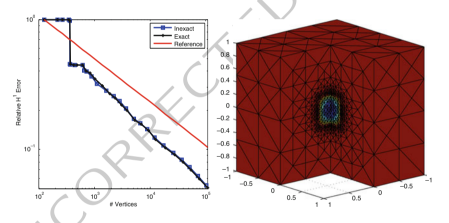


Fig. 3. Convergence plot and mesh cut-away for the Poisson-Boltzmann problem

we studied consisted of the standard SOLVE-ESTIMATE-MARK-REFINE proce- 183 dure common to many adaptive finite element algorithms, but where the SOLVE step 184 involves only a full solve on the coarsest level, and the remaining levels involve only 185 single Newton updates to the previous approximate solution. Our numerical results 186 indicate that the recently developed AFEM convergence theory for inexact solvers 187 in [3] does predict the actual behavior of the methods and can allow for significant 188 speedup in the approximation of solutions.

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# **Preconditioning for Mixed Finite Element Formulations of Elliptic Problems**

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Summary. In this paper, we discuss a preconditioning technique for mixed finite element 13 discretizations of elliptic equations. The technique is based on a block-diagonal approximation 14 of the mass matrix which maintains the sparsity and positive definiteness of the corresponding 15 Schur complement. This preconditioner arises from the multipoint flux mixed finite element 16 method and is robust with respect to mesh size and is better conditioned for full permeability 17 tensors than a preconditioner based on a diagonal approximation of the mass matrix.

1 Introduction 19

Consider the mixed formulation of a second order linear elliptic equation. Introduc- 20 ing a flux variable, we solve for a scalar potential p and a vector function **u** that 21 satisfy

$$\mathbf{u} = -\mathbb{K}\nabla p \quad \text{in } \Omega, \tag{1}$$

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$$\nabla \cdot \mathbf{u} = f \qquad \text{in } \Omega,$$

$$p = 0 \qquad \text{on } \partial \Omega,$$
(2)

$$p = 0$$
 on  $\partial \Omega$ . (3)

where  $\Omega$  is a polygonal domain with Lipschitz continuous boundary and  $\mathbb K$  is a 23 symmetric and uniformly positive definite tensor with  $L^{\infty}(\Omega)$  components. Homo- 24 geneous Dirichlet boundary conditions are considered for the simplicity of the pre- 25

Mixed finite element methods lead to the non-singular indefinite system:

$$\mathbb{M}\begin{pmatrix} U \\ P \end{pmatrix} := \begin{pmatrix} \mathbb{A} \ \mathbb{B}^T \\ \mathbb{B} \ 0 \end{pmatrix} \begin{pmatrix} U \\ P \end{pmatrix} = \begin{pmatrix} 0 \\ F \end{pmatrix}, \tag{4}$$

where the matrix  $\mathbb{A}$  is a symmetric and positive definite.

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In this paper, we consider preconditioners of the form:

$$\widetilde{\mathbb{M}} := \begin{pmatrix} \widetilde{\mathbb{A}} \ \mathbb{B}^T \\ \mathbb{B} \ 0 \end{pmatrix}. \tag{5}$$

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The applicability of this type preconditioner is due to the fact that

- $\widetilde{\mathbb{A}}$  is easily invertible.
- The Schur complement of the preconditioner  $\widetilde{M}$  is sparse and positive definite, and can be solved easily.

One way is choosing  $\widetilde{\mathbb{A}}$  as a diagonal matrix. In [1],  $\widetilde{\mathbb{A}}$  is given as  $\omega \mathbb{I}$ . The global 34 parameter  $\omega$  is chosen to minimize the spectral radius of  $\mathbb{I} - \mathbb{M}^{-1}\mathbb{M}$ . In [5], the diagonal matrix is optimally scaled at element level and a precise upper bound of the 36 spectral radius has been shown:  $\rho(\mathbb{I} - \widetilde{\mathbb{M}}^{-1}\mathbb{M}) \leq 1/2$ . In other words, the preconditioner is independent of both the mesh size and the tensor K. This uniformity is 38 derived when the problem has a diagonal K and is discretized by the lowest order 39 Raviart-Thomas [8] mixed finite element on rectangular grids. For other mixed fi- 40 nite element spaces or full tensor K, the uniformity result is not clearly understood. 41 Alternatively, a simple parameter-free choice for  $\mathbb{A}$ ,  $\mathbb{A} = \text{Diag}(\mathbb{A})$ , can be used.

Another approach is to take A as a block-diagonal matrix which guarantees that 43 the corresponding Schur complement matrix is sparse and positive definite. Multi- 44 point flux mixed finite element (MFMFE) methods [6, 9-12] give matrices of the 45 form (5), where the flux variable can be locally eliminated due to the block-diagonal 46 structure of A. The corresponding Schur complement gives a cell-centered stencil 47 for the scalar variable. In this paper, we study the performance of this MFMFE 48 operator as a preconditioner. The Schur complement of MFMFE has a 9-point 49 stencil on logically rectangular grids and with full tensor K in contrast to 5-point 50 stencil which arises if A is a diagonal matrix. Our numerical result indicates that 51 the MFMFE method gives a better preconditioner than the diagonal preconditioner 52 (A = Diag(A)). A natural extension of this work is the use of approximate preconditioners based on algebraic multigrid for MFMFE as described in [2, 7] and will be 54 the subject of future work.

The rest of the paper is organized as follows. Mixed finite element formulation 56 is described in Sect. 2. A block type preconditioner is discussed in Sect. 3. Finally in 57 Sect. 4, numerical experiments are given.

## 2 Mixed Finite Element Formulation

Define  $H(\operatorname{div};\Omega) := \{ \mathbf{v} \in (L^2(\Omega))^d : \nabla \cdot \mathbf{v} \in L^2(\Omega) \}$  and let  $(\cdot,\cdot)$  denote the inner 60 product in  $L^2(\Omega)$ . Let  $X \leq (\geq) Y$  denote that there exists a constant C, independent 61 of the mesh size h, such that  $X \leq (\geq)$  CY. The notation  $X \approx Y$  means that both  $X \lesssim Y$  62 and  $X \gtrsim Y$  hold.

Let  $\mathcal{T}_h$  be a finite element partition of the domain  $\Omega$  consisting of either triangles 64 or quadrilaterals. We assume that  $\mathcal{T}_h$  is shape-regular in the sense of Ciarlet [4]. 65

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The finite element spaces on any physical element  $E \in \mathcal{T}_h$  are defined via the Piola 66 transformation 67

$$\mathbf{v} \leftrightarrow \hat{\mathbf{v}} : \hat{\mathbf{v}} = \frac{1}{J_F} \mathbb{DF}_E \hat{\mathbf{v}} \circ F_E^{-1},$$
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and the scalar transformation

$$w \leftrightarrow \hat{w} : w = \hat{w} \circ F_F^{-1}, \tag{70}$$

where  $F_E$  denotes a mapping from the reference element  $\hat{E}$  to the physical element 71 E,  $\mathbb{DF}_E$  is the Jacobian of  $F_E$ , and  $J_E$  is its determinant. The finite element spaces  $V_h$  72 and  $W_h$  on  $\mathcal{T}_h$  are given by

$$V_h = \left\{ \mathbf{v} \in H(\operatorname{div}; \Omega) : \quad \mathbf{v}|_E \leftrightarrow \hat{\mathbf{v}}, \ \hat{\mathbf{v}} \in \hat{V}(\hat{E}), \quad \forall E \in \mathcal{T}_h \right\},$$

$$W_h = \left\{ w \in L^2(\Omega) : \quad w|_E \leftrightarrow \hat{w}, \ \hat{w} \in \hat{W}(\hat{E}), \quad \forall E \in \mathcal{T}_h \right\},$$

where  $V(\hat{E})$  and  $\hat{W}(\hat{E})$  are the lowest order Brezzi-Douglas-Marini (BDM<sub>1</sub>) spaces 74 on the reference element  $\hat{E}$ . Definitions of Piola transformation and BDM<sub>1</sub> spaces 75 yield  $V_h \subset H(\text{div}; \Omega)$  and  $W_h \subset L^2(\Omega)$ . 76

The finite element method reads: find  $\mathbf{u}_h \in V_h$  and  $p_h \in W_h$ , such that

$$(\mathbb{K}^{-1}\mathbf{u}_{h}, \mathbf{v}) - (p_{h}, \nabla \cdot \mathbf{v}) = 0, \qquad \forall \mathbf{v} \in V_{h},$$

$$-(\nabla \cdot \mathbf{u}_{h}, w) = -(f, w) \quad \forall w \in W_{h}.$$
(6)

$$-(\nabla \cdot \mathbf{u}_h, w) = -(f, w) \quad \forall w \in W_h. \tag{7}$$

The method (6) and (7) can have a second order convergence for the flux and first 78 order convergence for the scalar potential [3] if  $\mathbf{u}$  and p are sufficiently regular. 79

# 3 Preconditioning the Mixed Finite Element System

#### 3.1 Multipoint Flux Mixed Finite Element

A family of multipoint flux mixed finite element (MFMFE) methods on various grids 82 has been developed and analyzed [6, 9–12]. The method is defined as: find  $\mathbf{u}_h \in V_h$  83 and  $p_h \in W_h$ , such that

$$(\mathbb{K}^{-1}\mathbf{u}_h, \mathbf{v})_Q - (p_h, \nabla \cdot \mathbf{v}) = 0, \qquad \forall \mathbf{v} \in V_h,$$
(8)

$$-(\nabla \cdot \mathbf{u}_h, w) = -(f, w) \quad \forall w \in W_h, \tag{9}$$

where the finite element spaces are BDM<sub>1</sub> on triangular and rectangular meshes. 85 Compared to the BDM<sub>1</sub> finite element method, a specific numerical quadrature rule 86 is employed. It is defined as: 87

$$(\mathbb{K}^{-1}\mathbf{q},\mathbf{v})_{Q} = \sum_{E \in \mathcal{F}_{h}} (\mathbb{K}^{-1}\mathbf{q},\mathbf{v})_{Q,E} \equiv \sum_{E \in \mathcal{F}_{h}} \operatorname{Trap}(\mathscr{K}\hat{\mathbf{q}},\hat{\mathbf{v}})_{\hat{E}},$$
(10)

where  $\mathcal{K}$  on each  $\hat{E}$  is defined as

$$\mathcal{K} = \frac{1}{J_E} \mathbb{DF}_E^T \mathbb{K}^{-1}(F_E(\hat{x})) \mathbb{DF}_E, \tag{11}$$

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and the trapezoidal rule on  $\hat{E}$  is denoted as

$$\operatorname{Trap}(\hat{\mathbf{q}}, \hat{\mathbf{v}})_{\hat{E}} \equiv \frac{|\hat{E}|}{m} \sum_{i=1}^{m} \hat{\mathbf{q}}(\hat{\mathbf{r}}_i) \cdot \hat{\mathbf{v}}(\hat{\mathbf{r}}_i), \tag{12}$$

with  $\{\hat{\mathbf{r}}_i\}_{i=1}^m$  being vertices of  $\hat{E}$  and m being the number of vertices of  $\hat{E}$ 

The degrees of freedom for the flux variable are chosen as the normal components 91 at two vertices on each edge. More specifically, denote the basis functions associated 92 with  $\hat{\bf r}_i$  by  $\hat{\bf v}_{ij}$ , j=1,2:  $(\hat{\bf v}_{ij}\cdot\hat{\bf n}_{ij})(\hat{\bf r}_i)=1$ ,  $(\hat{\bf v}_{ij}\cdot\hat{\bf n}_{ik})(\hat{\bf r}_i)=0$ ,  $k\neq j$ , and  $(\hat{\bf v}_{ij}\cdot\hat{\bf n}_{lk})(\hat{\bf r}_l)=93$  0,  $l\neq i$ , k=1,2. As a consequence, the quadrature rule (10) couples only the two 94 basis functions associated with a vertex. For example, on the unit square 95

$$(\mathcal{K}\hat{\mathbf{v}}_{11},\hat{\mathbf{v}}_{11})_{\hat{Q},\hat{E}} = \frac{\mathcal{K}_{11}(\hat{\mathbf{r}}_1)}{4}, \quad (\mathcal{K}\hat{\mathbf{v}}_{11},\hat{\mathbf{v}}_{12})_{\hat{Q},\hat{E}} = \frac{\mathcal{K}_{21}(\hat{\mathbf{r}}_1)}{4},$$

$$(\mathcal{K}\hat{\mathbf{v}}_{11},\hat{\mathbf{v}}_{ij})_{\hat{Q},\hat{E}} = 0, \quad i \neq 1, j = 1, 2.$$

$$(13)$$

where  $\mathcal{K}_{ij}$  denotes *i*-th row and *j*-th column of the matrix function  $\mathcal{K}$ . This localization property on interactions between the flux basis functions gives the assembled mass matrix in (8) has a block diagonal structure with one block per grid vertex.

We denote the algebraic system arising from (8) and (9) as

$$\begin{pmatrix} \mathbb{A}_Q \ \mathbb{B}^T \\ \mathbb{B} \ 0 \end{pmatrix} \begin{pmatrix} U \\ P \end{pmatrix} = \begin{pmatrix} 0 \\ F \end{pmatrix}, \tag{14}$$

where  $\mathbb{A}_O$  is block diagonal. The approximate flux, U, can be easily eliminated via

$$U = -\mathbb{A}_Q^{-1} \mathbb{B}^T P. \tag{15}$$

The resulting Schur complement system

$$\mathbb{B}\mathbb{A}_O^{-1}\mathbb{B}^T P = -F,\tag{16}$$

is symmetric positive definite and sparse. On rectangular grids, Eq. (16) has a 102 5-point stencil for a diagonal tensor  $\mathbb{K}$  and 9-point stencil for the full tensor. The 103 Schur complement system can be solved using classical algebraic multigrid methods. 104 The flux variable is then obtained easily by (15) due to the block diagonal structure of  $\mathbb{A}_{\mathcal{O}}$ . 106

The following result concerns the convergence of the MFMFE methods. Let  $W^{k,\infty}_{\mathscr{T}_h}$  107 consist of functions  $\phi$  such that  $\phi|_E \in W^{k,\infty}(E)$  for all  $E \in \mathscr{T}_h$ .

**Theorem 1** ([6, 10–12]). Let  $\mathcal{T}_h$  consist of simplices,  $h^2$ -parallelograms,  $h^2$ 

$$\|\mathbf{u} - \mathbf{u}_h\| \lesssim h \|\mathbf{u}\|_1, \ \|\nabla \cdot (\mathbf{u} - \mathbf{u}_h)\| \lesssim h \|\nabla \cdot \mathbf{u}\|_1, \ \|p - p_h\| \lesssim h(\|\mathbf{u}\|_1 + \|p\|_1).$$

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Compared to the second order $L^2$ convergence of the flux variable in the BDM <sub>1</sub>	112
mixed method, the MFMFE has a first order convergence for the flux variable due to	113
the numerical quadrature. However the MFMFE method is a solver friendly scheme	114
since the MFMFE method can be reduced to a cell-centered stencil in terms of the	115
scalar variable without solving a saddle-point problem.	116

# 3.2 Multipoint Flux Mixed Finite Element as a Preconditioner

The MFMFE method may be used as a preconditioner to the BDM $_1$  mixed finite element method by choosing  $\widetilde{\mathbb{A}}=\mathbb{A}_Q$ .

**Lemma 1.** The condition number of  $\widetilde{\mathbb{A}}^{-1}\mathbb{A}$  is independent of the mesh size.

*Proof.* It has been shown [6, 11, 12] that the bilinear form  $(\mathbb{K}^{-1}\cdot,\cdot)_Q$  is an inner product in  $\mathbf{V}_h$  and  $(\mathbb{K}^{-1}\mathbf{q},\mathbf{q})_Q^{1/2}$  is a norm equivalent to the  $L^2$  norm. Thus

$$(\mathbb{K}^{-1}\mathbf{q},\mathbf{q})_{Q} \approx \|\mathbf{q}\|^{2} \approx (\mathbb{K}^{-1}\mathbf{q},\mathbf{q}), \quad \forall \mathbf{q} \in \mathbf{V}_{h}. \qquad \Box$$
 (17)

The preconditioner of the form (5) has been analyzed by Ewing, Lazarov, Lu and 123 Vassilevski.

**Theorem 2 ([5]).** The eigenvalues of  $\widetilde{\mathbb{M}}^{-1}\mathbb{M}$  are real and positive and lie in the 125 interval  $[\lambda_{min}, \lambda_{max}]$ , where  $\lambda_{min}$  and  $\lambda_{max}$  are the extreme eigenvalues of  $\widetilde{\mathbb{A}}^{-1}\mathbb{A}$ . 126

By Lemma 1 and Theorem 2, we have the following corollary.

**Corollary 1.** The preconditioned system of  $BDM_1$  mixed finite element method with 128 MFMFE as a preconditioner is positive definite. The condition number is independent of the mesh size. 129

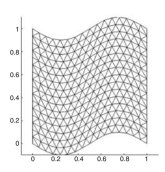
# **4 Numerical Results**

4.1 Example 1

In this example, we consider (1)–(3) on the computational domain shown in Fig. 1 133 (left) with p = 0 on  $\partial \Omega$  and f = 1.

First, we use the MFMFE method as a preconditioner for the BDM $_1$  mixed finite 135 element method with  $\mathbb{K} = \mathbb{I}$ . The result is presented in Table 1 where we can clearly 136 see that the preconditioner is robust with respect to the mesh size h. Next, we consider 137 the heterogeneous permeability field shown in Fig. 1 (right) which is generated using 138 geostatistical techniques (kriging) with a longer correlation length in the horizontal 139 direction. In Table 2 we see that the preconditioner is not only robust with respect to 140 mesh size, but also with respect to the heterogeneities in the permeability. 141

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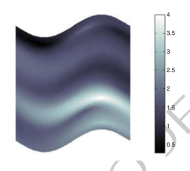


Fig. 1. The triangular mesh used in Example 1 with  $h \approx 1/16$  (*left*) and the log of the heterogeneous permeability field (right)

h	Degrees of Freedom	$cond(\widetilde{\mathbb{M}}^{-1}\mathbb{M})$
1/8	512	13.43
1/16	2048	15.84
1/32	8192	15.61
1/64	32768	15.63

**Table 1.** Performance of the MFMFE preconditioner with a homogeneous permeability field.

h	Degrees of Freedom	$\operatorname{cond}(\widetilde{\mathbb{M}}^{-1}\mathbb{M})$
1/8	512	20.07
1/16	2048	21.61
1/32	8192	16.61
1/64	32768	14.27

Table 2. Performance of the MFMFE preconditioner with a heterogeneous permeability field.

#### 4.2 Example 2

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In this example, we consider (1)–(3) with  $\Omega = [0,1] \times [0,1]$  and

$$\mathbb{K} = \begin{pmatrix} 1 + \alpha & 1 - \alpha \\ 1 - \alpha & 1 + \alpha \end{pmatrix},$$

with  $0 < \alpha \le 1$ . We use uniform rectangular meshes and our objective is to demonstrate that the MFMFE preconditioner is more robust as  $\alpha \to 0$ . In Tables 3 and 146 4 we present the results using the diagonal preconditioner ( $\mathbb{A} = \text{Diag}(\mathbb{A})$ ) and the MFMFE preconditioner respectively. We see that both preconditioners are robust 148 with respect to h, but degrade as  $\alpha \to 0$ , but the MFMFE preconditioner degrades at 149 a much slower rate.

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α	h = 1/4	h = 1/8	h = 1/16	h = 1/32
1	22.43	22.32	22.32	22.32
1E-1	1.06E2	9.95E2	1.06E2	1.06E2
1E-2	7.00E2	6.97E2	6.97E2	6.97E2
1E-3	9.51E3	9.41E3	9.75E3	8.42E3

**Table 3.** Performance of a diagonal preconditioner with respect to h and  $\alpha$ .

$\alpha$	h = 1/4	h = 1/8	h = 1/16	h = 1/32
1	22.42	22.32	22.32	22.32
1E-1	32.07	32.09	32.26	32.09
1E-2	51.01	50.06	50.39	50.39
1E-3	5.20E2	6.96E2	8.10E2	8.21E2

**Table 4.** Performance of the MFMFE preconditioner with respect to h and  $\alpha$ .

5 Conclusions 151

The purpose of this paper is to investigate the performance of the multipoint flux 152 mixed finite element as a preconditioner for the saddle-point system for the full 153 BDM<sub>1</sub> mixed finite element approximation. Numerical results indicate that the 154 MFMFE preconditioner is robust with respect to the mesh size and performs bet- 155 ter than the preconditioner based on the diagonal mass matrix.

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### **Multigrid Preconditioner for Nonconforming** Discretization of Elliptic Problems with Jump Coefficients

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Summary. In this paper, we present a multigrid preconditioner for solving the linear system 11 arising from the piecewise linear nonconforming Crouzeix-Raviart discretization of second 12 order elliptic problems with jump coefficients. The preconditioner uses the standard conforming subspaces as coarse spaces. Numerical tests show both robustness with respect to the jump 14 in the coefficient and near-optimality with respect to the number of degrees of freedom.

1 Introduction 16

The purpose of this paper is to present a multigrid preconditioner for solving the linear system arising from the  $\mathbb{P}^1$  nonconforming Crouzeix-Raviart (CR) discretization 18 of second order elliptic problems with jump coefficients. The multigrid preconditioner we consider here uses pointwise relaxation (point Gauss-Seidel/Jacobi itera- 20 tive methods) as a smoother, followed by a subspace (coarse grid) correction which 21 uses the standard multilevel structure for the nested  $\mathbb{P}_1$  conforming finite element 22 spaces. The subspace correction step is motivated by the observation that the stan- 23 dard  $\mathbb{P}^1$  conforming space is a subspace of the CR finite element space.

The idea of using conforming subspaces to construct preconditioners for CR dis- 25 cretization has been used in [6, 9, 11] in the context of smooth coefficients. To deal 26 with the jump coefficient problems, multilevel methods using conforming subspaces 27 were proposed and analyzed in [7, 8]. In particular, the author showed that if the 28 coefficients satisfy the quasi-monotone condition (cf. [5]), then the preconditioned 29 systems have condition numbers independent of the coefficients and depending on 30 the mesh size logarithmically. The author also showed that the same conclusions 31 hold for multilevel preconditioners with an additional exotic coarse space in case of 32 general coefficient distributions with cross points.

To avoid the implementation of the additional exotic coarse space, we take 34 another approach in this paper and show that the multigrid method (without the 35

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additional exotic coarse space) is a robust preconditioner for PCG algorithm. In particular, we show that the preconditioned system has only a few "bad eigenvalues" 37 (depending on the jumps of the coefficients), and the asymptotic convergence rate 38 of the PCG algorithm will be uniform with respect to the coefficient. The analysis 39 follows closely [12] with the help of special technical tools developed in [2]. Due 40 to space limitation we only state the main result (Theorem 1 in Sect. 3), and provide 41 numerical results that support it. Detailed analyses and further discussion of the al- 42 gorithm are presented in [13]. One of the main benefits of this algorithm is that it is 43 very easy to implement in practice. The procedure is the same as the standard multigrid algorithm on conforming spaces, and the only difference is the prolongation and 45 restriction matrices on the finest level. Since the spaces are nested, the prolongation 46 matrix is simply the matrix representation of the natural inclusion operator from the 47 conforming space to the CR space.

The paper is organized as follows. In Sect. 2, we give basic notation and the finite 49 element discretizations. In Sect. 3, we present the multigrid algorithm and discuss its 50 implementation and convergence. Finally, in Sect. 4 we verify numerically the the- 51 oretical results by presenting several numerical tests for two and three dimensional 52 model problems.

2 Preliminaries 54

Let  $\Omega \subset \mathbb{R}^d$  (d=2,3) be an open polygonal domain. Given  $f \in L^2(\Omega)$ , we consider 55 the following model problem: Find  $u \in H_0^1(\Omega)$  such that 56

$$a(u,v) := (\kappa \nabla u, \nabla v) = (f,v) \quad \forall v \in H_0^1(\Omega), \tag{1}$$

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where the diffusion coefficient  $\kappa \in L^{\infty}(\Omega)$  is assumed to be piecewise constant, 57 namely,  $\kappa(x)|_{\Omega_m} = \kappa_m$  is a constant for each (open) polygonal subdomain  $\Omega_m$  satisfying  $\bigcup_{m=1}^{M} \overline{\Omega}_m = \overline{\Omega}$  and  $\Omega_m \cap \Omega_n = \emptyset$  for  $m \neq n$ .

We assume that there is an initial (quasi-uniform) triangulation  $\mathcal{I}_0$ , with mesh 60 size  $h_0$ , such that for all  $T\in\mathscr{T}_0$   $\kappa_T:=\kappa(x)|_T$  is constant. Let  $\mathscr{T}_j:=\mathscr{T}_{h_j}$   $(j=1,\cdots,J)$ be a family of uniform refinement of  $\mathcal{T}_0$  with mesh size  $h_i$ . Without loss of generality, 62 we assume that the mesh size  $h_j \simeq 2^{-j} h_0$   $(j = 0, \dots, J)$  and denote  $h = h_J$ .

On each level  $j = 0, \dots, J$ , we define  $V_i$  as the standard  $\mathbb{P}^1$  conforming finite element space defined on  $\mathcal{I}_i$ . Then the standard conforming finite element discretization 65 of (1) reads:

Find 
$$u_j \in V_j$$
 such that  $a(u_j, v_j) = (f, v_j), \quad \forall v_j \in V_j.$  (2)

For each  $i = 0, \dots, J$ , we define the induced operator for (2) as

$$(A_i v_i, w_i) = a(v_i, w_i), \quad \forall v_i, w_i \in V_i.$$

We denote  $\mathcal{E}_h$  the set of all edges (in 2D) or faces (in 3D) of  $\mathcal{T}_h$ . Let  $V_h^{CR}$  be the piecewise linear nonconforming Crouzeix-Raviart finite element space defined by:

$$V_h^{CR} = \left\{ v \in L^2(\Omega) : v_{|_T} \in \mathbb{P}^1(T) \, \forall T \in \mathcal{T}_h \text{ and } \int_e [\![v]\!]_e ds = 0 \, \forall \, e \in \mathcal{E}_h \right\},$$

where  $\mathbb{P}^1(T)$  denotes the space of linear polynomials on T and  $[v]_e$  denotes the jump 71 across the edge/face  $e \in \mathscr{E}_h$  with  $[\![v]\!]_e = v$  when  $e \subset \partial \Omega$ . In the sequel, let us denote  $V_{J+1} := V_h^{CR}$  for simplicity. We remark that all these finite element spaces are nested, 73 that is,

$$V_0 \subset \cdots \subset V_J \subset V_{J+1}$$
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The  $\mathbb{P}^1$ -nonconforming finite element approximation to (1) reads:

Find 
$$u \in V_h^{CR}$$
:  $a_h(u, w) := \sum_{T \in \mathcal{T}_I} \int_T \kappa_T \nabla u \cdot \nabla w = (f, w), \forall w \in V_h^{CR}$ . (3)

The bilinear form  $a_h(\cdot,\cdot)$  induced a natural energy norm:  $|v|_{h,\kappa}:=\sqrt{a_h(v,v)}$  for any  $v\in V_h^{CR}$ . In operator form, we are going to solve the linear system 78

$$Au = f, (4)$$

where A is the operator induced by (3), namely

$$(Av, w) = a_h(v, w), \qquad \forall v, w \in V_h^{CR}.$$

#### 3 A Multigrid Preconditioner

The action of the standard multigrid *V*-cycle preconditioner  $B := B_{J+1} : V_{J+1} \mapsto V_{J+1}$  82 on a given  $g \in V_{J+1}$  is recursively defined by the following algorithm (cf. [3]):

Let  $g_{J+1} = g$ , and  $B_0 = A_0^{-1}$ . For  $j = 1, \dots, J+1$ , we define recursively  $B_j g_j$  for any  $g_i \in V_i$  by the following three steps:

- 1. Pre-smoothing :  $w_1 = R_i g_i$ ;
- 2. Subspace correction:  $w_2 = w_1 + B_{j-1}Q_{j-1}(g_j A_jw_1);$
- 3. Post-smoothing:  $B_j g_j := w_2 + R_i^* (g_i A_i w_2)$ .

In this algorithm,  $R_j$  corresponds to a Gauss-Seidel or a Jacobi iterative method 84 known as a smoother; and  $Q_i$  is the standard  $L^2$  projection on  $V_i$ :

$$(Q_j v, w_j) = (v, w_j), \qquad \forall w_j \in V_j, \ (j = 0, \cdots, J).$$

The implementation of Algorithm 3 is almost identical to the implementation of 87 the standard multigrid V-cycle (cf. [4]). Between the conforming spaces, we use the standard prolongation and restriction matrices (for conforming finite elements). The 89

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corresponding matrices between  $V_J$  and  $V_{J+1}$ , are however different. The prolongation matrix on  $V_I$  can be viewed as the matrix representation of the natural inclusion 91  $\mathscr{I}_J: V_J \to V_{J+1}$ , which is defined by

$$(\mathscr{I}_J v)(x) = \sum_{e \in \mathscr{E}_h} v(m_e) \psi_e(x),$$
 93

where  $\psi_e$  is the CR basis on the edge/face  $e \in \mathcal{E}_h$  and  $m_e$  is the barycenter of e. 94 Therefore, the prolongation matrix has the same sparsity pattern as the edge-to-vertex 95 (in 2D), or face-to-vertex (in 3D) connectivity, and each nonzero entry in this matrix 96 equals the constant 1/d where d is the space dimension. The restriction matrix is 97 simply the transpose of the prolongation matrix.

The efficiency and robustness of this preconditioner can be analyzed in terms of the effective condition number (cf. [12]) defined as follows:

**Definition 1.** Let V be a real N dimensional Hilbert space, and  $S: V \to V$  be a 101 symmetric positive definition operator with eigenvalues  $0 < \lambda_1 \leq \cdots \leq \lambda_N$ . The m-th 102 effective condition number of S is defined by 103

$$\mathscr{K}_m(S) := \lambda_N(S)/\lambda_{m+1}(S).$$
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Note that the standard condition number  $\mathcal{K}(BA)$  of the preconditioned system BA 105 will be large due to the large jump in the coefficient  $\kappa$ . However, there might be only 106 a small (fixed) number of small eigenvalues of BA, which cause the large condition 107 number; and the other eigenvalues are bounded nearly uniformly. In particular, we 108 have the following main result:

**Theorem 1.** Let B be the multigrid V-cycle preconditioner defined in Algorithm 3. 110 Then there exists a fixed integer  $m_0 < M$ , depending only on the distribution of the 111 coefficient K, such that

$$\mathscr{K}_{m_0}(BA) \le C^2 |\log h|^2 = C^2 J^2 ,$$

where the constant C > 0 is independent of the coefficients and mesh size.

The analysis is based on the subspace correction framework [10], but some technical 114 tools developed in [2] are needed to deal with nonconformity of the finite element 115 spaces. Due to space restriction, a detailed analysis will be reported somewhere else. 116

Thanks to Theorem 1 and a standard PCG convergence result (cf. [1, Sect. 13.2]), 117 the PCG algorithm with the multigrid V-cycle preconditioner defined in Algorithm 3 118 has the following convergence estimate: 119

$$|u-u_i|_{h,\kappa} \le 2(\mathcal{K}(BA)-1)^{m_0} \left(\frac{CJ-1}{CJ+1}\right)^{i-m_0} |u-u_0|_{h,\kappa}$$
,

where  $u_0$  is the initial guess, and  $u_i$  is the solution of *i*-th PCG iteration. Although the condition number  $\mathcal{K}(BA)$  might be large, the convergence rate of the PCG algorithm 122 is asymptotically dominated by  $\frac{CJ-1}{CJ+1}$ , which is determined by the effective condition 123 number  $\mathcal{K}_{m_0}(BA)$ . Moreover, this bound of asymptotic convergence rate convergence is independent of the coefficient  $\kappa$ , but depends on the mesh size logarithmically. 125

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#### 4 Numerical Results

In this section, we present several numerical tests in 2D and 3D which verify the 127 result in Theorem 1 on the performance of the multigrid V-cycle preconditioner de- 128 scribed in the previous sections. The numerical tests show that the effective condition numbers of the preconditioned linear systems (with V-cycle preconditioner) are 130 nearly uniformly bounded.

#### 4.1 A 2D Example

As a first model problem, we consider Eq. (1) in the square  $\Omega = (-1, 1)^2$  with coefficient such that,  $\kappa(x) = 1$  for  $x \in \Omega_1 = (-0.5, 0)^2 \cup (0, 0.5)^2$ , and  $\kappa(x) = \varepsilon$  for x in the remaining subdomain,  $x \in \Omega \setminus \Omega_1$  (see Fig. 1). By decreasing the value of  $\varepsilon$  we 135 increase the contrast in the PDE coefficients.

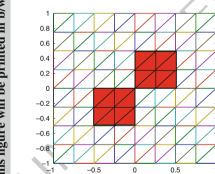
Our initial triangulation on level 0 has mesh size  $h_0 = 2^{-1}$  and resolves the interfaces where the coefficients have discontinuities. Then on each level, we uniformly 138 refine the mesh by subdividing each element into four congruent children. In this 139 example, we use 1 forward/backward Gauss-Seidel iteration as pre/post smoother 140 in the multigrid preconditioner, and the stopping criteria of the PCG algorithm is 141  $||r_k||/||r_0|| < 10^{-7}$  where  $r_k$  is the residual at k-th iteration.

0.9

0.8

0.7

0.6



0.5 0.4 0.3 0.2 0.1 0

Fig. 1. 2D computational domain

Fig. 2. Eigenvalue distribution of BA

Figure 2 shows the eigenvalue distribution of the multigrid V-cycle preconditioned system BA when  $h = 2^{-5}$  (level=4) and  $\varepsilon = 10^{-5}$ . As we can see from this 144 figure, there is only one small eigenvalue that deteriorates with respect to the jump 145 in the coefficient and the mesh size.

Table 4.1 shows the estimated condition number  $\mathcal{H}$  and the effective condition 147 number  $\mathcal{K}_1$  of BA. It can be observed that the condition number  $\mathcal{K}$  increases rapidly with respect to the increase of the jump in the coefficients and the number of de- 149 grees of freedom. On the other hand, the number of PCG iterations increases only a 150 small amount, and the corresponding effective condition number is nearly uniformly 151 bounded, as predicted by Theorem 1.

ε	levels	0	1	2	3	4	
1	$\mathscr{K}$	1.65 (8)	1.83 (10)	1.9 (10)	1.9 (10)	1.89 (10)	
1	$\mathscr{K}_1$	1.44	1.78	1.77	1.78	1.76	
$10^{-1}$	K	3.78 (10)	3.69 (11)	3.76 (12)	3.79 (12)	3.88 (12)	
10	$\mathscr{K}_1$	1.89	1.87	1.93	1.92	1.95	
$10^{-2}$	K	23.4 (12)	23.6 (13)	24.6 (13)	25.1 (14)	26 (15)	
10	$\mathscr{K}_1$	2.15	1.96	1.99	1.97	2.24	
$10^{-3}$	K	218 (13)	223 (14)	232 (15)	238 (16)	246 (16)	
10	$\mathscr{K}_1$	2.19	1.98	2	1.98	2.29	
$10^{-4}$	$ \mathscr{K} $	2.17e+3 (14)	2.21e+3 (15)	2.31e+3 (16)	2.37e+3(18)	2.45e+3 (18)	
10	$\mathscr{K}_1$	2.2	1.98	2	1.98	2.3	
$10^{-5}$	K	2.17e+4 (15)	2.21e+4 (16)	2.31e+4 (17)	2.37e+4(19)	2.76e+4(19)	
10	$\mathscr{K}_1$	2.2	1.98	2	1.98	2.64	

**Table 1.** Estimated condition number  $\mathcal{K}$  (number of PCG iterations) and the effective condition number  $\mathcal{K}_1$ 

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#### 4.2 A 3D Example

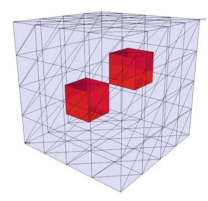
In this second example, we consider the model problem (1) in the open unit cube in 154 3D with a similar setting for the coefficient. We set  $\kappa(x) = 1$  for  $x \in \Omega_1 = (0.25, 0.5)^3$  155 or  $x \in \Omega_2 = (0.5, 0.75)^3$ , and  $\kappa(x) = \varepsilon$  for the remaining subdomain (that is, for 156  $x \in \Omega \setminus (\Omega_1 \cup \Omega_2)$ ). The domain  $\Omega$  and the subdomains just described are shown in 157 Fig. 3. The coarsest partition has mesh size  $h_0 = 2^{-2}$ , and it is set in a way so that it 158 resolves the interfaces where the coefficient has jumps.

To test the effects of the smoother, in this example we use 5 forward/backward 160 Gauss-Seidel as smoother in the multigrid preconditioner. In order to test more severe 161 jumps in the coefficients, we set the stopping criteria  $||r_k||/||r_0|| < 10^{-12}$  for the PCG 162 algorithm in this experiment.

Figure 4 shows the eigenvalue distribution of the multigrid V-cycle preconditioned system BA when  $h=2^{-5}$  (level=3) and  $\varepsilon=10^{-5}$ . As before, this figure 165 shows that there is only one small eigenvalue that even deteriorates with respect to 166 the jump in the coefficients and the mesh size.

Table 2 shows the estimated condition number  $\mathcal{K}$  (with the number of PCG 168 iterations), and the effective condition number  $\mathcal{K}_1$ . As is easily seen from the results 169 in this table, the condition number  $\mathcal K$  increases when  $\varepsilon$  decreases, i.e. the condition 170 number grows when the jump in the coefficients becomes larger. On the other hand, 171 the results in Table 2 show that the effective condition number  $\mathcal{K}_1$  remains nearly 172 uniformly bounded with respect to the mesh size and it is robust with respect to the 173 jump in the coefficient, thus confirming the result stated in Theorem 1: a PCG with 174

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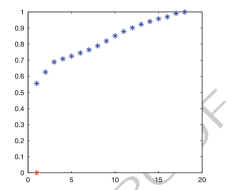


Fig. 3. 3D computational domain

Fig. 4. Eigenvalue distribution of BA

ε	levels	0	1	2	3
1	K	1.19 (8)	1.34 (11)	1.37 (11)	1.36 (11)
1	$\mathscr{K}_1$	1.16	1.26	1.31	1.29
$10^{-1}$	K	2.3 (10)	1.94(13)	1.75 (13)	1.67 (14)
10	$\mathscr{K}_1$	1.60	1.56	1.45	1.43
$10^{-3}$	K	86.01 (11)	63.07 (16)	52.67 (17)	48.19(17)
10	$\mathscr{K}_1$	2.4	2.12	1.89	1.78
$10^{-5}$	K	8.39+3 (13)	6.15e+3 (18)	5.13e+3 (19)	4.70e+3(19)
10	$\mathscr{K}_1$	2.44	2.14	1.91	1.80
$10^{-7}$	K	8.39+5 (14)	6.15e+5 (21)	5.13e+5 (23)	4.70e+5(21)
	$\mathscr{K}_1$	2.45	2.14	1.91	1.80

**Table 2.** Estimated condition number  $\mathcal{H}$  (number of PCG iterations) and effective condition number  $\mathcal{K}_1$ .

multigrid V-cycle preconditioner provides a robust, nearly optimal solver for the CR 175 approximation to (3).

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### **Domain Decomposition Methods of Stochastic PDEs**

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6

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1 Introduction 7

In conjunction with modern high performance computing systems, domain decomposition algorithms permit simulation of PDEs with extremely high resolution nu- 9 merical models. Such computational models substantially reduce discretization er- 10 rors. In realistic simulation of certain physical systems, it is however necessary to 11 consider the heterogeneities of the model parameters. Whenever sufficient statistical 12 information is available, such heterogeneities can be modeled by stochastic processes 13 (e.g. [2]). For uncertainty propagation, the traditional Monte Carlo simulation may 14 be impractical for these high resolution models. As an alternative, a domain decomposition algorithm for stochastic PDEs (SPDEs) is proposed [4] using the spectral 16 stochastic finite element method (SSFEM). The SSFEM discretization leads to a linear system with a block sparsity structure, and the size of the resulting system grows 18 rapidly with the spatial mesh resolution and the order of the stochastic dimension 19 [2]. The solution of this large-scale system constitutes a computationally challeng- 20 ing task and therefore efficient solvers are required. Extending the formulation in 21 [4], the iterative substructuring based non-overlapping domain decomposition methods are proposed to solve the large-scale linear system arising in the SSFEM. The 23 methodology is based on domain decomposition in the geometric space and a func- 24 tional decomposition in the stochastic space [4]. Firstly, we describe a primal version 25 of iterative substructuring methods of SPDEs. The method offers a straightforward 26 approach to formulate a two-level scalable preconditioner. In the proposed preconditioner, the continuity of the solution field is strictly enforced on the corner nodes of 28 the interface boundary, but weakly satisfied over the remaining interface nodes. This 29 approach naturally leads to a coarse grid connecting the subdomains globally and 30 provides a mechanism to propagate information across the subdomains which makes 31 the algorithm scalable. The proposed preconditioner may be viewed as an extension 32 of BDDC [3] for SPDEs. Secondly, a dual-primal iterative substructuring method is 33 introduced for SPDEs. In this approach, the continuity condition on the corner nodes 34 is strictly satisfied and Lagrange multipliers are used to weakly enforce the continu-

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ity on the remaining nodes of the interface boundary. This method may be construed to be an extension of FETI-DP [1] for SPDEs.

#### 2 Uncertainty Representation by Stochastic Processes

We briefly review the theories of stochastic processes, relevant to subsequent theoretical developments, by closely following [2, 4–6]. Assuming the input data (containing sufficient statistical information) permits a representation of the model parameters as stochastic processes that span the Hilbert space  $\mathcal{H}_G$ . Using Karhunen-Loeve expansion (KLE), a set of basis functions  $\{\xi_i(\theta)\}$  for the Hilbert space  $\mathcal{H}_G$  is identified. The KLE of a stochastic process  $\alpha(\mathbf{x}, \theta)$  is based on the spectral expansion of 44 its covariance function  $C_{\alpha\alpha}(\mathbf{x}, \mathbf{y})$ , and takes the following form [2]

$$\alpha(\mathbf{x}, \boldsymbol{\theta}) = \bar{\alpha}(\mathbf{x}) + \sum_{i=1}^{\infty} \sqrt{\lambda_i} \xi_i(\boldsymbol{\theta}) \phi_i(\mathbf{x}), \tag{1}$$

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where  $\bar{\alpha}(\mathbf{x})$  is the mean of the stochastic process,  $\{\xi_i(\theta)\}$  is a set of uncorrelated random variables and  $\{\lambda_i, \phi_i(\mathbf{x})\}$  are the eigenpairs of the covariance function, obtained from the following integral equation 48

$$\int_{\Omega} C_{\alpha\alpha}(\mathbf{x}, \mathbf{y}) \phi_i(\mathbf{y}) d\mathbf{y} = \lambda_i \phi_i(\mathbf{x}). \tag{2}$$

For a smooth stochastic process, only a finite number of KLE basis is sufficient to 49 represent the stochastic process. Given the covariance function of the solution is not 50 known a priori, the KLE cannot be used to represent solution process. Assuming the 51 solution process  $u(\mathbf{x}, \theta)$  belong to the Hilbert space  $\mathscr{H}_L$ , a generic basis of this space 52 can be identified using the Polynomial Chaos (PC) [2]. Consequently, the solution 53 process can be approximated as

$$u(\mathbf{x}, \boldsymbol{\theta}) = \sum_{j=0}^{N} \Psi_j(\boldsymbol{\theta}) u_j(\mathbf{x}), \tag{3}$$

where the polynomials  $\Psi_j(\theta)$  are orthogonal in the statistical sense, meaning 55  $\langle \Psi_j, \Psi_k \rangle = \langle \Psi_j^2 \rangle \delta_{jk}$  where  $\langle \cdot \rangle$  denotes the expectation operator and  $\delta_{jk}$  is the Kro- 56 necker delta, and  $u_j(\mathbf{x})$  are the PC coefficients to be determined by Galerkin projection.57

# 3 Review of Schur Complement Based Domain Decomposition Method of SPDEs

A review of the domain decomposition method for SPDEs based on [4–6] is provided in this section. For an elliptic SPDE defined on a domain  $\Omega$  with a prescribed boundary condition on  $\partial\Omega$ , the finite element discretization leads to the following linear system 63

$$\mathbf{A}(\boldsymbol{\theta})\mathbf{u}(\boldsymbol{\theta}) = \mathbf{f},\tag{4}$$

where  $\mathbf{A}(\theta)$  is the random stiffness matrix,  $\mathbf{u}(\theta)$  is the stochastic response and  $\mathbf{f}$  is 64 the applied force. The physical domain  $\Omega$  is split into  $n_s$  non-overlapping subdomains  $\{\Omega_s\}_{s=1}^{n_s}$ . For a typical subdomain  $\Omega_s$  the nodal vector  $\mathbf{u}^s(\theta)$  is partitioned 66 into interior  $\mathbf{u}_I^s(\theta)$  and interface  $\mathbf{u}_\Gamma^s(\theta)$  unknowns. This decomposition leads to the 67 following subdomain equilibrium equation 68

$$\begin{bmatrix} \mathbf{A}_{II}^{s}(\theta) & \mathbf{A}_{I\Gamma}^{s}(\theta) \\ \mathbf{A}_{\Gamma I}^{s}(\theta) & \mathbf{A}_{\Gamma \Gamma}^{s}(\theta) \end{bmatrix} \begin{pmatrix} \mathbf{u}_{I}^{s}(\theta) \\ \mathbf{u}_{\Gamma}^{s}(\theta) \end{pmatrix} = \begin{pmatrix} \mathbf{f}_{I}^{s} \\ \mathbf{f}_{\Gamma}^{s} \end{pmatrix}.$$
 (5)

Enforcing the transmission conditions and expanding the solution vector by the PCE 69 (as in Eq. (3)) and then performing Galerkin projection, we obtain the following 70 block linear systems of equations [4–6]:

$$\begin{pmatrix}
\mathbf{L} \\
\Psi_{l}(\theta) \\
\begin{bmatrix}
\mathbf{L} \\
\vdots \\
0 \\
\mathbf{R}_{I}^{T} \mathbf{A}_{II,i}^{1} \\
\vdots \\
0 \\
\mathbf{R}_{n_{s}}^{N} \mathbf{A}_{II,i}^{n_{s}} \\
\mathbf{R}_{n_{s}}^{N} \mathbf{A}_{II,i}^{n_{s}} \\
\mathbf{R}_{n_{s}}^{T} \mathbf{A}_{II,i}^{s} \\
\vdots \\
\mathbf{R}_{n_{s}}^{T} \mathbf{A}_{II,i}^{s}
\end{pmatrix}
\begin{pmatrix}
\mathbf{u}_{I,j}^{1} \\
\vdots \\
\mathbf{u}_{I,j}^{n_{s}} \\
\mathbf{u}_{I,j}
\end{pmatrix}
\Psi_{k}(\theta)$$

$$= \left\langle \left\{ \begin{array}{c}
\mathbf{f}_{I}^{1} \\
\vdots \\
\mathbf{f}_{n_{s}}^{n_{s}} \\
\mathbf{r}_{I}^{N} \\
\end{array} \right\} \Psi_{k}(\theta) \right\rangle, \quad k = 0, \dots, N. \tag{6}$$

where the restriction operator  $\mathbf{R}_s$  maps the global interface vector  $\mathbf{u}_{\Gamma}(\theta)$  to the local 73 interface unknown  $\mathbf{u}_{\Gamma}^s(\theta)$  as  $\mathbf{u}_{\Gamma}^s(\theta) = \mathbf{R}_s \mathbf{u}_{\Gamma}(\theta)$ . Compactly, Eq. (6) can be expressed 74 as

$$\begin{bmatrix} \mathcal{A}_{II}^{1} & \dots & 0 & \mathcal{A}_{I\Gamma}^{1} \mathcal{R}_{1} \\ \vdots & \ddots & \vdots & & \vdots \\ 0 & \dots & \mathcal{A}_{II}^{n_{s}} & \mathcal{A}_{I\Gamma}^{n_{s}} \mathcal{R}_{n_{s}} \\ \mathcal{R}_{1}^{T} \mathcal{A}_{\Gamma I}^{1} & \dots & \mathcal{R}_{n_{s}}^{T} \mathcal{A}_{\Gamma I}^{n_{s}} \sum_{s=1}^{n_{s}} \mathcal{R}_{s}^{T} \mathcal{A}_{\Gamma \Gamma}^{s} \mathcal{R}_{s} \end{bmatrix} \begin{pmatrix} \mathcal{U}_{I}^{1} \\ \vdots \\ \mathcal{U}_{I}^{n_{s}} \\ \mathcal{U}_{\Gamma} \end{pmatrix} = \begin{pmatrix} \mathcal{F}_{I}^{1} \\ \vdots \\ \mathcal{F}_{I}^{n_{s}} \\ \mathcal{I}_{\Gamma}^{n_{s}} \\ \mathcal{I}_{\Gamma}^{n_{s}} \end{pmatrix}, \quad (7)$$

where  $[\mathscr{A}_{\alpha\beta}^s]_{jk} = \sum_{i=0}^L \langle \Psi_i \Psi_j \Psi_k \rangle \mathbf{A}_{\alpha\beta,i}^s$ ,  $\mathscr{F}_{\alpha,k}^s = \langle \Psi_k \mathbf{f}_{\alpha}^s \rangle$ ,  $\mathscr{U}_I^m = (\mathbf{u}_{I,0}^m, \dots, \mathbf{u}_{I,N}^m)^T$  and 76

 $\mathcal{R}_s = blockdiag(\mathbf{R}_s^0, \dots, \mathbf{R}_s^N)$ . The subscripts  $\alpha$  and  $\beta$  represent the index I and  $\Gamma$ . 77 Performing Gaussian elimination in Eq. (7), we obtain the global *extended* Schur 78 complement system as

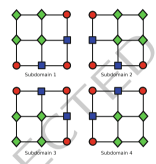
$$\mathscr{S} \mathscr{U}_{\Gamma} = \mathscr{G}_{\Gamma}, \tag{8}$$

where 
$$\mathscr{S} = \sum_{s=1}^{n_s} \mathscr{R}_s^T [\mathscr{A}_{\Gamma\Gamma}^s - \mathscr{A}_{\Gamma I}^s (\mathscr{A}_{II}^s)^{-1} \mathscr{A}_{I\Gamma}^s] \mathscr{R}_s, \ \mathscr{G}_{\Gamma} = \sum_{s=1}^{n_s} \mathscr{R}_s^T [\mathscr{F}_{\Gamma}^s - \mathscr{A}_{\Gamma:I}^s (\mathscr{A}_{II}^s)^{-1} \mathscr{F}_I^s].$$
 80

#### 4 Primal Iterative Substructuring Method of SPDEs

In this section, a two-level domain decomposition method is formulated in the context of SPDEs. The subdomain nodal vector, namely the primal variable, is partitioned into interior, remaining interface and corner nodes as schematically shown in 84 Fig. 1 [3]. Using PCE to represent the random coefficients of the system parame- 85 ters and performing Galerkin projection, lead to the following coupled deterministic 86 system

> $\left| egin{array}{c} \mathscr{A}_{ii}^{i} & \mathscr{A}_{ir}^{i} & \mathscr{A}_{ic}^{i} \ \mathscr{A}_{ri}^{s} & \mathscr{A}_{rc}^{s} & \mathscr{A}_{rc}^{s} \ \mathscr{A}_{s}^{s} & \mathscr{A}_{s}^{s} & \mathscr{A}_{s}^{s} \end{array} 
> ight| \left\langle egin{array}{c} \mathscr{U}_{i}^{s} \ \mathscr{U}_{r}^{s} \ \mathscr{U}_{s}^{s} \end{array} 
> ight\} = \left\langle egin{array}{c} \mathscr{F}_{i}^{s} \ \mathscr{F}_{s}^{s} \end{array} 
> ight\}$ (9)



**Fig. 1.** Partitioning domain nodes into: interior (♦), remaining (■) and corner (●)

Enforcing the transmission conditions along the boundary interfaces, the subdomain equilibrium equation can be written as

$$\begin{bmatrix} \mathcal{A}_{ii}^{s} & \mathcal{A}_{ir}^{s} \mathcal{B}_{r}^{s} & \mathcal{A}_{ir}^{s} \mathcal{B}_{r}^{s} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{r}^{sT} \mathcal{A}_{ri}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{r}^{sT} \mathcal{A}_{rr}^{s} \mathcal{B}_{r}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{r}^{sT} \mathcal{A}_{rc}^{s} \mathcal{B}_{c}^{s} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{ci}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{cr}^{s} \mathcal{B}_{r}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{cc}^{s} \mathcal{B}_{c}^{s} \end{bmatrix} \begin{pmatrix} \mathcal{U}_{i}^{s} \\ \mathcal{U}_{r} \\ \mathcal{U}_{c} \end{pmatrix} = \begin{pmatrix} \mathcal{F}_{i}^{s} \\ \mathcal{U}_{r}^{s} \\ \mathcal{U}_{r} \\ \mathcal{U}_{c} \end{pmatrix} = \begin{pmatrix} \mathcal{F}_{i}^{s} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{r}^{sT} \mathcal{F}_{r}^{s} \\ \mathcal{U}_{c} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{F}_{r}^{s} \\ \mathcal{E}_{c}^{sT} \mathcal{F}_{c}^{s} \end{pmatrix},$$

$$(10)$$

where  $\mathscr{B}_r^s$  and  $\mathscr{B}_c^s$  are Boolean rectangular matrices that extract the subdomain remaining interface and corner degrees of freedom from the corresponding global vec- 91 tors  $\mathcal{U}_r$  and  $\mathcal{U}_c$  as  $\mathcal{U}_r^s = \mathcal{B}_r^s \mathcal{U}_r$  and  $\mathcal{U}_c^s = \mathcal{B}_c^s \mathcal{U}_c$ . Eliminating  $\mathcal{U}_i^s$  from Eq. (10), we 92 obtain

$$\begin{bmatrix} \sum_{s=1}^{n_s} \mathcal{B}_r^{sT} \mathcal{S}_{rr}^s \mathcal{B}_r^s & \sum_{s=1}^{n_s} \mathcal{B}_r^{sT} \mathcal{S}_{rc}^s \mathcal{B}_c^s \\ \sum_{s=1}^{n_s} \mathcal{B}_c^{sT} \mathcal{S}_{cr}^s \mathcal{B}_r^s & \sum_{s=1}^{n_s} \mathcal{B}_c^{sT} \mathcal{S}_{cc}^s \mathcal{B}_c^s \end{bmatrix} \left\{ \begin{array}{c} \mathcal{U}_r \\ \mathcal{U}_c \end{array} \right\} = \left\{ \begin{array}{c} \sum_{s=1}^{n_s} \mathcal{B}_r^{sT} \mathcal{S}_r^s \\ \sum_{s=1}^{n_s} \mathcal{B}_c^{sT} \mathcal{S}_r^s \\ \sum_{s=1}^{n_s} \mathcal{B}_c^{sT} \mathcal{S}_c^s \end{array} \right\}, \quad (11)$$

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where  $\mathscr{S}_{\alpha\beta}^s = \mathscr{A}_{\alpha\beta}^s - \mathscr{A}_{\alpha i}^s [\mathscr{A}_{ii}^s]^{-1} \mathscr{A}_{i\beta}^s$  and  $\mathscr{G}_{\alpha}^s = \mathscr{F}_{\alpha}^s - \mathscr{A}_{\alpha i}^s [\mathscr{A}_{ii}^s]^{-1} \mathscr{F}_{i}^s$ . Eliminating 95  $\mathscr{U}_c$  from Eq. (11) leads to the following symmetric positive definite *reduced interface* 96 *problem* 97

$$(F_{rr} - F_{rc}[F_{cc}]^{-1}F_{cr})\mathcal{U}_r = d_r - F_{rc}[F_{cc}]^{-1}d_c, \tag{12}$$

where 
$$F_{\alpha\beta} = \sum_{s=1}^{n_s} \mathcal{B}_{\alpha}^{s} {}^T \mathcal{S}_{\alpha\beta}^s \mathcal{B}_{\beta}^s$$
 and  $d_{\alpha} = \sum_{s=1}^{n_s} \mathcal{B}_{\alpha}^s {}^T \mathcal{G}_{\alpha}^s$ .

#### 4.1 Two-Level Primal Preconditioner

The Preconditioned Conjugate Gradient Method (PCGM) can be used to solve the reduced interface problem in Eq. (12). At each iteration of the PCGM, the continuity of the solution field is enforced strictly on the corner nodes, but weakly satisfied on the remaining interface nodes. Consequently we obtain the following partially assembled Schur complement system:

$$\begin{bmatrix} \mathcal{S}_{rr}^{s} & \mathcal{S}_{rc}^{s} \mathcal{B}_{c}^{s} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{S}_{cr}^{s} \mathcal{B}_{r}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{S}_{cc}^{s} \mathcal{B}_{c}^{s} \end{bmatrix} \left\{ \begin{array}{c} \mathcal{U}_{r}^{s} \\ \mathcal{U}_{c} \end{array} \right\} = \left\{ \begin{array}{c} \mathcal{F}_{r}^{s} \\ 0 \end{array} \right\}, \tag{13}$$

where  $\mathscr{F}_r^s = \mathscr{D}_r^s \mathscr{B}_r^s \mathbf{r}_j$ , and  $\mathbf{r}_j$  is the residual of the *j*th iteration of PCGM, and  $\mathscr{D}_r^s$  is 105 a block diagonal weighting matrix which satisfies  $\sum_{s=1}^{n_s} \mathscr{B}_r^{sT} \mathscr{D}_r^s \mathscr{B}_r^s = \mathbf{I}$ . Next,  $\mathscr{U}_r^s$  can 106 be eliminated from Eq. (13) leading to the following coarse problem

$$\widetilde{F}_{cc}\mathcal{U}_c = \widetilde{d}_c,$$
 (14)

gg

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where 
$$\widetilde{F}_{cc} = \sum_{s=1}^{n_s} \mathscr{B}_c^{sT} (\mathscr{S}_{cc}^s - \mathscr{S}_{cr}^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{S}_{rc}^s) \mathscr{B}_c^s$$
 and  $\widetilde{d}_c = -\sum_{s=1}^{n_s} \mathscr{B}_c^{sT} \mathscr{S}_{cr}^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{F}_r^s$ . 108
The two-level preconditioner can be expressed as

n<sub>o</sub>

$$\mathcal{M}^{-1} = \sum_{s=1}^{n_s} \mathcal{B}_r^{sT} \mathcal{D}_r^s [\mathcal{S}_{rr}^s]^{-1} \mathcal{D}_r^s \mathcal{B}_r^s + R_0^T [\widetilde{F}_{cc}]^{-1} R_0, \tag{15}$$

where 
$$R_0 = \sum_{s=1}^{n_s} \mathcal{B}_c^{sT} \mathcal{S}_{cr}^s [\mathcal{S}_{rr}^s]^{-1} \mathcal{D}_r^s \mathcal{B}_r^s$$
.

#### 5 Dual-Primal Iterative Substructuring of SPDEs

In the dual-primal method [1], the continuity condition on the corner nodes is enforced strictly while Lagrange multipliers are used to weakly enforce the continuity
on the remaining interface. Partial assembly of the corner node unknowns leads to
the following system

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$$\begin{bmatrix} \mathcal{A}_{ii}^{s} & \mathcal{A}_{ir}^{s} & \mathcal{A}_{ic}^{s} \mathcal{B}_{c}^{s} & 0\\ \mathcal{A}_{ri}^{s} & \mathcal{A}_{ir}^{s} & \mathcal{A}_{rc}^{s} \mathcal{B}_{c}^{s} & \mathcal{B}_{r}^{s} \\ \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{ci}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{cr}^{s} & \sum_{s=1}^{n_{s}} \mathcal{B}_{c}^{sT} \mathcal{A}_{cc}^{s} \mathcal{B}_{c}^{s} & 0\\ 0 & \sum_{s=1}^{n_{s}} \mathcal{B}_{r}^{s} & 0 & 0 \end{bmatrix} \begin{pmatrix} \mathcal{U}_{i}^{s} \\ \mathcal{U}_{r}^{s} \\ \mathcal{U}_{c} \\ \Lambda \end{pmatrix} = \begin{pmatrix} \mathcal{F}_{i}^{s} \\ \mathcal{F}_{r}^{s} \\ \mathcal{W}_{c} \\ \Lambda \end{pmatrix},$$

$$(16)$$

where  $\sum_{s=1}^{n_s} \mathscr{B}_r^s \mathscr{U}_r^s = 0$  and  $\Lambda^T = \{\lambda_0, \dots, \lambda_N\}$ . The matrix  $\mathscr{B}_r^s$  is a block diagonal signed Boolean continuity operator and  $\lambda_j$  is the nodal force vector required to satisfy continuity on the remaining interface nodes. Eliminating  $\mathscr{U}_r^s$  and  $\mathscr{U}_r^s$  from 118

satisfy continuity on the remaining interface nodes. Eliminating  $\mathcal{U}_i^s$  and  $\mathcal{U}_r^s$  from 118 Eq. (16) leads to the following interface problem 119

$$\begin{bmatrix} \bar{F}_{cc} - \bar{F}_{cr} \\ \bar{F}_{rc} & \bar{F}_{rr} \end{bmatrix} \begin{Bmatrix} \mathcal{U}_c \\ \Lambda \end{Bmatrix} = \begin{Bmatrix} \bar{d}_c \\ \bar{d}_r \end{Bmatrix}, \tag{17}$$

where 120

$$\begin{split} \bar{F}_{cc} &= \sum_{s=1}^{n_s} \mathscr{B}_c^{sT} (\mathscr{S}_{cc}^s - \mathscr{S}_{cr}^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{S}_{rc}^s) \mathscr{B}_c^s, \quad \bar{F}_{cr} &= \sum_{s=1}^{n_s} \mathscr{B}_c^{sT} \mathscr{S}_{cr}^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{B}_r^{sT} \\ \bar{F}_{rc} &= \sum_{s=1}^{n_s} \mathscr{B}_r^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{S}_{rc}^s \mathscr{B}_c^s, \qquad \qquad \bar{F}_{rr} &= \sum_{s=1}^{n_s} \mathscr{B}_r^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{B}_r^{sT} \\ \bar{d}_c &= \sum_{s=1}^{n_s} \mathscr{B}_c^s (\mathscr{G}_c^s - \mathscr{S}_{cr}^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{G}_r^s), \qquad \bar{d}_r &= \sum_{s=1}^{n_s} \mathscr{B}_r^s [\mathscr{S}_{rr}^s]^{-1} \mathscr{G}_r^s \end{split}$$

Solving for  $\mathcal{U}_c$  from Eq. (17) gives the following coarse problem

$$\bar{F}_{cc}\mathcal{U}_c = (\bar{d}_c + \bar{F}_{cr}\Lambda) \tag{18}$$

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Substituting  $\mathcal{U}_c$  into Eq. (17) leads to the following symmetric positive definite Lagrange multiplier system

$$(\bar{F}_{rr} + \bar{F}_{rc}[\bar{F}_{cc}]^{-1}\bar{F}_{cr})\Lambda = \bar{d}_r - \bar{F}_{rc}[\bar{F}_{cc}]^{-1}\bar{d}_c.$$
(19)

The Lagrange multiplier system in Eq. (19) is solved using PCGM equipped with a 124 Dirichlet precondtioner defined as  $\widehat{\mathcal{M}} = \sum_{s=1}^{n_s} \mathscr{B}_r^s \mathscr{D}_r^s \mathscr{D}_r^s \mathscr{D}_r^s \mathscr{B}_r^{sT}$ . 125

#### 6 Numerical Results

For numerical illustrations, we consider the following elliptic SPDE

$$\nabla \cdot (\kappa(\mathbf{x}, \boldsymbol{\theta}) \nabla u(\mathbf{x}, \boldsymbol{\theta})) = f(\mathbf{x}), \quad \mathbf{x} \in \Omega, \tag{20}$$

$$u(\mathbf{x}, \boldsymbol{\theta}) = 0, \quad \mathbf{x} \in \partial \Omega.$$
 (21)

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The coefficient  $\kappa(\mathbf{x}, \theta)$  is modeled as a lognormal stochastic process, obtained from the underlying Gaussian process with an exponential covariance function given as

$$C_{\alpha\alpha}(\mathbf{x}, \mathbf{y}) = \sigma^2 \exp\left(-\frac{|x_1 - y_1|}{b_1} - \frac{|x_2 - y_2|}{b_2}\right).$$
 (22)

The lognormal process is approximated using four-dimensional second order PC expansion (L=15). Finite element discretization results in 375,444 elements and 131 186,925 nodes. The response is expressed using third order PCE (N=34) leading to 132 a linear system of order 6,542,375. The mean and standard deviation of the solution 133 process are shown in Fig. 2. The PCGM iteration counts for the primal and dual-primal methods for fixed problem size in the spatial domain is reported in Table 1 for 135 1st, 2nd and 3rd order of PCE. The results suggest that the methods are numerically 136 scalable with respect to number of subdomains. Table 2 shows the iteration counts of 137 the methods when we fix spatial problem size per subdomain and increase the overall 138 problem size by adding more subdomains. Again these results suggest that both the 139 methods are numerically scalable with respect to fixed problem size per subdomain. 140

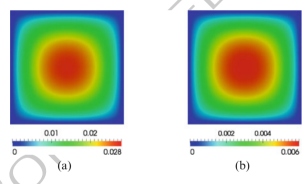


Fig. 2. The mean and standard deviation of the solution field. (a) Mean. (b) Standard deviation

Table 1. Iteration counts for fixed problem size in geometric space

Subdom	ain I	PP-DDN	Л	DP-DDM			
	1st	2nd	3rd	1 <i>st</i>	2nd	3rd	
8	11	12	12	9	9	9	
16	12	13	13	10	10	10	
32	14	14	14	11	11	11	
64	13	14	14	10	10	10	
128	14	14	14	10	10	10	
256	14	14	14	10	10	10	

**Table 2.** Iteration counts for fixed problem size per subdomain in geometric space

Subdoma	in I	PP-DDN	Л	Ι	OP-DDN	M
	1st	2nd	3rd	1 <i>st</i>	2nd	3rd
8	9	9	9	8	8	8
16	12	12	12	10	10	10
32	12	13	13	10	10	10
64	13	14	14	10	10	10
128	14	14	14	10	10	10
256	15	15	15	11	11	11

7 Conclusion 141

Primal and dual-primal domain decomposition methods are proposed to solve the 142 large-scale linear system arising from the finite element discretization of SPDEs. 143 The proposed techniques exploit a coarse grid in the geometric space which makes 144 the methods numerically scalable with respect to fixed geometric problem size, fixed 145 geometric size per subdomain and the order of PCE.

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# Improving the Convergence of Schwarz Methods for Helmholtz Equation

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1 Introduction 7

Various domain decomposition methods have been proposed for the Helmholtz equation, with the Optimized Schwarz Method (OSM) being one of them (see e.g. [7] 9 for a review of various domain decomposition methods, and [3] for the details of 10 OSM). In this paper, we focus on OSM, which is based on the idea of using approx-11 imated half-space Dirichlet-to-Neumann (DtN) maps to improve the convergence of 12 the Schwarz methods; current version of the OSM is based on polynomial approx-13 imation of the half-space DtN map. See [8] for a review of various approaches to 14 approximating the half-space DtN map (more commonly referred to as Absorbing 15 Boundary Conditions (ABCs)).

There are two approximations in the OSM that affect its convergence rate – the 17 first being the approximation of the rest of the domain as unbounded and the second 18 being the approximation of the half-space stiffness (square-root operator) as a poly- 19 nomial. In contrast with the polynomial approximation used in OSM, we utilize the 20 method of Perfectly Matched Discrete Layers (PMDL), which has close links to the 21 well-known Perfectly Matched Layers (PML) (see [1]) and the rational approximation of the square-root operator. The resulting PMDL-Schwarz method is shown to 23 converge faster than the second-order OSM. The rest of the paper contains a brief 24 review of OSM and PMDL concepts, followed by an outline of the new PMDL-Schwarz method and illustration of its effectiveness with the help of convergence 26 factor analysis and a numerical example.

Model Problem. We consider the governing equation,

$$-\frac{\partial^2 \hat{u}}{\partial x^2} - \frac{\partial^2 \hat{u}}{\partial y^2} - \omega^2 \hat{u} = \hat{f}, \quad (x, y) \in (-\infty, \infty) \times [0, L], \tag{1a}$$

$$\hat{u}(\cdot,0) = \hat{u}(\cdot,L) = 0. \tag{1b}$$

Applying Fourier Sine transform along the *y* direction, the above equation reduces 29 to a 1-D form:

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$$-\frac{\partial^2 u}{\partial x^2} - k^2 u = f, \quad x \in (-\infty, \infty), \tag{2}$$

where  $k = \sqrt{\omega^2 - k_y^2}$ ,  $k_y$  is the wavenumber along y and u, f are the Fourier symbols corresponding to  $\hat{u}$ ,  $\hat{f}$  respectively. For simplicity, we shall use the above 1-D 32 equation to discuss the main ideas in this paper, but note that the proposed method is 33 applicable to more complex equations and geometries. Also, since the focus of this 34 paper is to improve the treatment of the transmission condition at an interface, it is 35 sufficient to consider the case of two subdomains. Thus the domain is decomposed 36 into two subdomains:  $\Omega_1 \equiv (-\infty, 0)$  and  $\Omega_2 \equiv (0, \infty)$ , with the interface at x = 0.

#### 2 Optimized Schwarz Methods

Optimized Schwarz Method is a domain decomposition method that is a variant of 39 the Schwarz Alternating Method (see e.g. [7]). In the Schwarz Alternating Method, 40 the displacement and traction continuity across the artificial interface are enforced by 41 applying a mixed boundary condition of the form  $\mathcal{B}(\cdot) \equiv \partial(\cdot)/\partial \mathbf{n} + \Lambda(\cdot)$  where **n** is 42 the normal vector at the interface and the operator  $\Lambda$  is a parameter of the method. 43 The Schwarz iteration scheme for solving (2) is given by:

$$-\frac{\partial^{2} u_{1}^{j+1}}{\partial x^{2}} - k^{2} u_{1}^{j+1} = f_{1}, \quad x \in \Omega_{1}, \quad -\frac{\partial^{2} u_{2}^{j+1}}{\partial x^{2}} - k^{2} u_{2}^{j+1} = f_{2}, \quad x \in \Omega_{2}, \quad (3a)$$

$$\mathcal{B}_{1} u_{1}^{j+1} = \mathcal{B}_{1} u_{2}^{j}, \quad x = 0, \qquad \mathcal{B}_{2} u_{2}^{j+1} = \mathcal{B}_{2} u_{1}^{j+1}, \quad x = 0, \quad (3b)$$

$$\mathcal{B}_{1}(\cdot) \equiv \frac{\partial(\cdot)}{\partial \mathbf{n}_{1}} + \Lambda_{1}(\cdot), \qquad \mathcal{B}_{2}(\cdot) \equiv \frac{\partial(\cdot)}{\partial \mathbf{n}_{2}} + \Lambda_{2}(\cdot), \quad (3c)$$

$$\mathscr{B}_1 u_1^{j+1} = \mathscr{B}_1 u_2^j, \quad x = 0,$$
  $\mathscr{B}_2 u_2^{j+1} = \mathscr{B}_2 u_1^{j+1}, \quad x = 0,$  (3b)

$$\mathscr{B}_{1}(\cdot) \equiv \frac{\partial(\cdot)}{\partial \mathbf{n}_{1}} + \Lambda_{1}(\cdot), \qquad \qquad \mathscr{B}_{2}(\cdot) \equiv \frac{\partial(\cdot)}{\partial \mathbf{n}_{2}} + \Lambda_{2}(\cdot), \qquad (3c)$$

where the operators  $\Lambda_{1,2}$  are the parameters of the iteration that determine the convergence rate. The problem now reduces to choosing the parameters that lead to 46 optimal convergence of the iteration scheme. The parameters are commonly chosen 47 to be scalars but they can be operators that are optimized for convergence [3]. The 48 dependence of the convergence on the choice of parameters is better understood by 49 looking at the convergence factor  $\rho$ , which is defined as

$$\left| \hat{e}_i^{j+1} \right| = \rho \left| \hat{e}_i^{j} \right|, \tag{4}$$

where  $\hat{e}_i^J = |u - u_i^J|$  is the error in the solution in subdomain i at iteration j. Thus, 51 after one cycle of iteration, the error in solution reduces by  $\rho$  and the iterative scheme 52 converges to a solution as long as  $\rho < 1$ .

For the Schwarz method in (3), the convergence factor can be shown to be (see 54 for e.g. [3]) 55

$$\rho = \left| \left( \frac{\Lambda_1 - \mathcal{K}_2}{\Lambda_1 + \mathcal{K}_1} \right) \left( \frac{\Lambda_2 - \mathcal{K}_1}{\Lambda_2 + \mathcal{K}_2} \right) \right|, \tag{5}$$

where  $\mathscr{K}_1$  and  $\mathscr{K}_2$  are the DtN maps of the subdomains  $\Omega_1$  and  $\Omega_2$  respectively. 56 It is clear from (5) that the iterative scheme does not converge (because  $\rho = 1$ ) for 57

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a pure Neumann ( $\Lambda_i = 0$ ) or Dirichlet ( $\Lambda_i = \infty$ ) interface condition. Also, if  $\Lambda_1 = 58$  $\mathcal{K}_2$  or  $\Lambda_2 = \mathcal{K}_1$ , then  $\rho = 0$  and the Schwarz iterative scheme converges in two 59 iterations, i.e., the parameters are optimal. However, DtN maps are known only for 60 special cases and even then are usually non-local operators that are expensive to 61 compute accurately. Thus we look for local approximations to these DtN maps that 62 are accurate and computationally efficient.

Optimized Schwarz Methods [3] essentially approximate the DtN map of the subdomains by polynomial approximations of the DtN map of an unbounded domain, 65 e.g. the second-order OSM makes the approximation

$$\mathcal{K}_1 = -i\sqrt{\omega^2 - k_y^2} \approx p + q k_y^2, \tag{6}$$

where p, q are parameters that are found by minimizing the convergence factor over 67 the entire range of allowed vertical wavenumbers  $k_y$ . Note that there are other variants 68 of OSM based on zeroth-order approximation; in this paper, we focus on the best 69 available OSM, namely the second-order OSM.

#### 3 A Schwarz Method with Improved Convergence

It appears to us that OSM uses polynomial approximation for reasons of imple-72 mentability. A better approximation would be to use higher order rational approx- 73 imations, which have been investigated extensively in the context of Absorbing 74 Boundary Conditions (ABCs); it is now possible to implement these resulting ABCs 75 and can also be used in the context of Schwarz methods. In this paper, we propose 76 the use of a rational approximation in a recent ABC called Perfectly Matched Dis- 77 crete Layers (formerly known as Continued Fraction ABCs – see [4]) instead of the 78 polynomial approximation in (6).

The rational approximation corresponding to PMDL is given by:

$$\mathcal{X}_1 = -i\sqrt{\omega^2 - k_y^2} \approx \mathcal{S}_n^{pmdl}, \qquad (7)$$

where 81

$$\mathcal{S}_{n}^{pmdl} = p_{n} - \frac{q_{n}^{2}}{p_{n} + \left(p_{n-1} + \frac{q_{n-1}^{2}}{p_{n-1} + \left(p_{n-2} - \frac{q_{n-2}^{2}}{p_{n-2} + (\dots)}\right)\right)},$$
(8)

$$p_{i} = \frac{1}{4L_{i}} \left( 4 - k^{2} L_{i}^{2} \right)$$

$$q_{i} = \frac{1}{4L_{i}} \left( -4 - k^{2} L_{i}^{2} \right)$$

$$i = 1 \dots n.$$
(9)

where  $L_i$  are the parameters that determine the accuracy of the approximation.

The error in the approximation (7) is typically analyzed through the so-called 83 reflection coefficient, which has been shown to be (for details, see [4])

$$R = \prod_{i=1}^{n} \left| \frac{\mathcal{K}_1 - p_i}{\mathcal{K}_1 + p_i} \right|^2. \tag{10}$$

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If R = 0, then the approximation is exact, and the deviation from zero indicates magnitude of error in the approximation; smaller the value of R, better the approximation. 86 So from (10) and (9), it is clear that the accuracy of proposed approximation hinges 87 on the choice of  $L_i$ .

In general,  $L_i$  are chosen to be complex or imaginary to better approximate the 89 DtN map for propagating wave modes and are chosen to be real when evanescent 90 modes are important. While the parameters  $L_i$  can be optimized using the concepts 91 discussed in [5], in this paper we choose  $L_i$  based on the OSM parameters (see 92 Sect. 4).

**Implementation of PMDL.** While the rational form of the PMDL approxima- 94 tion in (8) is useful for analysis, the following matrix form proves to be useful for 95 implementation:

$$\begin{bmatrix}
\mathcal{S}_{n}^{pmdl}u_{b} \\
0 \\
0 \\
\vdots \\
0
\end{bmatrix} = \begin{bmatrix}
p_{1} & q_{1} & 0 & \cdots & 0 \\
q_{1} & p_{1} + p_{2} & q_{2} & & 0 \\
0 & \ddots & \ddots & \ddots & 0 \\
\vdots & & q_{n-1} & p_{n-1} + p_{n} & q_{n} \\
0 & \cdots & 0 & q_{n} & p_{n}
\end{bmatrix} \begin{bmatrix}
u_{b} \\
u_{a,1} \\
u_{a,2} \\
\vdots \\
u_{a,n-1}
\end{bmatrix}, (11)$$

where  $p_i, q_i$  are given by (9) and  $u_{a,i}$  are auxiliary variables that are introduced to 97 facilitate the implementation and have no direct physical relevance to the problem. 98 The equivalence between (8) and (11) can be easily seen by eliminating the auxil- 99 iary dof  $u_{a,i}$  from (11) to recover (8). The matrix form of PMDL enables an easy 100 implementation of the rational approximation as a simple tri-diagonal matrix.

PMDL, a link between Rational ABCs and Perfectly Matched Layers. While 102 the matrix form of the PMDL approximation in (11) is based on the rational approximation in (8), it is intimately linked to impedance-preserving discretization of PML 104 proposed in [4]. Unlike PML, the impedance is preserved even after discretization 105 and thus the approximation is named perfectly matched discrete layers, PMDL. This 106 link is substantial in that it provides a way to derive and easily implement PMDL ap- 107 proximations for more complex cases such as corners [4] and anisotropic elasticity 108 [**6**].

The ease of implementation of PMDL is in fact the impetus behind proposed 110 method. As implied by (10), the accuracy of approximation can be easily increased 111 by adding auxiliary variables, which is equivalent to adding lines of nodes parallel 112 to the interface. As will be shown in Sect. 4, addition of just one auxiliary variable, 113 which has minimal increase in computational cost per iteration, significantly reduces 114 the convergence factor and the number of iterations needed.

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**Implementation of the PMDL-Schwarz method.** The proposed 116 PMDL-Schwarz method is essentially the Schwarz Alternating method with the 117 operator  $\Lambda_1$  chosen to be the DtN map obtained using PMDL, i.e.,  $\Lambda_1 = \mathcal{S}_n^{pmdl}$  118 where  $\mathcal{S}_n^{pmdl}$  is given by (11). Thus the interface condition in (3) for  $\Omega_1$  can be 119 written as

$$\frac{\partial}{\partial \mathbf{n}_1} (u_1^{j+1} - u_2^j) + \mathcal{S}_n^{pmdl} (u_1^{j+1} - u_2^j) = 0.$$
 (12)

Substituting (11) in (12), we get the PMDL-Schwarz formulation as

$$\begin{bmatrix} \frac{\partial u_{1}^{j+1}}{\partial \mathbf{n}_{1}} \\ 0 \\ \vdots \\ 0 \end{bmatrix} + \begin{bmatrix} p_{1} & q_{1} & 0 & \cdots & 0 \\ q_{1} & p_{1} + p_{2} & q_{2} & & 0 \\ 0 & \ddots & \ddots & \ddots & 0 \\ \vdots & & q_{n-1} & p_{n-1} + p_{n} & q_{n} \\ 0 & \cdots & 0 & q_{n} & p_{n} \end{bmatrix} \begin{bmatrix} u_{1}^{j+1} \\ u_{a,1} \\ u_{a,2} \\ \vdots \\ u_{a,n-1} \end{bmatrix} = \begin{bmatrix} -\frac{\partial u_{2}^{j}}{\partial \mathbf{n}_{2}} + p_{1}u_{2}^{j} \\ q_{1}u_{2}^{j} \\ 0 \\ \vdots \\ 0 \end{bmatrix}.$$

$$(13)$$

Note that the formulation of the interface condition for  $\Omega_2$  can be derived in an identical manner and hence is not repeated here.

#### 4 Comparison Between OSM and PMDL-Schwarz Methods

In this section, we compare the performance of OSM and PMDL-Schwarz method both theoretically (using convergence factors) and in a numerical simulation involving multiple domains and closed boundaries.

Convergence Factors: Consider the stiffness approximation of the second-order 128 OSM (see [3]), 129

$$\mathscr{S}_{osm} = \frac{ab - \omega^2}{a+b} + \frac{1}{a+b}k_y^2. \tag{14}$$

Substituting  $\Lambda_1 = \Lambda_2 = \mathscr{S}_{osm}$  in (5), we get the convergence factor of OSM to be

$$\rho_{osm} = \left| \frac{ab + k_y^2 - \omega^2 + i(a+b)\sqrt{\omega^2 - k_y^2}}{ab + k_y^2 - \omega^2 - i(a+b)\sqrt{\omega^2 - k_y^2}} \right|^2.$$

To compare, we use a two-layer PMDL-Schwarz method with  $L_1=2/a$ , and 131  $L_2=2/b$ , where a,b are the OSM parameters in (14). The stiffness approximation 132 of the two-layer PMDL-Schwarz method is then given by

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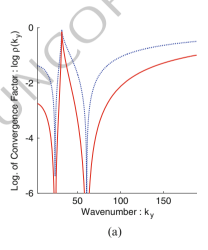
$$\begin{split} \mathscr{S}_n^{pmdl} &= p_2 - \frac{q_2^2}{p_2 + p_1} \,, \\ p_2 &= \frac{1}{L_2} - \frac{(\omega^2 - k_y^2) L_2}{4} \,, \quad q_2 = -\frac{1}{L_2} - \frac{(\omega^2 - k_y^2) L_2}{4} \,, \\ p_1 &= \frac{1}{L_1} - \frac{(\omega^2 - k_y^2) L_1}{4} \,. \end{split}$$

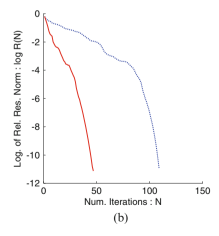
Substituting  $\Lambda_1 = \Lambda_2 = \mathscr{S}_n^{pmdl}$  in (5), we get the convergence factor of PMDL- 134 Schwarz that can be simplified to

$$\rho_{pmdl} = \left( \left| \frac{ab + k_y^2 - \omega^2 + i(a+b)\sqrt{\omega^2 - k_y^2}}{ab + k_y^2 - \omega^2 - i(a+b)\sqrt{\omega^2 - k_y^2}} \right|^2 \right)^2.$$

Clearly  $\rho_{pmdl} = \rho_{osm}^2$ , and so the parameters of PMDL-Schwarz are chosen such that its convergence factor is the square of that of OSM and the method performs uniformly better over the entire range of wavenumbers  $k_v$ .

It is easy to numerically verify the above result for the model problem (1a), with 139 the domain  $\Omega$  decomposed into two semi-infinite layers. We take  $a=20.741\,\mathrm{i}$  and 140 b=47.071 to be the OSM parameters as these were shown in [3] to be optimal 141 over the allowed wavenumber range  $k_y \in [\pi,60\pi]$ . Figure 1a compares the convergence factors of OSM and PMDL-Schwarz method (with  $L_1=2/a$  and  $L_2=2/b$ ) 143 and shows clearly that the proposed method performs better over the entire range of 144 wavenumbers for a slightly increased computational cost (there is only one auxiliary 145 variable introduced, which is similar to one line of nodes in 2-D).





**Fig. 1.** Comparison between OSM (*dotted line*) and PMDL-Schwarz method (*solid line*) . (a) Convergence Factor. (b) Convergence of Solution

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Numerical Example: In this example, Eq. (1a) is solved on a square domain 147  $(\Omega \equiv [0,1] \times [0,1])$  with  $\omega = 10\pi$  and a point source f = 1/2 is applied at (0,0.5). 148 Homogeneous Neumann boundary condition is applied on the left (x = 0), Dirichlet 149 condition at the top (y = 1) and bottom (y = 0), and an ABC on the right (x = 1). 150 The computational domain is discretized using 60 bilinear finite elements along each 151 direction. The domain is decomposed into nine subdomains with three subdomains 152 along each dimension. The convergence plot is shown in Fig. 1b. As expected, the 153 PMDL-Schwarz method converges twice as fast as the conventional OSM.

5 Discussion 155

We proposed a Schwarz method for Helmholtz equation based on the concepts of 156 perfectly matched discrete layers (PMDL), a recently developed absorbing boundary 157 condition that is related to the higher order rational approximations and the Per- 158 fectly Matched Layers. By examining the convergence factor and with the help of a 159 numerical example, PMDL-Schwarz method is shown to converge faster than existing Optimized Schwarz Methods. Although not treated in this paper, it is important 161 to mention that the PMDL is not just limited to the Helmholtz equation, but also 162 to more complicated vector equations such as the elastic and electromagnetic wave 163 equations. Thus, it is expected that the PMDL-Schwarz method would provide accelerated convergence in frequency domain computations in these contexts. Further- 165 more, as Waveform Relaxation Method in time domain share similar ideas with OSM 166 (see e.g. [2]), PMDL ideas can also be used to improve the convergence of existing 167 waveform relaxation methods. These extensions are subjects of ongoing research.

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## A Domain Decomposition Solver for the Discontinuous 2 **Enrichment Method for the Helmholtz Equation**

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1 Introduction 8

The discontinuous enrichment method (DEM) [4] for the Helmholtz equation approximates the solution as a sum of a piecewise polynomial continuous function and 10 element-wise supported plane waves [5]. A weak continuity of the plane wave part 11 is enforced using Lagrange multipliers. The plane wave enrichment improves the accuracy of solutions considerably. In the mid-frequency range, severalfold savings in 13 terms of degrees of freedom over comparable higher order polynomial discretizations 14 have been observed, which translates into even larger savings in compute time [6, 9]. 15 The partition of unity method [8] and the ultra weak variational formulation [1] also 16 employ plane waves in the construction of discretizations. It was shown recently in 17 [10] that DEM without the polynomial field is computationally more efficient than 18 these methods.

So far only direct solution methods have been used with DEM. This paper de- 20 scribes an iterative domain decomposition method which will enable to solve much 21 larger problems with DEM. The method is a generalization of the FETI-H version [3] 22 of the FETI method [2] and the domain decomposition method for DEM without the 23 polynomial part described in [7]. It is based on a non-overlapping decomposition of 24 the domain into subdomains. On the subdomain interfaces Lagrange multipliers are 25 introduced to enforce the continuity of the polynomial part strongly and the con- 26 tinuity of the enrichment weakly. An efficient iterative solution procedure with a 27 two-level preconditioner resembling that of the FETI-H method is constructed for 28 the Lagrange multipliers on the interfaces between the subdomains.

#### 2 Problem Formulation and Discretization

The solution  $u \in H^1(\Omega)$  of a Helmholtz problem modeling acoustic scattering from 31 a rigid obstacle, for example, satisfies the equations

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$$-\Delta u - k^2 u = f \qquad \text{in } \Omega$$

$$\frac{\partial u}{\partial v} = g_1 \qquad \text{on } \Sigma_1$$

$$\frac{\partial u}{\partial v} = iku + g_2 \qquad \text{on } \Sigma_2,$$
(1)

where k is the wavenumber,  $\Sigma_1$  is the boundary of a sound-hard scatterer,  $\Sigma_2$  is the 33 far-field boundary, and  $\nu$  denotes the unit outward normal.

Let the domain  $\Omega$  be split into  $n_e$  elements,  $\Omega = \bigcup_{e=1}^{n_e} \Omega_e$ . In DEM, the solution 35 is sought in the form  $u = u^P + u^E$ , where  $u^P$  is a standard continuous piecewise 36 polynomial finite element function, and  $u^E$  is an enrichment function discontinuous 37 across element interfaces. A weak inter-element continuity of the solution is enforced 38 by Lagrange multipliers  $\lambda^E$ . The following hybrid variational formulation is used: 39 Find  $u \in \mathcal{V}$  and  $\lambda^E \in \mathcal{W}^E$  such that

$$a(u,v)+b(\lambda^E,v)=r(v) \qquad \forall v\in \mathscr{V}$$
  $b(\mu^E,u) \qquad =0 \qquad \forall \mu^E\in \mathscr{W}^E.$ 

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The forms a, b, and r are defined by

$$\begin{split} a(u,v) &= \int_{\Omega} (\nabla u \cdot \nabla v - k^2 u v) d\Omega - \int_{\Sigma_2} i k u v \, d\Gamma, \\ b(\lambda^E,v) &= \sum_{e=1}^{n_e} \sum_{e'=1}^{e-1} \int_{\Gamma_{e,e'}} \lambda^E \left( v_{\Omega_e'} - v|_{\Omega_e} \right) d\Gamma, \quad \text{and} \\ r(v) &= \int_{\Omega} f v \, d\Omega + \int_{\Sigma_1} g_1 v \, d\Gamma + \int_{\Sigma_2} g_2 v \, d\Gamma, \end{split}$$

where  $\Gamma_{e,e'} = \partial \Omega_e \cap \partial \Omega_{e'}$ . For the considered discretization, the space  $\mathscr{V}$  consists of 44 functions of the form  $u = u^P + u^E$ , where  $u^E$  is a superposition of  $n_\theta$  planar waves, 45 i.e.

$$u^{E}(\mathbf{x}) = \sum_{p=1}^{n_{\theta}} e^{ik\theta_{p} \cdot \mathbf{x}} u_{e,p}^{E}, \qquad \mathbf{x} \in \Omega_{e}.$$

In two dimensions,  $\theta_p = (\cos \vartheta_p, \sin \vartheta_p)^T$ ,  $\vartheta_p = 2\pi(p-1)/n_\theta$ ,  $p = 1, \dots, n_\theta$ . The 48 Lagrange multipliers space  $\mathscr{W}^E$  is then chosen using functions of the form 49

$$\lambda^{E}(\mathbf{x}) = \sum_{p=1}^{n_{\lambda}} e^{ik\eta_{p}\tau_{e,e'}\cdot\mathbf{x}} \lambda_{e,e',p}, \qquad \mathbf{x} \in \Gamma_{e,e'},$$

where  $\tau_{e,e'}$  is a unit tangent vector and  $\eta_p$  is a scalar. This choice yields a family 51 of quadrilateral elements, denoted by  $Q - n_\theta - n_\lambda$ . In particular, the elements Q-8-2 52 and Q-16-4 used in the numerical experiments in this paper use  $\eta_1 = -\eta_2 = 0.5$  and 53  $\{\eta_p\}_{p=1}^4 = \{\pm 0.2, \pm 0.75\}$ , respectively. For details on stability, implementation, and 54 accuracy, the reader is referred to [5, 6].

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#### 3 Domain Decomposition Formulation

The elements are divided into  $n^d$  disjoint subsets  $E^j$  defining subdomains  $\Omega^j$  such 57 that  $\bar{\Omega}^j = \bigcup_{e \in E^j} \bar{\Omega}_e$ . Subdomain problems are given by regularized bilinear forms 58

$$\begin{split} \widetilde{a}^{j}(u^{j}, v^{j}) &= \int_{\Omega^{j}} (\nabla u^{j} \cdot \nabla v^{j} - k^{2} u^{j} v^{j}) d\Omega - \int_{\Sigma_{2} \cap \partial \Omega^{j}} iku^{j} v^{j} d\Gamma \\ &- \gamma \sum_{\substack{j'=1\\j' \neq j}}^{n^{d}} \int_{\Gamma^{j,j'}} s^{j,j'} iku^{j} v^{j} d\Gamma, \end{split}$$

where  $\Gamma^{j,j'}=\partial\Omega^j\cap\partial\Omega^{j'}$ . The functions  $u^j$  and  $v^j$  belong to the restriction of  $\mathscr V$  60 into  $\Omega^j$  and the last term ensures the subdomain problems cannot be singular; for 61 details see [7]. The coefficients  $s^{j,j'}$  are chosen so that the regularization terms cancel 62 out for a continuous function. The continuity of the polynomial part of the solution 63  $\tilde u^P=\sum_{j=1}^{n^d}u^{P,j}$  across the subdomain interfaces is enforced using a Lagrange multiplier 64

 $\lambda^{P}$ . For this purpose, a bilinear form

$$c(\lambda^{P}, \tilde{v}) = \sum_{j=1}^{n^{d}} \sum_{j'=1}^{j-1} \sum_{l} \lambda_{j,j',l}^{P} \left( \tilde{v}^{P}|_{\Omega^{j'}} - \tilde{v}^{P}|_{\Omega^{j}} \right) \left( \mathbf{x}_{j,j',l} \right)$$

$$66$$

is defined, where  $\mathbf{x}_{j,j',l}$  is the location of the lth mesh node on  $\Gamma^{j,j'}$ . The mesh nodes are given by the Lagrange interpolation points of the piecewise polynomial functions. 68 The domain decomposition formulation then reads:

Find 
$$\tilde{u} \in \widetilde{\mathscr{V}}$$
,  $\lambda^E$ , and  $\lambda^P$  such that

$$\widetilde{a}(\widetilde{u},\widetilde{v}) + b(\lambda^{E},\widetilde{v}) + c(\lambda^{P},\widetilde{v}) = \widetilde{r}(\widetilde{v}) \qquad \forall \widetilde{v} \in \widetilde{\mathscr{V}}$$

$$b(\mu^{E},\widetilde{u}) = 0 \qquad \forall \mu^{E} \in \mathscr{W}^{E}$$

$$c(\mu^{P},\widetilde{u}) = 0 \qquad \forall \mu^{P} \in \mathscr{W}^{P},$$
(2)

where  $\widetilde{\mathscr{V}}$  is spanned by  $\sum\limits_{j=1}^{n^d} v_j$ ,  $\widetilde{a}(\widetilde{u},\widetilde{v}) = \sum\limits_{j=1}^{n^d} a^j(u^j,v^j)$ , and  $\widetilde{r}$  is the sum of subdomain 71 contributions of r.

#### 4 Linear Systems and Condensations

The formulation (2) leads to the saddle point system of linear equations

$$\begin{pmatrix} \mathbf{r}A^{PP} & \mathbf{r}A^{PE} & 0 & \mathbf{C}^{PL} \\ \mathbf{r}A^{EP} & \mathbf{r}A^{EE} & \mathbf{B}^{EL} & 0 \\ 0 & \mathbf{B}^{LE} & 0 & 0 \\ \mathbf{C}^{LP} & 0 & 0 & 0 \end{pmatrix} \begin{pmatrix} \mathbf{u}^{P} \\ \mathbf{u}^{E} \\ \lambda^{P} \\ \lambda^{P} \end{pmatrix} = \begin{pmatrix} \mathbf{r}^{P} \\ \mathbf{r}^{E} \\ 0 \\ 0 \end{pmatrix}, \tag{3}$$

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where the superscripts P, E, and L refer to the polynomial part, the enrichment 75 part, and the Lagrange multiplier, respectively, and  $u^P$ ,  $u^E$ ,  $\lambda^E$ ,  $\lambda^P$  are vectors of the 76 subdomain-by-subdomain polynomial degrees of freedom (depicted by black dots 77 in Fig. 1), the element-by-element enrichment degrees of freedom (magenta arrows), 78 the enrichment element-to-element continuity Lagrange multipliers (red arrows), and 79 the polynomial subdomain-to-subdomain continuity Lagrange multipliers (black ar- 80 rows), respectively. The enrichment unknowns  $u^E$  can be condensed out on the element level (Fig. 1 top and left) to obtain

$$\begin{pmatrix} \mathbf{r}\bar{A} \ \mathbf{B}^T \ \mathbf{\bar{C}}^T \\ \mathbf{\bar{B}} \ \mathbf{\bar{D}} \ 0 \\ \mathbf{\bar{C}} \ 0 \ 0 \end{pmatrix} \begin{pmatrix} \mathbf{u}^P \\ \lambda^E \\ \lambda^P \end{pmatrix} = \begin{pmatrix} \mathbf{\bar{r}} \\ \bar{\mu} \\ 0 \end{pmatrix}, \tag{4}$$

where 83

$$\mathbf{r}\bar{A} = \mathbf{r}A^{PP} - \mathbf{r}A^{PE} (\mathbf{r}A^{EE})^{-1} \mathbf{r}A^{EP}, \ \bar{\mathbf{B}} = -\mathbf{B}^{LE} (\mathbf{r}A^{EE})^{-1} \mathbf{r}A^{EP}, 
\bar{\mathbf{C}} = \mathbf{C}^{LP}, \qquad \bar{\mathbf{D}} = -\mathbf{B}^{LE} (\mathbf{r}A^{EE})^{-1} \mathbf{B}^{EL}, 
\bar{\mathbf{r}} = \mathbf{r}^P - \mathbf{r}A^{PE} (\mathbf{r}A^{EE})^{-1} \mathbf{r}^E, \qquad \bar{\mu} = -\mathbf{B}^{LE} (\mathbf{r}A^{EE})^{-1} \mathbf{r}^E.$$

The enrichment Lagrange multipliers  $\lambda^E$  can be divided into two parts—those on 85 the boundaries between the subdomains and those inside the subdomains, denoted by 86 the subscript B and I, respectively. The system (4) can then be written in the block 87 form

$$\begin{pmatrix} \bar{\mathbf{r}}A & \bar{\mathbf{B}_{II}}^T & \bar{\mathbf{B}_{BB}}^T & \bar{\mathbf{C}}^T \\ \bar{\mathbf{B}}_{II} & \bar{\mathbf{D}}_{II} & \bar{\mathbf{D}}_{IB} & 0 \\ \bar{\mathbf{B}}_{BB} & \bar{\mathbf{D}}_{BI} & \bar{\mathbf{D}}_{BB} & 0 \\ \bar{\mathbf{C}} & 0 & 0 & 0 \end{pmatrix} \begin{pmatrix} \boldsymbol{u}^P \\ \lambda_I^E \\ \lambda_B^E \\ \lambda^P \end{pmatrix} = \begin{pmatrix} \bar{\mathbf{r}} \\ \bar{\mu}_I \\ \bar{\mu}_B \\ 0 \end{pmatrix}.$$

Finally, the elimination on the subdomain level of the unknowns  $\mathbf{u}^P$  and the interior

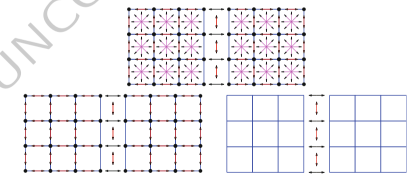


Fig. 1.  $2 \times 1$  domain decomposition of a DEM discretization with bilinear polynomials and O-8-2 elements resulting in the system (3) (top); variables left after condensation of enrichment dofs (4) (*left*); and elimination of the subdomain interior dofs (5) (*right*)

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enrichment Lagrange multipliers  $\lambda_I^E$  gives the Schur complement system (cf. Fig. 1 91 right)

 $\mathbf{F} \begin{pmatrix} \lambda_B^E \\ \lambda^P \end{pmatrix} = \mathbf{b}. \tag{5}$ 

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It is noted that the matrix  $\mathbf{F}$  is a sum of subdomain matrices. Once the Lagrange multipliers  $\lambda_B^E$  and  $\lambda^P$  have been solved from (5), the rest of the unknowns is recovered by post-processing, first to obtain  $\mathbf{u}^P$  and  $\lambda_L^E$ , then to obtain  $\mathbf{u}^E$ .

#### 5 Preconditioning

The system (5) is solved efficiently using a Krylov iterative method with a two-level 97 preconditioner which is a generalization of those described in [3, 7]. 98

Here, the subdomain preconditioners are based on the bilinear forms

$$\hat{a}^{j}(u^{j}, v^{j}) = \int_{\Omega^{j}} (\nabla u^{j} \cdot \nabla v^{j} - k^{2}u^{j}v^{j}) d\Omega - \int_{\partial\Omega^{j} \setminus \Sigma_{1}} iku^{j}v^{j} d\Gamma,$$

$$\hat{b}^{j}(\lambda^{E}, v^{j}) = \sum_{e \in E^{j}} \sum_{e'=e+1}^{n_{e}} \int_{\Gamma_{e,e'}} \lambda^{E}v|_{\Omega_{e}} d\Gamma - \sum_{e \in E^{j}} \sum_{e'=1}^{e-1} \int_{\Gamma_{e,e'}} \lambda^{E}v|_{\Omega_{e}} d\Gamma, \text{ and}$$

$$\hat{c}^{j}(\lambda^{P}, v^{j}) = \sum_{i'=j+1}^{n^{d}} \sum_{l} \lambda_{j,j',l}^{P} v^{P}|_{\Omega^{j}} (\mathbf{x}_{j,j',l}) - \sum_{i'=1}^{j-1} \sum_{l} \lambda_{j,j',l}^{P} v^{P}|_{\Omega^{j}} (\mathbf{x}_{j,j',l}).$$

Repeating the same steps described above for obtaining  ${\bf F}$  in (5) but with matrices based on  $\hat a^j$ , and restricting the resulting matrix to the unknowns corresponding to the interfaces of the subdomain  $\Omega^j$ , a matrix denoted by  ${\bf F}^j$  is obtained (cf. [7]). An additive subdomain-by-subdomain preconditioner is then defined by

$$\mathbf{K} = \sum_{i=1}^{n^d} \left( \mathbf{R}^j \right)^T \left( \mathbf{F}^j \right)^{-1} \mathbf{R}^j,$$
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where  $\mathbf{R}^j$  is the restriction on the interfaces associated with  $\Omega^j$ . Linear systems with 106  $\mathbf{F}^j$  can be solved efficiently using an LU decomposition.

The system (5) is solved iteratively on the orthogonal complement of a coarse space spanned by the columns of a matrix  $\mathbf{Q}$  (cf. [3, 7]). A projector to the orthogonal complement of the coarse space is given by

$$\mathbf{P} = \mathbf{I} - \mathbf{Q}(\mathbf{Q}^T \mathbf{F} \mathbf{Q})^{-1} \mathbf{Q}^T \mathbf{F}.$$

The solution  $\lambda = [\lambda_B^E, \lambda^P]^T$  of (5) can be decomposed into two parts  $\lambda = \lambda^0 + \mathbf{P}\lambda^1$ , 112 where  $\lambda^0 = \mathbf{Q}(\mathbf{Q}^T\mathbf{F}\mathbf{Q})^{-1}\mathbf{Q}^T\mathbf{b}$  and  $\lambda^1$  satisfies

$$\mathbf{P}^T \mathbf{F} \lambda^1 = \mathbf{P}^T \mathbf{b}.$$

Including the preconditioner **K** leads to the following equation

$$\mathbf{P}\mathbf{K}\mathbf{P}^{T}\mathbf{F}\lambda^{1} = \mathbf{P}\mathbf{K}\mathbf{F}\lambda^{1} = \mathbf{P}\mathbf{K}\mathbf{P}^{T}\mathbf{b},$$

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which is solved by GMRES.

The coarse space is based on plane waves propagating in  $n_q$  uniformly distributed 116 directions. Each set of  $n_a$  plane waves are supported by one subdomain interface  $\Gamma^{j,j'}$  117 and their normal derivatives on the interface are approximated using an  $L^2$ -projection 118 into the space of Lagrange multipliers giving rise to  $n_a$  columns of **Q**. Currently, the 119 coarse space acts only on the interface enrichment Lagrange multipliers  $\lambda_R^E$ . The 120 maximum dimension of the coarse space is  $n_q n_i$ , where  $n_i$  is the number of nonzero measure interfaces  $\Gamma^{j,j'}$ . A **QR** factorization is used to remove nearly linearly dependent vectors. More details are given in Sect. 3.4 of [7].

#### 6 Numerical Results

The model problem considered here is given by (1) with the computational domain 125  $\Omega = \{\mathbf{x} \in \mathbb{R}^2 : 1 < ||\mathbf{x}|| < 2\}$ , and the boundaries  $\Gamma_1 = \{\mathbf{x} \in \mathbb{R}^2 : ||\mathbf{x}|| = 1\}$  and  $\Gamma_2 = 126$   $\{\mathbf{x} \in \mathbb{R}^2 : ||\mathbf{x}|| = 2\}$ . The right-hand side function and the boundary functions are 127 chosen as 128

$$f(\mathbf{x}) = (-\Delta - k^2)(x_1^2 + x_2^2) = -4 - k^2(x_1^2 - x_2^2),$$

$$g_1(\mathbf{x}) = -\frac{\partial e^{-ikx_1}}{\partial v} + \frac{\partial (x_1^2 + x_2^2)}{\partial v} = -ikx_1e^{ikx_1} - 2(x_1^2 + x_2^2), \text{ and}$$

$$g_2(\mathbf{x}) = \frac{\partial (x_1^2 + x_2^2)}{\partial v} - ik(x_1^2 + x_2^2) = (1 - ik)(x_1^2 + x_2^2).$$

The solution is a sum of that given by the scattering of the plane wave  $e^{-ikx_1}$  by 130 a sound-hard disk inside  $\Gamma_1$  and the polynomial  $x_1^2 + x_2^2$ . Two wavenumbers, k = 131 $8\pi$  and  $16\pi$  are considered, in which case the diameter of the scatterer is 8 and 132 16 wavelengths, respectively. The solution at  $k = 16\pi$  is shown in Fig. 2. Meshes 133 of  $96 \times 8$   $(k = 8\pi)$  and  $192 \times 16$   $(k = 16\pi)$  elements result in two elements per 134 wavelength in the radial direction. 135

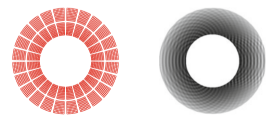


Fig. 2. The  $24 \times 2$  domain decomposition for the  $192 \times 16$  mesh (*left*) and the real part of the solution at  $k = 16\pi$  (right)

t1 1 t1.2 t1 3 t1 4 t1.5

t1 7 t1.8 t1.9 t1 10 +1 11

149

<b>Table 1.</b> Results for the $96 \times 8$ mesh with the wavenumber $k = 8\pi$ .							
	12 x 1 subdomains	24 x 2 subdomains					

		12 x 1 subdomains			24 x 2 subdomains			
			$n_q = 0$	$n_q = 8$		$n_q = 0$	$n_q = 8$	
poly	enrich	N	iter.	iter.	N	iter.	iter.	error
$\overline{Q_1}$	none	108	49		336	213		0.683405
$Q_2$	none	204	33		624	195		0.141341
none	Q-8-2	192	35	31	576	163	7	0.438341
$Q_1$	Q-8-2	300	34	31	912	184	28	0.004677
$Q_2$	Q-8-2	396	34	31	1200	206	48	0.004472
none	Q-16-4	384	35	30	1152	151	39	0.019767
$Q_1$	Q-16-4	492	36	31	1488	160	54	0.000024
$Q_2$	Q-16-4	588	36	31	1776	176	73	0.000013

**Table 2.** Results for the  $192 \times 16$  mesh with the wavenumber  $k = 16\pi$ .

		12 :	12 x 1 subdomains			x 2 subd	omains		t2.1
			$n_q = 0$	$n_q = 16$		$n_q = 0$	$n_q = 16$		t2.2
poly	enrich	N	iter.	iter.	N	iter.	iter.	error	t2.3
$\overline{Q_1}$	none	204	79		624	350		0.568750	t2.4
$Q_2$	none	396	40		1200	368		0.174451	t2.5
none	Q-8-2	384	44	34	1152	264	16	0.478914	t2.6
$Q_1$	Q-8-2	588	42	34	1776	281	31	0.007441	t2.7
$Q_2$	Q-8-2	780	42	34	2352	295	56	0.007826	t2.8
none	Q-16-4	768	42	33	2304	233	42	0.021694	t2.9
$Q_1$	Q-16-4	972	42	35	2928	238	52	0.000011	t2.10
$Q_2$	Q-16-4	1164	42	33	3504	253	123	0.000010	t2.11

Bilinear  $(Q_1)$  and biquadratic  $(Q_2)$  bases are used for the polynomial part  $u^P$ . 136 Q-8-2 and Q-16-4 elements are used for the enrichment  $u^E$  and its Lagrange multipliers  $\lambda^E$ . The domain is decomposed into  $12 \times 1$  and  $24 \times 2$  subdomains (Fig. 2). 138 The GMRES iterations are terminated once the norm of the residual is reduced by 139 10<sup>-8</sup>. Tables 1 and 2 summarize the performance results obtained for various element 140 types. In these tables, N is the size of the system (5), i.e. the number of Lagrange multipliers enforcing continuity between subdomains. The error is the relative  $l_2$  error of 142 the averaged nodal values with respect to the analytical solution of the problem.

The errors in the last column of Tables 1 and 2 clearly show the benefit of discretizations with both polynomial and enrichment fields for this problem. The com- 145 bined discretizations increase the accuracy by at least two orders of magnitude. The 146 iteration counts without a coarse space  $(n_a = 0)$  are roughly the same for all discretizations and not quite satisfactory for the 24 × 2 decomposition. However, these 148 are reduced substantially when the coarse space is added.

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# Domain Decomposition Methods for the Helmholtz Equation: A Numerical Investigation

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1 Introduction 7

We are interested in solving the Helmholtz equation

$$\begin{cases} -\triangle u(x, y, z) - k^{2}(x, y, z) \ u(x, y, z) = g(x, y, z), \ (x, y, z) \in \Omega, \\ \partial_{n} u(x, y, z) - \mathbf{i}k(x, y, z) \ u(x, y, z) = 0, \ (x, y, z) \in \partial \Omega, \end{cases}$$
(1)

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where  $k := 2\pi f/c$  is the wavenumber with frequency  $f \in \mathbf{R}$  and c := c(x,y,z) is 9 the velocity of the medium, which varies in space. The geophysical model SEG- 10 SALT is used as a benchmark problem on which we will test some existing domain 11 decomposition methods in this paper. In this model, the domain  $\Omega$  is defined as 12  $(0,13,520) \times (0,13,520) \times (0,4,200) \,\mathrm{m}^3$ , the velocity is described as piecewise constants on  $676 \times 676 \times 210$  cells and varies from 1,500 to 4,500 m/s, and the source 14 g is a Dirac function at the point (6,000,6,760,10).

To discretize the problem (1) on a coarser mesh, the velocity is sub-sampled to 16 less number of cells such that every cell has a constant velocity and contains one 17 or more mesh elements. Then the problem (1) is discretized with Q1 finite elements 18 (i.e. trilinear local basis functions on brick elements).

We first test the direct solver  $A \setminus b$  in Matlab; the results are listed in Table 1 where 20 nw is the number of wavelength along the x-direction at the lowest velocity. At f=2, 21 the direct solver runs out of memory after 6 h on a computer with 64 GB of memory. The inefficiency in both memory and time of the direct solver for large scale 23 problems calls for cheaper iterative methods. For a review of current iterative methods for the Helmholtz equation, we refer to [6]. In this work, we focus on domain 25 decomposition methods which are easily parallelized.

#### 2 Overview of Some Existing Methods

Due to the indefiniteness of the Helmholtz equation, the classical Schwarz method 28 with Dirichlet transmission conditions fails to converge. As a remedy, [5] introduced 29

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$\overline{f}$	1/4	1/2	1	2
nw	2.25	4.5	9	18
mesh	$24 \times 24 \times 8$	$48\times48\times16$	$96 \times 96 \times 32$	$192\times192\times64$
CPU	1.28s	27.51s	829.91s	> 6h

**Table 1.** Test of the direct solver (backslash in Matlab)

first-order absorbing transmission conditions to replace the Dirichlet transmission 30 conditions. This type of interface condition was also adopted in [7] to regularize 31 subdomain problems. More general local transmission conditions of zero or second 32 order were proposed and analyzed in [10, 11] with parameters optimized for accelerating convergence. More advanced and even non-local transmission conditions can 34 be used, see [3, 12, 18], and also [2, 13] in this volume. In this paper, however, we 35 will restrict ourselves to local transmission conditions.

Another remedy is to modify the usual coarse problem, which probably originated from the multigrid context, first suggested by Achi Brandt and presented in 38 [19]. In their paper [7], Farhat et al. used plane waves on the interface as basis of the 39 coarse space. The idea turns out to be very successful and was followed by Farhat 40 et al. [8], Kimn and Sarkis [15], and Li and Tu [17], and will also be used for the 41 optimized Schwarz methods in this paper. Note that, however, the coarse problem 42 does not change the underlying subdomain problems.

In the following paragraphs, we will give a brief introduction to these methods at 44 the (almost) continuous level. 45

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#### 2.1 The Non-overlapping Methods

We partition the domain into non-overlapping subdomains denoted by  $\overline{\Omega} := \cup_i \overline{\Omega_i}$ , 47 and we call the set of points shared by more than two subdomains (or shared by two 48 subdomains and the outer boundary  $\partial \Omega$ ) corners. In three dimensions, this includes 49 vertices and edges. We call all the points shared by exactly two subdomains the 50 interface  $\Gamma$ , and in particular a connected component of the interface shared by  $\Omega_i$  51 and  $\Omega_j$  is called interface segment  $\Gamma_{ij}$ .

If we know the Neumann, Dirichlet or Robin data (denoted by  $\lambda$ ) of the exact 53 solution on the interface, then we can recover the exact solution from the corresponding boundary value problems defined on subdomains (as long as they are well-posed) with *continuous constraints at corners*. Since on every subdomain there is a 56 recovered solution that gives Dirichlet, Neumann or Robin traces on the interface, 57 we expect for each interface segment  $\Gamma_{ij}$  the traces from  $\Omega_i$  and  $\Omega_j$  to be equal. The 58 above process indeed sets up an equation, denoted by  $F\lambda = d$ , for the interface data 59  $\lambda$  of the exact solution. For the Helmholtz equation, an additional coarse problem is 60 introduced such that  $(I - FQ(Q^*FQ)^{-1}Q^*)F\lambda = (I - FQ(Q^*FQ)^{-1}Q^*)d$  is solved, 61 where the columns of Q are traces of plane waves on the interface.

From the above point of view, we summarize some existing non-overlapping 63 domain decomposition methods in Table 2. The (first-order) absorbing boundary data 64

t2.1 t2.2 t2.3 t2.4

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is defined as  $\lambda := \partial_{\bf n} u - {\bf i} k u$ . The lumped preconditioner is the stiffness submatrix 65  $A_{\Gamma\Gamma}$  corresponding to the interface. The first three methods share interface data (up 66 to a sign for the normal derivative) on their common interface segments, and are 67 therefore one-field methods. This is in contrast to the last method, since optimized 68 Schwarz methods have two sets of unknowns on each interface segment, and thus 69 belong to the class of two-field methods. Note also that we do not have suitable 70 preconditioners for the last two methods, which can be a subject for future study.

**Table 2.** The non-overlapping methods

Algorithms	Unknowns	Matching	Precond.
FETI-DPH ([8])	Neumann	Dirichlet	DtN/lumped
BDDC-H ([17])	Dirichlet	Neumann	NtD
FETI-H ([7])	Absorbing	Dirichlet	(none)
Optimized Schwarz ([10])	two-field Robin	two-field Robin	(none)

### 2.2 The Overlapping Methods

We partition the domain into overlapping subdomains. We will use the *substructured* 73 *form*<sup>3</sup> as for the non-overlapping methods in Sect. 2.1. Note that in an overlapping 74 setting, subdomains can not share the same interface data, since the interfaces are 75 in different locations, and therefore all overlapping methods are in some sense two 76 field methods, like the non-overlapping optimized Schwarz methods. The interface 77 data used (both as unknowns and matching conditions) and related references are: 78 Dirichlet [16], absorbing [4, 15], Neumann [14], Robin [9]. A coarse problem as in 79 Sect. 2.1 is adopted but without corner constraints.

# **3 Numerical Experiments**

All the experiments were done in Matlab with sequential codes. We use GMRES with 82 zero initial guess to solve the substructured systems until the relative residual is less 83 than  $10^{-6}$  or the maximum iteration number is attained. The domain is partitioned in 84 a Cartesian way. If we vary the mesh size, then the velocity in (1) is sub-sampled on 85 the coarsest mesh of  $24 \times 24 \times 8$ . 86

We introduce the following acronyms:

FL/FD: FETI-DPH with the lumped/DtN preconditioner
FH: FETI-H with corner constraints
00/O2: non-overlapping optimized Schwarz of zero/second order
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<sup>&</sup>lt;sup>3</sup> Though most of the overlapping methods in the literature are not in this form, we found by numerical experiments it may be cheaper in both time and memory.

OD/ON/OR: overlapping method with Dirichlet/Neumann/absorbing data OO0/OO2: overlapping optimized Schwarz of zero/second order

For the overlapping methods, the overlapping region has a thickness of two mesh 93 elements and the matching conditions are imposed on faces, edges and vertices, re- 94 spectively, without repeats on any degrees of freedom. Due to the absence of relevant 95 results, the parameters for the optimized Schwarz methods are not respecting overlapping (except OO0), coarse problem and medium heterogeneity. The plane waves 97 used are along six directions that are normal to the x-y, y-z and z-x planes, respec- 98 tively.

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We found that all the methods outperform the direct solver in CPU time (see 100 Table 1) on the  $96 \times 96 \times 32$  mesh. We are interested in how the convergence of these methods depends on the frequency f in (1), the mesh size h, the partition  $N_x \times N_y \times N_z$  102 or the subdomain size H and medium heterogeneity. At f = 1 the domain contains 103 nine wavelength along the x-direction, which corresponds to the problem on the unit 104 cube with the wavenumber  $18\pi$ .

In the following tables, the numbers outside/inside parentheses are the iteration 106 numbers with/without plane waves, respectively, and a bar is used instead of 200 107 when the maximum iteration number is reached. We use e/w to represent the number 108 of elements per wavelength at the lowest velocity. The smallest iteration numbers 109 among the non-overlapping methods and those among the overlapping methods are 110 in bold. Note that for the FETI-DPH method with DtN preconditioner the amount 111 of work per iteration is about 1.5 times that for the others, and construction of the 112 preconditioner also leads to double LU factorizations in the setup stage.

In Tables 3 and 4, we increase the frequency with fh or  $f^3h^2$  [1] kept constant.

f FL FD FH 00O2 OD OR ON OO<sub>2</sub> 000 t3.1 partition  $3 \times 3 \times 1$ 6 (15) 4 (8) 9 (15) 15 (21) 8 (14) 8 (20) 8 (12) 9 (20) 7 (15) **6** (14) 15 (30) 9 (12) 18 (33) 29 (34) 19 (20) 23 (34) 12 (15) 24 (37) 12 (17) **11** (13) t3.3 **20** (23) 75 (93) 43 (48) 25 (25) 51 (58) 17 (17) 57 (66) 22 (25) **14** (15) t3.4 partition scaling with mesh: H/h = 8 (see also the first row for  $f = \frac{1}{4}$ ) 10 (73) 17 (71) 10 (50) 14 (73) 11 (33) 21 (103) **8** (55) **8** (51) 9 (183) 7 (-) 21 (-) 12 (-) 27 (-) **15** (74) 152 (-) 16 (-) **15** (-) 11 (-) t3.6 partition scaling with mesh: H/h = 16 (see also the second row for  $f = \frac{1}{2}$ ) 1 39 (127) 32 (103) 74 (-) 59 (113) **27** (39) 76 (171) 26 (38) 114 (-) 26 (53) **22** (32) t3.7

**Table 3.** Dependence on the frequency (fh = constant)

We see that more iterations are usually needed for larger frequency except in the 115 middle of Table 4.

In Table 5, the frequency is fixed and the mesh is refined. From the table, the 117 iteration numbers with plane waves almost remain constant. 118

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**Table 4.** Dependence on the frequency  $(f^3h^2 = \text{constant})$ 

f	FL	FD	FH	O0	O2	OD	OR	ON	OO0	OO2	t4.1			
	partition $3 \times 3 \times 1$ (see also the first row in Table 3 for $f = 0.25$ )													
0.40	12(25)	<b>6</b> (11)	14(25)	30 (33)	18(21)	18 (29)	11(14)	19 (32)	<b>9</b> (15)	<b>9</b> (13)	t4.2			
0.63	27 (41)	<b>11</b> (15)	33 (49)	37 (42)	25 (26)	38 (46)	16(17)	39 (50)	15 (20)	<b>13</b> (14)	t4.3			
	partitio	n scaling	with mes	sh: H/h =	8(see al	so the firs	st row in	Table 4 f	for $f = 0$	.25)				
0.40	7 (36)	<b>5</b> (23)	10 (54)	15 (58)	9 (40)	12 (60)	10(29)	13 (73)	7 (40)	7 (40)	t4.4			
0.63	7 (127	<b>5</b> (100)	9 (149)	14 (156)	8(112)	14 (160)	11 (65)	20(-)	7 (123)	<b>7</b> (117)	t4.5			
	par	tition sca	ling with	mesh: H	/h = 16 (	see also t	he first r	ow for f	= 0.40)					
0.63	15 (89)	<b>8</b> (53)	18 (119)	43 (125)	18 (75)	33(113)	16 (35)	36 (112)	<b>13</b> (75)	<b>13</b> (75)	t4.6			
			Table 5	Depende	ence on tl	he mesh s	ize (f =	$\frac{1}{2}$						
	<b>Table 5.</b> Dependence on the mesh size $(f = \frac{1}{4})$													
e/w	FL	FD	FH	O0	O2	OD	OR	ON	OO0	OO2	t5.1			
		partition	$13 \times 3 \times$	1 (see alse	o the first	t row in T	able 4 fo	or $e/w =$	10)					
20	10 (19	5 (9)	13 (20)	17 (26)	9 (17)	14 (28)	11 (15	) 13 (27	8 (16)	<b>6</b> (16)	t5.2			
40				21 (32)			. //				t5.3			
		partition	nH/h=1	8 (see also	o the first	row in T	able 4 fo	or $e/w =$	10)					
20	7 (21	5 (12)	10 (32)	14 (47)	8 (32)	10 (46)	9 (25	) 10 (44	7 (29)	<b>6</b> (30)	t5.4			
40	6 (19	<b>4</b> (13)	9 (36)	14 (92)	7 (63)	9 (90)	9 (46	9 (91	) 7 (56)	6 (59)	t5.5			
		par	tition $H/$	h = 16 (s	ee also th	ne first rov	w for $e/1$	w = 20)						
40	11 (34	6 (15)	14 (47)	17 (60)	10 (38)	15 (63)	12 (28	) 13 (52	7 (33)	7 (35)	t5.6			
					-									

Next, we compare the iteration numbers for different partitions with both the 119 frequency and the mesh size fixed in Table 6. One can see that with plane waves

**Table 6.** Dependence on the partition

	FL	FD	FH	00	O2	OD	OR	ON	OO0	002
$\frac{H}{H_0}$		$f = \frac{1}{2}$	, mesh an	d velocity	48 × 48	× 16 and	$H_0$ part	ition $3 \times 3$	× 1	
1	15 (30)	<b>9</b> (12)	18 (33)	28 (35)	19 (21)	22 (34)	12 (15)	23 (37)	<b>11</b> (17)	<b>11</b> (14)
$\frac{1}{2}$	8 (47)	<b>5</b> (30)	10 (73)	16 (72)	9 (51)	14 (75)	11 (34)	21 (105)	8 (62)	<b>7</b> (57)
$\frac{1}{4}$	4(22)	<b>4</b> (21)	7 (48)	10 (95)	7 (72)	7 (97)	8 (52)	11 (-)	6 (83)	<b>5</b> (78)
		f = 1	, mesh an	d velocity	$96 \times 96$	$\times$ 32 and	$H_0$ part	ition $3 \times 3$	$\times 1$	
1	46 (54)	<b>22</b> (24)	79 (97)	45 (49)	26 (26)	54 (61)	17 (18)	60 (69)	22 (26)	<b>15</b> (16)
$\frac{1}{2}$	43 (133)	35 (109)	82 (-)	63 (117)	<b>28</b> (40)	82 (176)	27 (39)	136 (-)	28 (56)	<b>24</b> (34)
$\frac{I}{A}$	10 (184)	8 (-)	14 (-)	26 (-)	16 (40)	32 (-)	17 (-)	- (-)	25 (-)	22 (-)
$N_{\chi}$		f =	1, mesh a	nd velocity	$96 \times 96$	$6 \times 32$ an	d partitio	on $N_x \times 1$	< 1	
8	117 (125)	79 (75)	171 (184)	66 (70)	<b>28</b> (28)	94 (99)	<b>23</b> (24)	100 (104)	51 (46)	<b>23</b> (25)
16	184 (-)	192 (199)	- (-)	131 (137)	<b>45</b> (47)	- (-)	46 (47)	- (-)	72 (81)	<b>43</b> (45)
32	- (-)	- (-)	- (-)	172(173)	<b>87</b> (90)	- (-)	86 (90)	182 (88)	148 (136)	<b>84</b> (87)

using more subdomains can both increase and decrease the iteration numbers. It is 121 interesting that for the strip-wise partition only the methods based on transmission 122 conditions (O0, O2, OR, OO0 and OO2) work reliably, though with substantial iter- 123 ation numbers, and the plane waves do not help much.

Last, we study the influence of the heterogeneity in the velocity. The experiments 125 are carried out on artificial velocity models to have high contrasts. The frequency is 126 fixed as  $f = \frac{1}{2}$ . The lowest velocity is fixed as  $c_{\min} = 1,500$  and different levels of highest velocity  $c_{\rm max}=\rho c_{\rm min}$  are considered. It can be seen from Table 7 that the 128 iteration numbers vary only little.

Table 7. Influence of medium heterogeneity

								_		
ρ	FL	FD	FH	00	O2	OD	OR	ON	OO0	002
	r	nesh 48 ×	48 × 16, pa	artition 8	$\times$ 1 $\times$ 1 a	$\operatorname{nd} c = c_{\operatorname{m}}$	$_{\mathrm{in}},c_{\mathrm{max}}$ O	n subdom	ains	
1	58 (76)	37 (46)	83 (94)	60 (64)	<b>28</b> (29)	70 (81)	27 (26)	69 (79)	37 (44)	<b>24</b> (24)
$10^{2}$	28 (36)	42 (58)	30 (37)	37 (55)	<b>26</b> (31)	37 (53)	27 (29)	63 (75)	15 (26)	<b>13</b> (22)
$10^{4}$	32 (36)	49 (58)	33 (37)	45 (55)	<b>26</b> (31)	43 (53)	29 (30)	71 (75)	19 (26)	<b>17</b> (22)
			as	above exc	ept parti	tion $6 \times 6$	$\times 2$			
1	9 (90)	7 (62)	12 (124)	26 (79)	15 (39)	18 (97)	14 (35)	22 (117)	<b>10</b> (46)	12 (34)
$10^{2}$	12 (59)	<b>10</b> (104)	17 (51)	25 (78)	15 (46)	17 (67)	12 (34)	29 (100)	8 (42)	9 (37)
$10^{4}$	14 (58)	<b>11</b> (104)	19 (51)	27 (79)	17 (47)	19 (68)	12 (34)	33 (100)	8 (42)	10 (37)
	m	esh $48 \times 4$	$8 \times 16$ , par	tition 1 ×	$8 \times 1$ and	$d c = c_{\min}$	$c_{\max}$ on	$8\times1\times1$	cells	
1	70 (81)	40 (50)	105 (114)	73 (75)	<b>27</b> (28)	74 (80)	28 (27)	62 (66)	34 (37)	<b>24</b> (24)
$10^{2}$	51 (59)	30 (34)	69 (84)	58 (67)	<b>26</b> (28)	56 (67)	<b>23</b> (26)	51 (59)	26 (28)	<b>23</b> (26)
$10^{4}$	52 (59)	30 (34)		58 (67)						
	m	esh $84 \times 8$	$4 \times 24$ , par	tition 6 ×	$6 \times 2$ and	$d c = c_{\min}$	$c_{\max}$ on	$7 \times 7 \times 3$	cells	
1	12 (105)	8 (65)	16 (144)	34 (96)	19 (41)	24 (121)	17 (37)	25 (111)	<b>12</b> (46)	15 (34)
$10^{2}$	10 (68)	7 (34)	14 (107)	29 (109)	17 (48)	26 (111)	13 (45)	21 (106)	<b>11</b> (47)	12 (40)
$10^{4}$	11 (68)	7 (34)		31 (109)						
	m	esh $48 \times 4$	$8 \times 16$ , par							` ′
$10^{2}$	7 (16)	5 (10)	10 (21)	14 (61)	9 (41)	14 (60)	11 (37)	12 (59)	7 (35)	8 (38)
$10^{4}$	8 (15)	6 (9)	11 (20)	12 (67)	8 (46)	14 (67)	15 (61)	25 (86)	8 (39)	
		. ,	as	above exc	ept parti	tion $3 \times 3$	×1	. ,	` ,	` /
1	22 (38)	<b>10</b> (16)	26 (45)					27 (36)	15 (21)	<b>12</b> (14)
$10^{2}$	11 (17)	<b>6</b> (8)	15 (20)					16 (42)		
	12 (17)	<b>6</b> (8)	16 (21)		. ,			17 (52)		
	-2 (17)	0 (0)	10 (21)	10 (07)	> (2 T)	10 (10)	10 (31)	1 (32)	0 (20)	/ (22)

4 Conclusions

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For the SEG-SALT model on the cube domain, we get the following conclusions: 131 among the non-overlapping methods, the FETI-DPH method with DtN precondi- 132 tioner performs best in terms of iteration numbers. Among the overlapping methods, 133

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the optimized Schwarz method of second order is usually the best. With a fixed number of plane waves, all the methods can slow down for larger frequencies on properly 135 refined meshes. They also deteriorate for fixed frequency on finer meshes, unless 136 when using plane waves and more subdomains. A smaller subdomain size can both 137 increase and decrease the iteration numbers, and the experiments indicate the existence of some optimal choice. For strip-wise partitions, only the methods based on 139 transmission conditions work well, and plane waves do not help much. We also find 140 the performance of all the method is only little affected by the heterogeneity in the 141 velocity we considered, but other kinds of heterogeneity still need to be investigated. 142

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# **Stable BETI Methods in Electromagnetics**

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Summary. In this paper we present a stable boundary element tearing and interconnecting 6 domain decomposition method for the parallel solution of the electromagnetic wave equation 7 with piecewise constant wave numbers. In particular we consider stable boundary integral 8 formulations and generalized Robin type transmission conditions to ensure unique solvability 9 of the local subproblems. Numerical results confirm the robustness of the proposed approach. 10

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1 Introduction 11

The application of standard finite and boundary element tearing and interconnecting 12 domain decomposition methods [4, 5] may fail in the case of the acoustic or electromagnetic wave equation due to a possible occurence of spurious modes which 14 are related to local Dirichlet or Neumann boundary value problems. For the acoustic wave equation we have introduced in [9, 10] a boundary element tearing and 16 interconnecting domain decomposition approach which is stable for all local wave 17 numbers. The aim of this paper is to extend these results when considering the electromagnetic wave equation. Although the general concept is rather similar in both 19 cases, the numerical analysis of boundary integral equations and boundary element 20 methods for the Maxwell system requires advanced techniques, in particular appro- 21 priate space splitting approaches. For the definition of Sobolev spaces which are 22 related to the Maxwell equation, see, e.g., [2], for the analysis of Maxwell boundary 23 integral equations, see, for example, [7], and for related boundary element methods, 24 see, e.g., [1].

## 2 Formulation of the Domain Decomposition Approach

As a model problem we consider the Neumann boundary value problem of the elec- 27 tromagnetic wave equation 28

$$\mathbf{curl}\,\mathbf{curl}\,\mathbf{U}(x) - [k(x)]^2\mathbf{U}(x) = \mathbf{0} \qquad \text{for } x \in \Omega, \tag{1}$$

$$\gamma_N \mathbf{U}(x) := \mathbf{curl} \, \mathbf{U}(x) \times \mathbf{n} = \mathbf{f}(x) \quad \text{for } x \in \Gamma,$$
 (2)

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where  $\Omega \subset \mathbb{R}^3$  is a Lipschitz polyhedron with boundary  $\Gamma = \partial \Omega$ . We assume that 29 the boundary value problems (1) and (2) admits a unique solution. Since the wave 30 number k(x) is assumed to be piecewise constant, i.e.  $k(x) = k_i$  for  $x \in \Omega_i$ , instead of (1) and (2) we consider local boundary value problems to find  $U_i = U_{|\Omega_i|}$  satisfying

**curl curl U**<sub>i</sub>
$$(x) - k_i^2 \mathbf{U}_i(x) = \mathbf{0}$$
 for  $x \in \Omega_i$ ,  $\gamma_N \mathbf{U}_i(x) = \mathbf{g}(x)$  for  $x \in \Gamma_i \cap \Gamma$ 

with respect to a non-overlapping domain decomposition

$$\overline{\Omega} = \bigcup_{i=1}^p \overline{\Omega}_i, \quad \Omega_i \cap \Omega_j = \emptyset \quad \text{for } i \neq j, \quad \Gamma_i = \partial \Omega_i,$$
 35

together with the transmission or interface boundary conditions

$$\gamma_{D,i}\mathbf{U}_i(x) = \gamma_{D,i}\mathbf{U}_i(x) \quad \text{for } x \in \Gamma_{ij} = \Gamma_i \cap \Gamma_j,$$
 (3)

$$\begin{split} \gamma_{D,i}\mathbf{U}_i(x) &= \gamma_{D,j}\mathbf{U}_j(x) &\quad \text{for } x \in \varGamma_{ij} = \varGamma_i \cap \varGamma_j, \\ \gamma_{N,i}\mathbf{U}_i(x) &+ \gamma_{N,j}\mathbf{U}_j(x) = \mathbf{0} &\quad \text{for } x \in \varGamma_{ij}, \end{split} \tag{4} \end{split}$$
 where the Dirichlet trace operator is given by 
$$\gamma_D\mathbf{U} = \mathbf{n} \times (\mathbf{U}_{|\varGamma} \times \mathbf{n}).$$

$$\gamma_D \mathbf{U} = \mathbf{n} \times (\mathbf{U}_{|\Gamma} \times \mathbf{n}).$$
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Since the local Dirichlet or Neumann boundary value problems may exhibit spu- 39 rious modes, instead of the Neumann transmission condition in (4) we consider a 40 generalized Robin interface condition 41

$$\gamma_{N,i}\mathbf{U}_i(x) + \gamma_{N,j}\mathbf{U}_j(x) + i\eta_{ij}\mathsf{R}_{ij}[\gamma_{D,i}\mathbf{U}_i(x) - \gamma_{D,j}\mathbf{U}_j(x)] = \mathbf{0} \quad \text{for } x \in \Gamma_{ij}, i < j. \quad (5)$$

The operators  $R_{ij}$  are assumed to be strictly positive, i.e.  $\langle R_{ij} \mathbf{u}, \mathbf{u} \rangle_{\Gamma_{ij}} > 0$  for all  $\mathbf{u} \in {}^{42}$  $\mathbf{H}_{\perp}^{-1/2}(\operatorname{curl}_{\Gamma}, \varGamma_{ij})$ , and  $\eta_{ij} \in \mathbb{R} \backslash \{0\}$ . We define 43

$$(\mathsf{R}_{i}u_{\mid \Gamma_{i}})(x) := (\mathsf{R}_{ij}u_{\mid \Gamma_{ij}})(x) \quad \text{for } x \in \Gamma_{ij}$$

$$\eta_{i}(x) := \begin{cases}
\eta_{ij} & \text{for } x \in \Gamma_{ij}, \ i < j, \\
-\eta_{ij} & \text{for } x \in \Gamma_{ij}, \ i > j, \\
0 & \text{for } x \in \Gamma_{i} \cap \Gamma,
\end{cases}$$

where we assume that  $\eta_i(x)$  for  $x \in \Gamma_i$  does not change its sign, see also [9]. In 47 this case we can ensure unique solvability [11] of the local Robin boundary value 48 problems

$$\mathbf{curl} \, \mathbf{curl} \, \mathbf{U}_i(x) - k_i^2 \mathbf{U}_i(x) = \mathbf{0} \qquad \text{for } x \in \Omega_i,$$

$$\gamma_N \mathbf{U}_i(x) + i \eta_i \mathsf{R} \gamma_D \mathbf{U}_i(x) = \mathbf{g}(x) \qquad \text{for } x \in \Gamma_i \cap \Gamma.$$

$$\tag{7}$$

$$\gamma_N \mathbf{U}_i(x) + i \eta_i \mathsf{R} \gamma_D \mathbf{U}_i(x) = \mathbf{g}(x) \quad \text{for } x \in \Gamma_i \cap \Gamma.$$
 (7)

For the solution of local Dirichlet and Robin boundary value problems we will apply 50 boundary element methods which are based on the use of the Stratton-Chu represen- 51 tation formula for  $x \in \Omega$ , see [3], 52

58

$$\mathbf{U}(x) = \mathbf{\Psi}_k^M(\gamma_D \mathbf{U})(x) + \mathbf{\Psi}_k^A(\gamma_N \mathbf{U})(x) + \frac{1}{k^2} \mathbf{grad} \, \mathbf{\Psi}_k^S \mathrm{div}_{\Gamma}(\gamma_N \mathbf{U})(x).$$

Here, 53

$$\Psi_k^A(\lambda)(x) := \int_{\Gamma} g_k(x,y)\lambda(y)ds_y \quad \text{for } x \notin \Gamma, \quad g_k(x,y) = \frac{1}{4\pi} \frac{e^{ik|x-y|}}{|x-y|},$$

is the vector-valued single layer potential with the fundamental solution of the 54 Helmholtz equation, and 55

$$\Psi_k^M(\lambda)(x) := \mathbf{curl}\,\Psi_k^A(\lambda \times \mathbf{n})(x) \quad \text{for } x \notin \Gamma$$

is the Maxwell double layer potential. In addition,

$$\Psi_k^V(\lambda)(x) := \int_{\Gamma} g_k(x, y) \lambda(y) ds_y \quad \text{for } x \notin \Gamma$$

is the scalar single layer potential. By introducing the Maxwell single layer potential 57

$$\Psi_k^S(\lambda)(x) := \Psi_k^A(\lambda)(x) + \frac{1}{k^2} \operatorname{grad} \Psi_k^S \operatorname{div}_{\Gamma}(\lambda)(x) \quad \text{for } x \notin \Gamma,$$

we can write the Straton-Chu representation formula as

$$\mathbf{U}(x) = \mathbf{\Psi}_k^M(\gamma_D \mathbf{U}(x)) + \mathbf{\Psi}_k^S(\gamma_N \mathbf{U}(x)) \quad \text{for } x \in \Omega.$$
 (8)

The application of the Maxwell trace operators gives the boundary integral equations 59 [7, 11]

$$\gamma_N \mathbf{U} = \mathsf{N}_k(\gamma_D \mathbf{U}) + (\frac{1}{2}I + \mathsf{B}_k)(\gamma_N \mathbf{U}), 
\gamma_D \mathbf{U} = (\frac{1}{2}I + \mathsf{C}_k)(\gamma_D \mathbf{U}) + \mathsf{S}_k(\gamma_N \mathbf{U}).$$
(9)

Now we are in a position to derive different approaches to solve local boundary 61 value problems with generalized Robin boundary conditions. Here we consider an 62 approach which is based on the use of the Steklov–Poincaré operator 63

$$T_k = N + (\frac{1}{2}I + B_k)S_k^{-1}(\frac{1}{2}I + C_k) = S_k^{-1}(\frac{1}{2}I + C_k)$$
 (10)

which requires the invertibility of the single layer operator  $S_k$ . Since  $S_k$  is not invertible for all wave numbers k, instead of (10) we consider a system of boundary integral equations to find  $\mathbf{u} \in \mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma)$  and  $\mathbf{t} \in \mathbf{H}_{\parallel}^{-1/2}(\operatorname{curl}_{\Gamma}, \Gamma)$  such that

$$\begin{pmatrix} \mathsf{N}_k + i\eta \,\mathsf{R} \, \frac{1}{2}I + \mathsf{B}_k \\ -\frac{1}{2}I + \mathsf{C}_k \, \mathsf{S}_k \end{pmatrix} \begin{pmatrix} \mathbf{u} \\ \mathbf{t} \end{pmatrix} = \begin{pmatrix} \mathbf{g} \\ \mathbf{0} \end{pmatrix} \tag{11}$$

is satisfied. The unique solvability of (11) follows from a generalized Garding in- 67 equality

$$\begin{split} \operatorname{Re}\left(\left\langle \left( \mathbf{N}_{k} + i\eta \mathbf{R} \ \frac{1}{2}I + \mathbf{B}_{k} \right) \left( \mathbf{u} \atop \mathbf{t} \right), \left( \frac{\mathscr{Y}\mathbf{u}}{\mathscr{X}\mathbf{t}} \right) \right\rangle_{\Gamma} + C((\mathbf{u}, \mathbf{t}), (\mathbf{u}, \mathbf{t})) \right) \\ & \geq c \left( \left\| \mathbf{u} \right\|_{\mathbf{H}_{\perp}^{-1/2}(\operatorname{curl}_{\Gamma}, \Gamma)}^{2} + \left\| \mathbf{t} \right\|_{\mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma)}^{2} \right) \end{split}$$

for some appropriate bijective operators  $\mathscr{X}$  and  $\mathscr{Y}$ , and from injectivity which is in 69 fact related to the unique solvability of the local Robin boundary value problems (6) 70 and (7), see [11]. Since the proof of the generalized Grarding inequality requires a 71 comprehensive study of the trace spaces  $\mathbf{H}_{\perp}^{-1/2}(\text{curl}_{\Gamma},\Gamma)$  and  $\mathbf{H}_{\parallel}^{-1/2}(\text{div}_{\Gamma},\Gamma)$ , and 72 of the corresponding Hodge–type splittings, we refer to [2, 11] for a detailed presentation.

By summing up all local boundary integral equation systems with respect to the 75 transmission conditions (5) we finally obtain the following variational formulation 76 to find  $\mathbf{u} \in \mathbf{H}_{\perp}^{-1/2}(\operatorname{curl}_{\Gamma}, \Gamma_{S})$  and  $\mathbf{t}_{i} \in \mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma_{i})$  satisfying 77

$$\sum_{i=1}^{p} \left[ \langle \mathsf{N}_{i} \mathbf{u}_{|\Gamma_{i}}, \mathbf{v}_{|\Gamma_{i}} \rangle_{\Gamma_{i}} + \langle (\frac{1}{2}I + \mathsf{B}_{i}) \mathbf{t}_{i}, \mathbf{v}_{|\Gamma_{i}} \rangle_{\Gamma_{i}} + i \eta_{i} \langle \mathsf{R}_{i} \mathbf{u}_{|\Gamma_{i}}, \mathbf{v}_{|\Gamma_{i}} \rangle_{\Gamma_{i}} \right] = \langle \mathbf{f}, \mathbf{v} \rangle_{\Gamma}$$
(12)

for all  $\mathbf{v} \in \mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{arGamma}\,,arGamma_{\!S})$  and

$$\langle \mathsf{S}_{i} \mathbf{t}_{i}, \mu_{i} \rangle_{\Gamma_{i}} + \langle (-\frac{1}{2}I + \mathsf{C}_{i}) \mathbf{u}_{|\Gamma_{i}}, \mu_{i} \rangle_{\Gamma_{i}} = 0 \tag{13}$$

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for all  $\mu_i \in \mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma_i)$ ,  $i = 1, \dots, p$ . The variational formulation (12), (13) admits a unique solution iff the original problems (1) and (2) has a unique solution, see 80 [11].

A boundary element discretization of the Sobolev spaces  $\mathbf{H}_{\perp}^{-1/2}(\operatorname{curl}_{\Gamma}, \Gamma_{S})$  and 82  $\mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma_{i})$  by using Raviart–Thomas elements [8, 11], i.e. 83

$$\mathscr{E}_h := \mathscr{E}_h(\Gamma_S) = \operatorname{span}\{\phi_k\}_{k=1}^{M_S} \subset \mathbf{H}_{\perp}^{-1/2}(\operatorname{curl}_{\Gamma}, \Gamma_S)$$

and 84

$$\mathscr{F}_{i,h} = \operatorname{span}\{\psi_k^i\}_{k=1}^{N_i} \subset \mathbf{H}_{\parallel}^{-1/2}(\operatorname{div}_{\Gamma}, \Gamma_i),$$

then results in a linear system of algebraic equations,

$$\begin{pmatrix}
S_{1,h} & \widetilde{C}_{1,h}A_{i} \\
\dots & \vdots \\
S_{p,h} & \widetilde{C}_{p,h}A_{p} \\
A_{1}^{\top}\widetilde{B}_{1,h} \dots A_{p}^{\top}\widetilde{B}_{p,h} \sum_{i=1}^{p} A_{i}^{\top}[N_{i,h} + i\eta_{i}R_{i,h}]A_{i}
\end{pmatrix}
\begin{pmatrix}
\underline{t}_{1} \\
\vdots \\
\underline{t}_{p} \\
\underline{u}
\end{pmatrix} = \begin{pmatrix}
\underline{0} \\
\vdots \\
\underline{0} \\
\sum_{i=1}^{p} A_{i}^{\top}\underline{f}_{i}
\end{pmatrix}, (14)$$

where the block matrices are given by

$$\begin{split} \mathsf{S}_{i,h}[\ell,k] &= \langle \mathsf{S}_i \psi_k^i, \psi_\ell^i \rangle_{\Gamma_i}, \\ \widetilde{\mathsf{C}}_{i,h}[\ell,n] &= \langle (-\frac{1}{2}I + \mathsf{C}_i) \phi_n^i, \psi_\ell^i \rangle_{\Gamma_i}, \\ \widetilde{\mathsf{B}}_{i,h}[m,k] &= \langle (\frac{1}{2}I + \mathsf{B}_i) \psi_k^i, \phi_m^i \rangle_{\Gamma_i}, \\ \mathsf{N}_{i,h}[m,n] &= \langle \mathsf{N}_i \phi_n^i, \phi_m^i \rangle_{\Gamma_i}, \\ \mathsf{R}_{i,h}[m,n] &= \langle \mathsf{R}_i \phi_n^i, \phi_m^i \rangle_{\Gamma_i} \end{split}$$

for 
$$k, \ell = 1, ..., N_i, m, n = 1, ..., M_i$$
, and  $i = 1, ..., p$ .

In what follows we will discuss an efficient and parallel solution of the linear 88 system (14). Although the computation of all block matrices can be done in parallel, 89 the construction of an appropriate preconditioner is more challenging. A possible approach is to design preconditioners as in tearing and interconnecting methods which 91 are well established for a wide range of applications. A first step into this direction 92 is the formulation of stable tearing and interconnecting methods. 93

The idea of the tearing and interconnecting approach is to tear the global degrees 94 of freedom, which are given by  $\underline{u}$ , into local degrees of freedom  $\underline{u}_i$ . To ensure global 95 continuity, we need to glue them together by using Langrange multipliers [10, 11], 96 see also Fig. 1. Note, that instead of Neumann transmission condition we use the 97 generalized Robin transmission conditions as given in (5). As in the standard tearing 98 and interconnecting approach this leads to the extended linear system 99

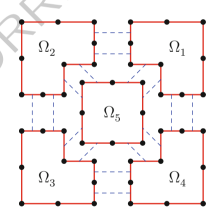


Fig. 1. Tearing and Interconnecting for edge based trial functions

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$$\begin{pmatrix}
\mathsf{N}_{1,h} + i\eta_1 \mathsf{R}_{i,h} \ \widetilde{\mathsf{B}}_{1,h} & -B_1^\top \\
\widetilde{\mathsf{C}}_{1,h} & \mathsf{S}_{1,h} & & \vdots \\
& & \ddots & & \vdots \\
& & \mathsf{N}_{p,h} + i\eta_p \mathsf{R}_{p,h} \ \widetilde{\mathsf{B}}_{p,h} - B_p^\top \\
& & \widetilde{\mathsf{C}}_{p,h} & \mathsf{S}_{p,h}
\end{pmatrix}
\begin{pmatrix}
\underline{u}_1 \\ \underline{t}_1 \\ \vdots \\ \underline{u}_p \\ \underline{t}_p \\ \underline{\lambda}
\end{pmatrix} = \begin{pmatrix}
\underline{f}_1 \\ \underline{0} \\ \vdots \\ \underline{f}_p \\ \underline{0} \\ \underline{0}
\end{pmatrix}$$
(15)

where the sparse and Boolean matrices  $B_i$  ensure the continuity of the global solution. 100 Since the local Robin boundary value problems (6) and (7) are uniquely solvable, 101 the local block matrices are invertible, and we can consider the Schur complement 102 system 103

$$\sum_{i=1}^{p} (0 B_{i}) \begin{pmatrix} \mathsf{N}_{i,h} + i\eta_{i} \mathsf{R}_{i,h} & \widetilde{\mathsf{B}}_{i,h} \\ \widetilde{\mathsf{C}}_{i,h} & \mathsf{S}_{i,h} \end{pmatrix}^{-1} \begin{pmatrix} B_{i}^{\top} \underline{\lambda} \\ \underline{0} \end{pmatrix} \\
= -\sum_{i=1}^{p} (B_{i} 0) \begin{pmatrix} \mathsf{N}_{i,h} + i\eta_{i} \mathsf{R}_{i,h} & \widetilde{\mathsf{B}}_{i,h} \\ \widetilde{\mathsf{C}}_{i,h} & \mathsf{S}_{i,h} \end{pmatrix}^{-1} \begin{pmatrix} \underline{f}_{i} \\ \underline{0} \end{pmatrix}.$$
(16)

Note that (16) corresponds to the adjoint system of standard tearing and interconnecting approaches [4, 5].

### 3 Numerical Results

As a first example we consider the Neumann boundary value problem

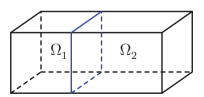
curl curl 
$$\mathbf{U} - k^2 \mathbf{U} = \mathbf{0}$$
 in  $\Omega$ ,  
 $\gamma_N \mathbf{U} = \mathbf{f}$  on  $\Gamma$  (17)

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where the domain  $\Omega$  is given by  $(-1.0, 1.5) \times (0.0, 1.0) \times (0.0, 1.0)$ , and  $\Omega$  is divided 108 into two subdomains  $\Omega_i$  by the yz-plane, see Fig. 2.



**Fig. 2.** Computational domain  $\Omega$  and domain decomposition

As an analytical solution for both examples we use

$$\mathbf{U}(x) = \begin{bmatrix} \frac{1 + ikr - k^2r^2}{r^3} \begin{pmatrix} 1\\0\\0 \end{pmatrix} - \frac{3 + 3ikr - k^2r^2}{r^5} (x_1 - \hat{x}_1) \begin{pmatrix} x_1 - \hat{x}_1\\x_2 - \hat{x}_2\\x_3 - \hat{x}_3 \end{pmatrix} \end{bmatrix} e^{ikr}$$

t1 1 t1 2 t1.3 t1 4 t1.5

t3.1 t3 2 t3.3 t3 4

123

132

135

with  $r = |x - \hat{x}|$  and  $\hat{x} = (-3.0, 2.1, 1.1)^{\top}$ . The boundary element discretization of the coupled variational formulation (12) and (13) is done with respect to a globally 112 uniform boundary element mesh with  $E_i$  edges per subdomain  $\Omega_i$ , and by using first 113 order Raviart-Thomas elements. The number of Lagrange multipliers is denoted by 114  $\Lambda$ . The linear system (16) is solved by a GMRES method with a relative residuum 115 reduction of  $\varepsilon = 10^{-7}$ . For our numerical tests we consider two different wave numbers: The first one is k = 1.0 and the second one is the first Dirichlet and Neumann 117 eigenfrequency of the unit cube  $\Omega_1$ ,  $k = \sqrt{2}\pi \approx 4.44288$ . The results are given in Table 1, where the error is the relative  $L_2(\Gamma_1)$  error of the lowest order Rayiart-Thomas 119 approximation of the local Dirichlet datum  $\mathbf{u}_1$ .

$E_i$	$\Lambda$	iter	error	$E_i$	Λ	iter	error
36	8	5	0.1824189	36	8	5	0.7042192
144	28	17	0.0895037	144	28	19	0.3055468
576	104	49	0.0440296	576	104	47	0.1472184
2304	400	142	0.0234164	2304	400	104	0.0772003

**Table 1.** Iteration numbers and errors for k = 1 (left) and  $k = \sqrt{2}\pi$  (right).

In a second example we consider the Neumann boundary value problem (17) for the 121 unit cube  $\Omega = (0,1)^3$  which is divided into eight subcubes  $\Omega_i$ . The results for two 122 different wave numbers k = 1.0, 8.0 are given in Table 2.

$E_i$	$\Lambda$	iter	error	$E_i$	$\Lambda$	iter	error
36	90	60	0.1133393	36	90	60	0.9432815
144	324	147	0.0550944	144	324	153	0.3776120
576	1224	476	0.0266769	576	1224	397	0.1769975

**Table 2.** Iteration numbers and errors for k = 1 (left) and k = 8 (right).

Both numerical experiments confirm the stability and robustness of the proposed 124 approach, and the theoretical error estimate as given in [11], i.e. we expect a linear 125 order of convergence when using lowest order Raviart-Thomas elements. Note that 126 the linear system (16) is solved by a GMRES method without preconditioner. Hence 127 we observe a rapidly increasing number of required iterations. Therefore, the use 128 of local and global preconditioners is mandatory for the solution of problems of 129 practical interest. Probably, possible preconditioners can be constructed as in the 130 acoustic scattering case see [11]. Another possibility is to consider a dual-primal 131 approach as in [6].

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# Preconditioning High-Order Discontinuous Galerkin **Discretizations of Elliptic Problems**

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1 Introduction 9

In recent years, attention has been devoted to the development of efficient iterative 10 solvers for the solution of the linear system of equations arising from the discontinuous Galerkin (DG) discretization of a range of model problems. In the frame- 12 work of two level preconditioners, scalable non-overlapping Schwarz methods have 13 been proposed and analyzed for the h-version of the DG method in the articles 14 [1, 2, 6, 7, 9]. Recently, in [3] it has been proved that the non-overlapping Schwarz 15 preconditioners can also be successfully employed to reduce the condition number 16 of the stiffness matrices arising from a wide class of high-order DG discretizations 17 of elliptic problems. In this article we aim to validate the theoretical results derived 18 in [3] for the multiplicative Schwarz preconditioner and for its symmetrized variant 19 by testing their numerical performance.

#### 2 Model Problem and DG Discretization

In this section we introduce the model problem under consideration and its DG ap- 22 proximation, working, for the sake of simplicity, with the SIPG formulation proposed 23 in [4].

We consider, for simplicity, the weak formulation of the Poisson problem with 26 homogeneous Dirichlet boundary conditions: find  $\mathcal{U} \in H_0^1(\Omega)$  such that

$$(\nabla \mathcal{U}, \nabla v)_{\Omega} = (f, v)_{\Omega} \quad \forall v \in H_0^1(\Omega), \tag{1}$$

where  $\Omega$  is a bounded polygonal domain in  $\mathbb{R}^d$ ,  $d=2,3,f\in L^2(\Omega)$  is a given source 28 term and  $(\cdot,\cdot)_{\Omega}$  is the standard inner product in  $[L^2(\Omega)]^d$ .

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Let  $\mathscr{T}_h$  be a shape-regular, not necessarily matching partition of  $\Omega$  into disjoint 31 open elements  $\mathscr{K}$  (with diameter  $h_{\mathscr{K}}$ ), where each  $\mathscr{K}$  is the affine image of a fixed 32 master element  $\widehat{\mathscr{K}}$ , i.e.,  $\mathscr{K} = F_{\mathscr{K}}(\widehat{\mathscr{K}})$ , where  $\widehat{\mathscr{K}}$  is either the open unit d-simplex or 33 the d-hypercube in  $\mathbb{R}^d$ , d=2,3. We define the mesh-size h by  $h:=\max_{\mathscr{K}\in\mathscr{T}_h}h_{\mathscr{K}}$ , 34 and assume that  $\mathscr{T}_h$  satisfies a bounded local variation property: for any pair of 35 neighboring elements  $\mathscr{K}_1, \mathscr{K}_2 \in \mathscr{T}_h, h_{\mathscr{K}_1} \approx h_{\mathscr{K}_2}$ .

For a given approximation order  $p \ge 1$ , we define the DG space

$$V_{h,p} := \{ v \in L^2(\Omega) : v |_{\mathscr{K}} \circ F_{\mathscr{K}} \in \mathscr{M}^p(\widehat{\mathscr{K}}) \ \forall \ \mathscr{K} \in \mathscr{T}_h \},$$
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where  $\mathscr{M}^p(\widehat{\mathscr{K}})$  is either the space of polynomials of degree at most p on  $\widehat{\mathscr{K}}$ , if  $\widehat{\mathscr{K}}$  39 is the reference d-simplex, or the space of polynomials of degree at most p in each 40 variable on  $\widehat{\mathscr{K}}$ , if  $\widehat{\mathscr{K}}$  is the reference d-hypercube.

Next, for any internal face  $\overline{F} = \overline{\partial \mathcal{K}^{+}} \cap \overline{\partial \mathcal{K}^{-}}$  shared by two adjacent elements 42  $\mathcal{K}^{\pm}$ , with outward unit normal vectors  $\mathbf{n}^{\pm}$ , respectively, we define 43

$$[\![\tau]\!] := \tau^+ \cdot \mathbf{n}^+ + \tau^- \cdot \mathbf{n}^-, \qquad [\![v]\!] := v^+ \mathbf{n}^+ + v^- \mathbf{n}^-, \{\![\tau]\!] := (\tau^+ + \tau^-)/2, \qquad \{\![v]\!] := (v^+ + v^-)/2,$$

where  $\tau^{\pm}$  and  $v^{\pm}$  denote the traces on  $\partial \mathscr{K}^{\pm}$  taken from the interior of  $\mathscr{K}^{\pm}$  of the 44 (sufficiently regular) functions  $\tau$  and v, respectively (cf. [5]). On a boundary face 45  $\overline{F} = \overline{\partial \mathscr{K}} \cap \overline{\partial \Omega}$ , we set  $[\![\tau]\!] := \tau \cdot \mathbf{n}$ ,  $[\![v]\!] := v \cdot \mathbf{n}$ ,  $[\![\tau]\!] := \tau$ , and  $\{\![v]\!] := v$ .

We collect all interior (respectively, boundary) faces in the set  $\mathscr{F}_h^I$  (respectively, 48  $\mathscr{F}_h^B$ ), define  $\mathscr{F}_h:=\mathscr{F}_h^I\cup\mathscr{F}_h^B$ , and introduce on  $V_{h,p}\times V_{h,p}$  the following the bilinear 49 form

$$\begin{aligned} \mathscr{A}(u,v) &:= \sum_{\mathscr{K} \in \mathscr{T}_h} \int_{\mathscr{K}} \nabla u \cdot \nabla v \, d\mathbf{x} + \sum_{\mathscr{K} \in \mathscr{T}_h} \int_{\mathscr{K}} \nabla u \cdot \mathscr{R}(\llbracket v \rrbracket) \, d\mathbf{x} \\ &+ \sum_{\mathscr{K} \in \mathscr{T}_h} \int_{\mathscr{K}} \mathscr{R}(\llbracket u \rrbracket) \cdot \nabla v \, d\mathbf{x} + \sum_{F \in \mathscr{F}_h} \int_{F} \alpha \frac{p^2}{|F|} \llbracket u \rrbracket \cdot \llbracket v \rrbracket \, d\mathbf{s}, \end{aligned}$$

where  $\alpha > 0$  is a parameter at our disposal. The lifting operator  $\mathscr{R}(\cdot)$  is defined as: 51  $\mathscr{R}(\tau) := \sum_{F \in \mathscr{F}_h} r_F(\tau)$ , where  $r_F : [L^2(F)]^d \to [V_{h,p}]^d$  is given by

$$\int_{\Omega} r_F(\tau) \cdot \eta \, d\mathbf{x} := -\int_F \tau \cdot \{\!\!\{\eta\}\!\!\} \, d\mathbf{s} \quad \forall \eta \in [V_{h,p}]^d \quad \forall F \in \mathscr{F}_h.$$

The DG discretization of problem (1) reads:

Find 
$$u \in V_{h,p}$$
 such that  $\mathscr{A}(u,v) = \int_{\Omega} fv \, dx \quad \forall v \in V_{h,p}.$  (2)

Let  $\varphi_j$ ,  $j = 1, ..., N_h^p := \dim(V_{h,p})$ , be a set of basis functions that span  $V_{h,p}$ , then 54 (2) can be written in the following equivalent form: Find  $\mathbf{u} \in \mathbb{R}^{N_h^p}$  such that  $\mathbf{A}\mathbf{u} = \mathbf{f}$ , 55 where here (and in the following) we use the bold notation to denote the spaces of 56

degrees of freedom (vectors) and discrete linear operators (matrices). The following 57 result provides an estimate for the spectral condition number of **A**; we refer to [3] for 58 the proof. 59

**Proposition 1** ([3]). For a set of basis functions which are orthonormal on the reference element  $\widehat{\mathcal{K}} \subset \mathbb{R}^d$ , d=2,3, the condition number  $\kappa(A)$  of the stiffness matrix A 61 can be bounded by

$$\kappa(A) \lesssim \alpha \frac{p^4}{h^2}.$$

Remark 1. We are working, for the sake of simplicity, with the SIPG formulation 63 proposed in [4], but the results shown in Proposition 1 and in Theorem 1 below also 64 hold for a wide class of DG methods; we refer to [3] for details.

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## 3 Two Level Non-overlapping Schwarz Preconditioners

In this section we introduce the non-overlapping Schwarz preconditioners.

**Subdomain partition.** We decompose the domain  $\Omega$  into N non-overlapping subdomains  $\Omega_i$ , i.e.,  $\overline{\Omega} = \bigcup_{i=1}^N \overline{\Omega}_i$ . Next, we consider two levels of *nested* partitions of 70 the domain  $\Omega$ : (i) a coarse partition  $\mathcal{T}_H$  (with mesh-size H); (ii) a fine partition  $\mathcal{T}_h$  71 (with mesh-size h). We will suppose that the subdomain partition does not cut any 72 element of  $\mathcal{T}_H$  (and therefore of  $\mathcal{T}_h$ ).

**Local solvers.** For i = 1, ..., N, we define the local DG spaces as

$$V_{h,p}^i := \{ v \in L^2(\Omega_i) \ : \ v |_{\mathscr{K}} \circ F_{\mathscr{K}} \in \mathscr{M}^p(\widehat{\mathscr{K}}) \quad \forall \ \mathscr{K} \in \mathscr{T}_h, \mathscr{K} \subset \Omega_i \}.$$

Denoting by  $R_i^T: V_{h,p}^i \longrightarrow V_{h,p}$  the classical injection operator from  $V_{h,p}^i$  to  $V_{h,p}$ , the 76 local solvers  $\mathscr{A}_i: V_{h,p}^i \times V_{h,p}^i \longrightarrow \mathbb{R}$  are defined as

$$\mathscr{A}_i(u_i, v_i) := \mathscr{A}(R_i^{\mathsf{T}} u_i, R_i^{\mathsf{T}} v_i) \quad \forall u_i, v_i \in V_{h,p}^i, \quad i = 1, \dots, N.$$
(3)

**Coarse solver.** For an integer  $0 \le q \le p$ , we define the coarse space  $V_{H,q}^0$  as

$$V_{H,q}^0 := \{ v \in L^2(\Omega) : v|_{\mathscr{D}} \circ F_{\mathscr{D}} \in \mathbb{M}^{q_{\mathscr{D}}}(\widehat{\mathscr{K}}) \quad \forall \ \mathscr{D} \in \mathscr{T}_H \},$$

and the *coarse solver*  $\mathcal{A}_0: V^0_{H,q} \times V^0_{H,q} \longrightarrow \mathbb{R}$  as

$$\mathscr{A}_0(u_0, v_0) := \mathscr{A}(R_0^{\mathsf{T}} u_0, R_0^{\mathsf{T}} v_0) \quad \forall u_0, v_0 \in V_{H,a}^0, \tag{4}$$

where  $R_0^{\mathrm{T}}: V_{H,q}^0 \longrightarrow V_{h,p}$  is the classical injection operator from  $V_{H,q}^0$  to  $V_{h,p}$ .

Let the *local* projection operators be defined as

$$\widetilde{P}_{i}: V_{h,p} \to V_{h,p}^{i}: \qquad \mathscr{A}_{i}(\widetilde{P}_{i}u, R_{i}^{T}v_{i}) := \mathscr{A}(u, R_{i}^{T}v_{i}) \qquad \forall v_{i} \in V_{h,p}^{i}, \quad i = 1, \dots, N, 
\widetilde{P}_{0}: V_{h,p} \to V_{H,q}^{0}: \qquad \mathscr{A}_{0}(\widetilde{P}_{0}u, R_{0}^{T}v_{0}) := \mathscr{A}(u, R_{0}^{T}v_{0}) \qquad \forall v_{0} \in V_{H,q}^{0},$$
(5)

and define the projection operators as  $P_i := R_i^T \widetilde{P_i} : V_{h,p} \longrightarrow V_{h,p}, i = 0, 1, ..., N$ . The 83 multiplicative Schwarz operator and its symmetrized variant are then defined as

$$P_{\text{mu}} := I - (I - P_N)(I - P_{N-1}) \cdots (I - P_0),$$

$$P_{\text{mu}}^{S} := I - (I - P_0)^T \cdots (I - P_N)^T (I - P_N) \cdots (I - P_0),$$
(6)

respectively (cf. [10]). The Schwarz method consists in solving either  $P_{\text{mu}}u = g_{\text{mu}}$  85 or  $P_{\text{mu}}^{\text{S}}u = g_{\text{mu}}^{\text{S}}$ , for suitable right hand sides  $g_{\text{mu}}$  and  $g_{\text{mu}}^{\text{S}}$ , respectively. It can be 86 shown that the operator defined in (7) is symmetric and positive definite; we therefore 87 consider the conjugate gradient (CG) algorithm for the solution of  $P_{\text{mu}}^{\text{S}}u = g_{\text{mu}}^{\text{S}}$ . An 88 estimate of the condition number of  $P_{\text{mu}}^{\text{S}}$  is

$$\kappa(P_{\mathrm{mu}}^{\mathrm{S}}) := \frac{\lambda_{\mathrm{max}}(P_{\mathrm{mu}}^{\mathrm{S}})}{\lambda_{\mathrm{min}}(P_{\mathrm{mu}}^{\mathrm{S}})},$$

where  $\lambda_{\max}(P_{\min}^S)$  and  $\lambda_{\min}(P_{\min}^S)$  are the extremal eigenvalues of the operator  $P_{\min}^S$ . 90 On the other hand, the multiplicative operator  $P_{\min}$  is non-symmetric; we therefore 91 consider a Richardson iteration applied to  $P_{\min}u = g_{\min}$ , and show that the norm of 92 the error propagation operator  $E_{\min} := (I - P_N)(I - P_{N-1}) \cdots (I - P_0)$  is strictly less 93 than one, i.e., 94

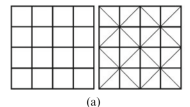
$$||E_{\mathrm{mu}}||_{\mathscr{A}}^{2} := \sup_{\substack{v \in V_{h,p} \\ v \neq 0}} \frac{\mathscr{A}(E_{\mathrm{mu}}v, E_{\mathrm{mu}}v)}{\mathscr{A}(v,v)} < 1,$$

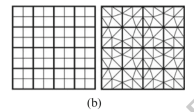
and therefore a Richardson iteration applied to the preconditioned system converges. 95 The following result provides a bound for the norm of the error propagation operator of the multiplicative Schwarz operator, and for the condition number of the symmetrized Schwarz operator (we refer to [3] for the proof). 98

**Theorem 1** ([3]). There exists constants  $C_1, C_2 \ge 1$ , independent of the mesh-size 99 and the polynomial degree, such that

$$||E_{mu}||_{\mathscr{A}}^2 \leq 1 - \frac{h}{C_1 \alpha p^2 H}, \quad \kappa(P_{mu}^S) \leq C_2 \alpha p^2 \frac{H}{h}.$$

Theorem 1 also guarantees that the multiplicative Schwarz method can be accelerated with the GMRES iterative solver. Indeed, according to [8], the GMRES method applied to the preconditioned system  $P_{\text{mu}}u=g_{\text{mu}}$  does not stagnate (i.e., 103 the iterative method makes some progress in reducing the residual at each iteration step) provided that: (i)  $\|P_{mu}\|_{\mathscr{A}}$  is bounded; (ii) the symmetric part of  $P_{mu}$  is 105 positive definite, i.e., there exists  $c_p>0$  such that  $\mathscr{A}(v,P_{\text{mu}}v)>c_p\mathscr{A}(v,v)$  for all 106  $v\in V_{h,p}$ . Condition (i) follows directly from the definition of  $P_{\text{mu}}$  and Theorem 1: 107  $\|P_{mu}\|_{\mathscr{A}}=\|I-E_{\text{mu}}\|_{\mathscr{A}}\leq 1+\|E_{\text{mu}}\|_{\mathscr{A}}<2$ . To prove condition (ii), it can be shown 108 that





**Fig. 1.** Initial Cartesian and triangular coarse and fine grids on a 16 subdomain partition. (a) Initial coarse grids (mesh-size  $H_0$ ) and (b) initial fine grids (mesh-size  $h_0$ )

$$\mathscr{A}(P_{\mathrm{mu}}v,v) = \mathscr{A}(v,v) - \mathscr{A}(E_{\mathrm{mu}}v,v) \ge (1 - ||E_{\mathrm{mu}}||_{\mathscr{A}}) \mathscr{A}(v,v).$$

Therefore, condition (ii) holds true with  $c_p = 1 - ||E_{\text{mu}}||_{\mathscr{A}}$  which is positive due to 110 Theorem 1.

#### 4 Numerical Results

In this section we present some numerical experiments to highlight the practical performance of the multiplicative and symmetrized non-overlapping Schwarz preconditioners. From the algebraic point of view, the Schwarz operators (6) and (7) canbe written as the product of a suitable preconditioner, namely  $\mathbf{B}_{\text{mu}}$ ,  $\mathbf{B}_{\text{mu}}^{\text{S}}$ , respectively, and  $\mathbf{A}$ . Indeed, the local components can be constructed as  $\mathbf{A}_i = \mathbf{R}_i \mathbf{A} \mathbf{R}_i^T$ , see(3) for i = 1..., N, and (4) for i = 0. From the definition (5) of the local projection- $\widetilde{\mathbf{P}}_i = \mathbf{A}_i^{-1} \mathbf{R}_i \mathbf{A}$ , and therefore  $\mathbf{P}_i = \mathbf{R}_i^T \widetilde{\mathbf{P}}_i = \mathbf{R}_i^T \mathbf{A}_i^{-1} \mathbf{R}_i \mathbf{A}$ . In practice, only the actionof the preconditioner on a vector is needed. Algorithm 2 shows how to computethe action of  $\mathbf{B}_{\text{mu}}$  on a vector  $\mathbf{x} \in \mathbb{R}^{N_h^p}$ . Throughout this section we have set the

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\begin{aligned} & \overline{\mathbf{Algorithm}} \ \mathbf{2} \ \mathbf{z} = \mathbf{B}_{\text{mu}} \mathbf{x} \\ & \mathbf{z} = \mathbf{R}_0^T \mathbf{A}_0^{-1} \mathbf{R}_0 \mathbf{x} \\ & \mathbf{for} \ i = 1 \rightarrow N \ \mathbf{do} \\ & \mathbf{z} = \mathbf{z} + \mathbf{R}_i^T \mathbf{A}_i^{-1} \mathbf{R}_i (\mathbf{x} - \mathbf{Az}) \\ & \mathbf{end} \ \mathbf{for} \end{aligned}
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penalty parameter  $\alpha := 10$  (see (2)). We consider a subdomain partition consisting of N = 16 squares, and consider the initial Cartesian and unstructured triangular partitions shown in Fig. 1, and denote by  $H_0$  and  $h_0$  the corresponding initial coarse and fine mesh-sizes, respectively. We consider n successive global uniform refinements of these initial grids so that the resulting mesh-sizes are  $H_n = H_0/2^n$  and  $h_n = h_0/2^n$ , with n = 0, 1, 2, 3, respectively. The (relative) tolerance is set equal to  $10^{-9}$  (respectively,  $10^{-6}$ ) for the CG (respectively, GMRES) iterative solver. We first address the performance of the multiplicative Schwarz preconditioner by keeping the mesh fixed, and varying the polynomial approximation degree p. In Table 1 we compare the GMRES iteration counts for both the preconditioned and non-preconditioned (in 131)

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**Table 1.** GMRES iteration counts. Multiplicative Schwarz preconditioner with a piecewise constant coarse solver (q = 0). Unstructured triangular grids.

	$h = h_0/2$	$h = h_0/4$	$h = h_0/4$
	$H = H_0$	$H = H_0$	$H = H_0/2$
p=1	23 (94)	33 (199)	25 (199)
p = 2	45 (259)	64 (540)	49 (540)
p = 3	66 (470)	93 (996)	74 (996)
p = 4	85 (713)	124 (1546)	97 (1546)
p = 5	105 (1004)	153 (2187)	123 (2187)
p = 6	124 (1342)	183 (2924)	144 (2924)
p = 7	143 (1727)	209 (3742)	167 (3742)
p = 8	162 (2148)	235 (4673)	189 (4673)
p-rate	0.93 (1.63)	0.88 (1.66)	0.93 (1.66)

parenthesis) systems, for different polynomial approximation degrees and different 132 mesh configurations. These results have been obtained on unstructured triangular 133 grids (cf. Fig. 1). Comparing the iteration counts of the preconditioned systems with 134 the unpreconditioned ones for a fixed p, it is clear that the proposed preconditioner is 135 very efficient. Indeed, we observe a reduction in the number of iterations needed to 136 achieve convergence of around one order of magnitude when the proposed preconditioner is employed. The last row of Table 1 shows the computed growth rate in the 138 number of iterations: we observe that the number of iterations needed to obtain convergence increases linearly as a function of p for the preconditioned system of equations, whereas this quantity grows almost quadratically for the non-preconditioned 141 problem. In Fig. 2 we report the condition number estimates of the symmetrized 142 Schwarz operator and the corresponding iteration counts versus the polynomial de- 143 gree p. The solid lines refer to the mesh configuration  $h = h_0/2$ ,  $H = H_0$ , whereas 144 the dashed lines refer to the mesh configuration  $h = h_0/4$ ,  $H = H_0/2$ . This set of numerical experiments has been obtained on Cartesian meshes, employing a piecewise 146 linear coarse solver. As predicted by the theoretical estimates, the condition number of the preconditioned system grows quadratically as a function of p. Moreover, 148 we clearly observe that, for fixed p, by refining both the fine and the coarse grid, 149 but keeping the ratio of the fine and coarse mesh-sizes constant, the condition number (and therefore the number of iterations needed to obtain convergence) remains 151 constant.

Next, we consider the performance of the symmetrized Schwarz preconditioner 153 when varying the coarse and fine mesh-size, and keeping the polynomial approxima- 154 tion degree p fixed. In Table 2 (left) we report the condition number estimates for the symmetrized Schwarz operator employing piecewise biquadratic elements (p = 2) 156 and a piecewise constant coarse solver (q = 0); whereas, in Table 2 (right) the analogous results obtained with piecewise bicubic elements (p = 3) and a piecewise linear 158 coarse solver (q = 1) are shown. We clearly observe that the condition number grows 159

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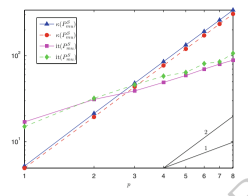


Fig. 2. Condition number estimates of the symmetrized Schwarz operator and corresponding iteration counts versus the polynomial degree p on Cartesian grids for different discretization steps (solid line:  $h = h_0/2$ ,  $H = H_0$ ; dashed line  $h = h_0/4$ ,  $H = H_0/2$ ). Piecewise linear coarse solver

**Table 2.** Condition number estimates for the symmetrized Schwarz operator with p = 2, q = 0(left) and p = 3, q = 1 (right). Cartesian grids.

$h \downarrow H \rightarrow$	$H_0$	$H_0/2$	$H_0/4$	$H_0/8$	$H_0$	$H_0/2$	$H_0/4$	$H_0/8$
$h_0$	5.32e2	1.12e3	4.01e3	7.08e3	4.81e1	9.5925e1	1.92e2	3.91e2
$h_0/2$	2.74e2	4.71e2	2.80e3	5.59e3	2.14e1	4.35e1	8.70e1	1.75e2
$h_0/4$	_	2.60e2	1.18e3	3.42e3	_	2.09e1	4.24e1	8.44e1
$h_0/8$	-	-	3.45e2	1.75e3	_	_	2.05e1	4.26e1
$\kappa(\mathbf{A})$	2.88e5	1.18e6	4.89e6	1.99e7	7.44e5	2.81e6	1.11e7	4.55e7

as  $O(Hh^{-1})$ , as predicted by Theorem 1. Moreover, we clearly observe that employing a piecewise linear coarse solver (q = 1) rather than a piecewise constant coarse solver (q = 0) significantly improves the performance of the preconditioner. Indeed, 162 comparing the condition number estimates of the preconditioned system with the 163 analogous ones obtained for the non-preconditioned problem (last row of Table 2) 164 we clearly observe that the condition number of the non-preconditioned system is 165 reduced with respect to the condition number of the preconditioned system by approximately 5 orders of magnitude for q = 1 and 4 orders of magnitude for q = 0.

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# A Block Solver for the Exponentially Fitted IIPG-0 Method

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Summary. We consider an exponentially fitted discontinuous Galerkin method for advection 10 dominated problems and propose a block solver for the resulting linear systems. In the case of 11 strong advection the solver is robust with respect to the advection direction and the number of 12 unknowns. 13

1 Introduction 14

Let  $\Omega \subset \mathbb{R}^2$  be a polygon,  $f \in L^2(\Omega)$ ,  $g \in H^{1/2}(\partial \Omega)$  and let  $\varepsilon > 0$  be constant. We 15 consider the advection-diffusion problem 16

$$-\operatorname{div}(\varepsilon \nabla u - \beta u) = f \quad \text{in } \Omega, \qquad u = g \quad \text{on } \partial \Omega, \tag{1}$$

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 $-\operatorname{div}(\varepsilon\nabla u-\beta u)=f\quad\text{in }\Omega,\qquad u=g\quad\text{on }\partial\Omega,\tag{1}$  where  $\beta\in[W^{1,\infty}(\Omega)]^2$  derives from a potential  $\beta=\nabla\psi$ . In applications to semiconductor devices, u is the electron density,  $\psi$  the electrostatic potential and the electric 18 field  $\nabla \psi$  might be fairly large in some parts of  $\Omega$ , so that (1) becomes advection 19 dominated. Its robust numerical approximation and the design of efficient solvers, 20 are still a challenge. Exponential fitting [2] and discontinuous Galerkin (DG) are two 21 approaches that have been combined in [3] to develop exponentially fitted DG meth- 22 ods (in primal and mixed formulation). In this note, we consider a variant of these 23 schemes, based on the use of the Incomplete Interior Penalty IIPG-0 method and 24 propose an efficient solver for the resulting linear systems.

The change of variable  $\rho := e^{-\frac{\psi}{\varepsilon}} u$  in the problem (1) leads to

$$-\nabla \cdot (\kappa \nabla \rho) = f \text{ in } \Omega, \quad \rho = \chi \text{ on } \partial \Omega,$$
 (2)

where  $\kappa := \varepsilon e^{\frac{\psi}{\varepsilon}}$  and  $\chi := e^{-\frac{\psi}{\varepsilon}} g$ . An IIPG-0 approximation to (2) gives rise to the EF- 27 IIPG-0 scheme for (1). We propose a block solver that uses ideas from [1] and reduce 28 the solution to that of an exponentially fitted Crouziex-Raviart (CR) discretization, 29

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which has much less degrees of freedom. The associated (CR) matrix is further reduced to an approximate block lower triangular form, which is efficiently solved by a block Gauss-Siedel algorithm.

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In our description we focus on the case  $\beta = \nabla \psi$  piecewise constant; although 34 we include some numerical results for a more general case (cf. Test 2). Extensions 35 of the method (allowing  $\psi$  to be discontinuous) and further analysis of the proposed 36 solvers are topics of current research.

### 2 The Exponentially Fitted IIPG-0 Method

Let  $\mathscr{T}_h$  be a shape-regular family of partitions of  $\Omega$  into triangles T and let h=39  $\max_{T\in\mathscr{T}_h}h_T$  with  $h_T$  denoting the diameter of T for each  $T\in\mathscr{T}_h$ . We assume  $\mathscr{T}_h$  40 does not contain hanging nodes. We denote by  $\mathscr{E}_h^o$  and  $\mathscr{E}_h^\partial$  the sets of all interior and 41 boundary edges, respectively, and we set  $\mathscr{E}_h=\mathscr{E}_h^o\cup\mathscr{E}_h^\partial$ . 42 Let  $T^+$  and  $T^-$  be two neighboring elements, and  $\mathbf{n}^+$ ,  $\mathbf{n}^-$  be their outward normal 43

Let  $T^+$  and  $T^-$  be two neighboring elements, and  $\mathbf{n}^+$ ,  $\mathbf{n}^-$  be their outward normal 43 unit vectors, respectively ( $\mathbf{n}^{\pm} = \mathbf{n}_{T^{\pm}}$ ). Let  $\zeta^{\pm}$  and  $\boldsymbol{\tau}^{\pm}$  be the restriction of  $\zeta$  and  $\boldsymbol{\tau}$  to 44  $T^{\pm}$ . We define the average and jump trace operators:

$$\begin{split} &2\{\zeta\} = (\zeta^+ + \zeta^-), \quad [\![\zeta]\!] = \zeta^+ \mathbf{n}^+ + \zeta^- \mathbf{n}^- & \text{on } E \in \mathcal{E}_h^o, \\ &2\{\boldsymbol{\tau}\} = (\boldsymbol{\tau}^+ + \boldsymbol{\tau}^-), \quad [\![\boldsymbol{\tau}]\!] = \boldsymbol{\tau}^+ \cdot \mathbf{n}^+ + \boldsymbol{\tau}^- \cdot \mathbf{n}^- & \text{on } E \in \mathcal{E}_h^o, \end{split}$$

and on  $e \in \mathscr{E}_h^{\partial}$  we set  $[\![\zeta]\!] = \zeta \mathbf{n}$  and  $\{\boldsymbol{\tau}\} = \boldsymbol{\tau}$ . We will also use the notation

$$(u,w)_{\mathcal{I}_h} = \sum_{T \in \mathcal{T}_h} \int_T uw dx \qquad \langle u,w \rangle_{\mathcal{E}_h} = \sum_{e \in \mathcal{E}_h} \int_e uw ds \quad \forall u,w, \in V^{DG} \;,$$

where  $V^{DG}$  is the discontinuous linear finite element space defined by:

$$V^{DG} = \left\{ u \in L^2(\Omega) : u_{|_T} \in \mathbb{P}^1(T) \, \forall T \in \mathscr{T}_h \right\},\,$$

Here,  $\mathbb{P}^1(T)$  is the space of linear polynomials on T. Similarly,  $\mathbb{P}^0(T)$  and  $\mathbb{P}^0(e)$  are 48 the spaces of constant polynomials on T and e, respectively. For each  $e \in \mathscr{E}_h$ , let  $\mathscr{P}^0_e$ : 49  $L^2(e) \mapsto \mathbb{P}^0(e)$  (resp.  $\mathscr{P}^0_T: L^2(T) \mapsto \mathbb{P}^0(T)$ , for each  $T \in \mathscr{T}_h$ ) be the  $L^2$ -orthogonal 50 projections defined by

$$\mathscr{P}_e^0(u) := \frac{1}{|e|} \int_e u, \quad \forall u \in L^2(e) \;, \quad \mathscr{P}_T^0(v) := \frac{1}{|T|} \int_T v, \quad \forall v \in L^2(T) \;.$$

We denote by  $V^{CR}$  the classical Crouziex-Raviart (CR) space:

$$V^{CR} = \left\{ v \in L^2(\Omega) : v_{\mid_T} \in \mathbb{P}^1(T) \, \forall T \in \mathcal{T}_h \text{ and } \mathscr{P}_e^0[\llbracket v \rrbracket] = 0 \, \forall e \in \mathscr{E}_h \right\}.$$

Note that v=0 at the midpoint  $m_e$  of each  $e\in\mathcal{E}_h^\partial$ . To represent the functions in  $V^{DG}$  53 we use the basis  $\{\varphi_{e,T}\}_{T\in\mathscr{T}_h,e\in\mathcal{E}_h}$ , defined by

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$$\forall T \in \mathcal{T}_h \quad \varphi_{e,T}(x) \in \mathbb{P}^1(T) \quad e \subset \partial T \quad \varphi_{e,T}(m_{e'}) = \delta_{e,e'} \quad \forall e' \in \mathcal{E}_h . \tag{3}$$

In particular, any  $w \in \mathbb{P}^1(T)$  can be written as  $w = \sum_{e \subset \partial T} w(m_e) \varphi_{e,T}$ .

We first consider the IIPG-0 approximation to the solution of (2): Find  $\rho \in V^{DG}$  57 such that  $\mathscr{A}(\rho,w)=(f,w)_{\mathscr{T}_h}$  for all  $w\in V^{DG}$  with

$$\mathscr{A}(\rho, w) = (\kappa_T^* \nabla \rho, \nabla w)_{\mathscr{T}_h} - \langle \{\kappa_T^* \nabla \rho\}, [[w]] \rangle_{\mathscr{E}_h} + \langle S_e \{[[\rho]]\}, \mathscr{P}^0([[w]]) \rangle_{\mathscr{E}_h}.$$
(4)

Here,  $S_e$  is the penalty parameter and  $\kappa_T^* \in \mathbb{P}^0(T)$  the harmonic average approximation to  $\kappa = \varepsilon e^{\psi/\varepsilon}$  both defined in [3] by:

$$\kappa_T^* := \frac{1}{\mathscr{P}_T^0(\kappa^{-1})} = \frac{\varepsilon}{\mathscr{P}_T^0(e^{-\frac{\psi}{\varepsilon}})}, \qquad S_e := \alpha_e h_e^{-1} \{\kappa_T^*\}_e, \tag{5}$$

Next, following [3] we introduce the local operator  $\mathfrak{T}: V^{DG} \mapsto V^{DG}$  that approximates the change of variable introduced before (2):

$$\mathfrak{T}w := \sum_{T \in \mathscr{T}_h} (\mathfrak{T}w)|_T = \sum_{T \in \mathscr{T}_h} \sum_{e \subset \partial T} \mathscr{P}_e^0(e^{-\frac{\Psi}{\varepsilon}}) w(m_e) \varphi_{e,T} \quad \forall w \in V^{DG} .$$
 (6)

By setting  $\rho := \mathfrak{T}u$  in (4), we finally get the EF-IIPG-0 approximation to (1):

Find 
$$u_h \in V^{DG}$$
 s.t.  $\mathscr{B}(u_h, w) := \mathscr{A}(\mathfrak{T}u_h, w) = (f, w)_{\mathscr{T}_h} \ \forall w \in V^{DG}$  with

$$\mathscr{B}(u,w) = (\kappa_T^* \nabla \mathfrak{T} u, \nabla w)_{\mathcal{T}_h} - \langle \{\kappa_T^* \nabla \mathfrak{T} u\}, [\![w]\!] \rangle_{\mathcal{E}_h} + \langle S_e \{ [\![\mathfrak{T} u]\!] \}, \mathscr{P}^0 [\![w]\!] \rangle_{\mathcal{E}_h} . \tag{7}$$

It is important to emphasize that the use of harmonic average to approximate  $\kappa = 66$   $\varepsilon e^{\psi/\varepsilon}$  as defined in (5) together with the definition of the local approximation of the 67 change of variables prevents possible overflows in the computations when  $|\nabla \psi|$  is 68 large and  $\varepsilon$  is small. (See [3] for further discussion).

Also, these two ingredients are essential to ensure that the resulting method has 70 an automatic upwind mechanism built-in that allows for an accurate approximation 71 of the solution of (1) in the advection dominated regime. We will discuss this in more 72 detail in Sect. 3.

Prior to close this section, we define for each  $e \in \mathcal{E}_h$  and  $T \in \mathcal{T}_h$ :

$$\psi_{m,e} := \min_{x \in e} \psi(x) \quad \psi_{m,T} := \min_{x \in T} \psi(x); \quad \psi_{m,T} \le \psi_{m,e} \text{ for } e \subset \partial T.$$

In the advection dominated regime  $\varepsilon \ll |\beta|h = |\nabla \psi|h$ 

$$\mathscr{P}^0_T(e^{-(\psi/\varepsilon)}) \simeq \varepsilon^2 e^{-\frac{\psi_{m,T}}{\varepsilon}} \qquad \qquad \mathscr{P}^0_{e_i}(e^{-\psi/\varepsilon}) \simeq \varepsilon e^{-\frac{\psi_{m,e}}{\varepsilon}} \ . \tag{8}$$

The first of the above scalings together with the definitions in (5) implies

$$\kappa_T^* \simeq \frac{1}{\varepsilon} e^{\frac{\psi_{m,T}}{\varepsilon}}, \qquad S_e \simeq \frac{\alpha}{2\varepsilon} |e|^{-1} e^{\frac{(\psi_{m,T_1} + \psi_{m,T_2})}{\varepsilon}} \qquad e = \partial T_1 \cap \partial T_2.$$
(9)

### 3 Algebraic System and Properties

Let A and B be the operators associated to the bilinear forms  $\mathscr{A}(\cdot,\cdot)$  (4) and  $\mathscr{B}(\cdot,\cdot)$  79 (7), respectively. We denote by A and B their matrix representation in the basis 80  $\{\varphi_{e,T}\}_{T\in\mathscr{T}_h,e\in\mathscr{E}_h}$  (3). In this basis, the operator  $\mathfrak{T}$  defined in (6) is represented as a 81 diagonal matrix,  $\mathbb{D}$ , and  $\mathbb{B} = \mathbb{AD}$ . Thus, the approximation to (2) and (1) amounts to 82 solve the linear systems (of dimension  $2n_e - n_b$ ; with  $n_e$  and  $n_b$  being the cardinality 83 of  $\mathcal{E}_h$  and  $\mathcal{E}_h^{\partial}$ , respectively): 84

$$\mathbb{A}\boldsymbol{\rho} = \boldsymbol{F}$$
, and  $\mathbb{D}\boldsymbol{u} = \boldsymbol{\rho}$  or  $\mathbb{B}\boldsymbol{u} = \widetilde{\boldsymbol{F}}$ , (10)

where  $\rho$ , u, F and  $\widetilde{F}$  are the vector representations of  $\rho$ , u and the right hand sides 85 of the approximate problems. From the definition (6) of T it is easy to deduce the 86 scaling of the entries of the diagonal matrix  $\mathbb{D} = (d_{i,i})_{i=1}^{2n_e-n_b}$ 

$$\mathbb{D} = (d_{i,j})_{i,j=1}^{2n_e-n_b} \quad d_{i,i} = \mathscr{P}^0_{e_i}(e^{-\psi/arepsilon}) \simeq arepsilon e^{-rac{\psi m_e}{arepsilon}}, \quad d_{i,j} \equiv 0 \quad i 
eq j \ .$$

We now revise a result from [1]:

**Proposition 1.** Let  $\mathscr{Z} \subset V^{DG}$  be the space defined by

$$\mathscr{Z} = \{ z \in L^2(\Omega) : z_{|_T} \in \mathbb{P}^1(T) \, \forall T \in \mathscr{T}_h \text{ and } \mathscr{P}_e^0 \{ v \} = 0 \, \forall e \in \mathscr{E}_h^o \}.$$

Then, for any  $w \in V^{DG}$  there exists a unique  $w^{cr} \in V^{CR}$  and a unique  $w^z \in \mathcal{Z}$  such 90 that  $w = w^{cr} + w^z$ , that is:  $V^{DG} = V^{CR} \oplus \mathcal{Z}$ . Moreover,  $\mathcal{A}(w^{cr}, w^z) = 0 \ \forall w^{cr} \in V^{CR}$ , 91 and  $\forall w^z \in \mathcal{Z}$ .

Proposition 1 provides a simple *change of basis* from  $\{\varphi_{e,T}\}$  to canonical basis in 93  $V^{CR}$  and  $\mathscr{Z}$  that results in the following algebraic structure for (10):

$$\boldsymbol{\rho} = \begin{bmatrix} \boldsymbol{\rho}^z \\ \boldsymbol{\rho}^{cr} \end{bmatrix}, \qquad \mathbb{A} = \begin{bmatrix} \mathbb{A}^{zz} & \mathbf{0} \\ \mathbb{A}^{vz} & \mathbb{A}^{vv} \end{bmatrix}, \qquad \mathbb{B} = \begin{bmatrix} \mathbb{B}^{zz} & 0 \\ \mathbb{B}^{vz} & \mathbb{B}^{vv} \end{bmatrix}. \tag{11}$$

Due to the assumed continuity of  $\psi$ ,  $\mathbb{D}$  is still diagonal in this basis. The algebraic 95 structure (11) suggests the following exact solver:

The solution  $u = u^z + u^{cr}$  satisfying  $\mathcal{B}(u, w) = (f, w)_{\mathcal{T}_n}$ , for all  $w \in V^{DG}$  is then obtained by

- 1. Solve for  $u^z$ :  $\mathcal{B}(u^z, w^z) = (f, w^z)_{\mathscr{T}_h} \quad \forall w^z \in \mathscr{Z}$ . 2. Solve for  $u^{cr}$ :  $\mathcal{B}(u^{cr}, w^{cr}) = (f, w^{cr})_{\mathscr{T}_h} \mathcal{B}(u^z, w^{cr}) \quad \forall w^{cr} \in V^{CR}$ .

Next, wet discuss how to solve efficiently each of the above steps.

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Step 1: Solution in the  $\mathscr{Z}$ -space. In [1] it was shown that  $A^{zz}$  is a diagonal posi- 99 tive definite matrix. This is also true for  $\mathbb{B}^{zz}$  since it is the product of two diagonal 100 matrices. The continuity of  $\psi$  implies

$$\mathscr{B}(u^z, w^z) = \langle S_e \mathfrak{T}[[u^z]], \mathscr{P}_e^0([[w^z]]) \rangle_{\mathscr{E}_h} \quad \forall u^z, w^z \in \mathscr{Z}.$$
 (12)

Using (8) and (5) we observe that the entries of  $\mathbb{B}^{zz}$  scale as:

$$\mathbb{B}^{zz}=(b_{i,j})_{i=1}^{n_e}\quad b_{i,j}=S_{e_i}|e_i|d_j\boldsymbol{\delta}_{i,j}\simeq\boldsymbol{\delta}_{i,j}\frac{\alpha}{2}\,e^{-(\psi_{m,e}-\psi_{m,T_1}-\psi_{m,T_2})/\varepsilon}$$

which are always positive, so in particular  $\mathbb{B}^{zz}$  it is also an *M*-matrix.

**Step 2: Solution in**  $V^{CR}$ . In [1] it was shown that the block  $\mathbb{A}^{\nu\nu}$  coincides with the stiffness matrix of a CR discretization of (2), and so it is an s.p.d. matrix. However, 106 this is no longer true for  $\mathbb{B}^{\nu\nu}$  which is positive definite but non-symmetric. 107

$$\mathscr{B}(u^{cr}, w^{cr}) = (\kappa_T^* \nabla \mathfrak{T} u^{cr}, \nabla w^{cr})_{\mathscr{T}_h} \quad \forall \ u^{cr}, w^{cr} \in V^{CR} \ .$$

In principle, the sparsity pattern of  $\mathbb{B}^{\nu\nu}$  is that of a symmetric matrix. Using (8) and 108 (5), we find that the entries of the matrix scale as: 109

$$\mathbb{B}^{vv} = \left(b_{i,j}^{cr}\right)_{i,j}^{n_{cr}:=n_e-n_b} \quad b_{i,j}^{cr} := \kappa_T^* \frac{|e_i||e_j|}{|T|} \mathbf{n}_{e_i} \cdot \mathbf{n}_{e_j} d_j \simeq e^{-\frac{(\psi_{m,e}-\psi_{m,T})}{\varepsilon}}$$
(13)

Since  $\psi$  is assumed to be piecewise linear, for each T, it attains its minimum (and 110 also its maximum) at a vertex of T, say  $x_0$  and  $\psi_{m,e}$  is attained at one of the vertex 111 of the edge e, say  $x_e$ . In particular, this implies that

$$\psi_{m,e} - \psi_{m,T} \approx \nabla \psi \cdot (\mathbf{x_e} - \mathbf{x_0}) = \beta \cdot (\mathbf{x_e} - \mathbf{x_0}) = \begin{cases} 0 & \mathbf{x_e} = \mathbf{x_0} \\ |\beta|h & \mathbf{x_e} \neq \mathbf{x_0} \end{cases}$$

Hence, in the advection dominated case  $\varepsilon \ll |\beta|h$  some of the entries in (13) vanish (up to machine precision) for  $\varepsilon$  small; this is the automatic upwind mechanism 114 intrinsic of the method. As a consequence, the sparsity pattern of  $\mathbb{B}^{\nu\nu}$  is no longer 115 symmetric and this can be exploited to re-order the unknowns so that  $\mathbb{B}^{\nu\nu}$  can be 116 reduced to block lower triangular form.

Notice also that for  $\mathcal{T}_h$  acute, the block  $\mathbb{A}^{\nu\nu}$  being the stiffness matrix of the 118 Crouziex-Raviart approximation to (2), is an M-matrix. Hence, since the block  $\mathbb{B}^{\nu\nu}$ is the product of a positive diagonal matrix and  $\mathbb{A}^{\nu\nu}$ , it will also be an M-matrix if the 120 triangulation is acute (see [2]). 121

## 4 Block Gauss-Siedel Solver for V<sup>CR</sup>-Block

We now consider re-orderings of the unknowns (dofs), which reduce  $\mathbb{B}^{\nu\nu}$  to block 123 lower triangular form. For such reduction, we use the algorithm from [4] which 124 roughly amounts to partitioning the set of dofs into non-overlapping blocks. In the 125 strongly advection dominated case the size of the resulting blocks is small and a 126 block Gauss-Seidel method is an efficient solver. Such techniques have been studied 127 in [5] for conforming methods.

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The idea is to consider the *directed* graph G = (V, E) associated with  $\mathbb{B}^{\nu\nu} \in 130$   $\mathbb{R}^{n_{cr} \times n_{cr}}$ ; G has  $n_{cr}$  vertices labeled  $V = \{1, \dots, n_{cr}\}$  and its set of *edges* E has car-131 dinality equal to the number of nonzero entries of  $\mathbb{B}^{\nu\nu}$ . By definition,  $(i, j) \in E$  iff 132  $b_{ij}^{cr} \neq 0$ . Note that in the advection dominated case, the built-in upwind mechanism 133 results in a non-symmetric sparsity pattern for  $\mathbb{B}^{\nu\nu}$  (see the last two paragraphs of 134 Sect. 3). Thus, we may have  $(i, j) \in E$ , while  $(j, i) \notin E$ . Then, the problem of reducing  $\mathbb{B}^{\nu\nu}$  to block lower triangular form of  $\mathbb{B}^{\nu\nu}$  is equivalent to partitioning G as a 136 union of strongly connected components.

Such partitioning induces non-overlapping partitioning of the set of dofs,  $\mathbf{V} = 138$   $\bigcup_{i=1}^{N_b} \boldsymbol{\omega}_i$ . For  $i=1,\dots,N_b$ , let  $m_i$  denote the cardinality of  $\boldsymbol{\omega}_i$ ; let  $\mathbb{I}_i \in \mathbb{R}^{n_{cr} \times m_i}$  be 139 the matrix that is identity on dofs in  $\boldsymbol{\omega}_i$  and zero otherwise; and  $\mathbb{B}_i^{vv} = \mathbb{I}_i^T \mathbb{B}^{vv} \mathbb{I}_i$  is the 140 block corresponding to the dofs in  $\boldsymbol{\omega}_i$ . The block Gauss–Seidel algorithm reads: Let 141  $\boldsymbol{u}_0^{cr}$  be given, and assume  $\boldsymbol{u}_k^{cr}$  has been obtained. Then  $\boldsymbol{u}_{k+1}^{cr}$  is computed via: For 142  $i=1,\dots N_b$  143

$$\boldsymbol{u}_{k+i/N_b}^{cr} = \boldsymbol{u}_{k+(i-1)/N_b}^{cr} + \mathbb{I}_i(\mathbb{B}_i^{vv})^{-1} \mathbb{I}_i^T \left( \boldsymbol{F} - \mathbb{B}^{vv} \boldsymbol{u}_{k+(i-1)/N_b}^{cr} \right) . \tag{14}$$

As we report in Sect. 5, the action of  $(\mathbb{B}_i^{\nu\nu})^{-1}$  can be computed exactly since in the advection dominated regime the size of the blocks  $\mathbb{B}_i^{\nu\nu}$  is small.

#### **5 Numerical Results**

We present a set of numerical experiments to assess the performance of the proposed block solver. The tests refer to problem (2) with  $\varepsilon = 10^{-3}, 10^{-5}, 10^{-7},$  and  $\Omega$  148 is triangulated with a family of unstructured triangulations  $\mathscr{T}_h$ . In the tables given 149 below J=1 corresponds to the coarsest grid and each refined triangulation on level 150 J, J=2,3,4 is obtained by subdividing each of the  $T\in\mathscr{T}_h$  on level (J-1) into four 151 congruent triangles. From the number of triangles  $n_T$  the total number of dofs for the 152 DG approximation is  $3n_T$ .

**Test 1. Boundary Layer:**  $\Omega = (-1,1)^2$ ,  $\beta = [1,1]^t$ ,  $n_T = 112$  for the coarsest mesh and f is such that the exact solution is given by

$$u(x,y) = \left(x + \frac{1 + e^{-2/\varepsilon} - 2e^{(x-1)/\varepsilon}}{1 - e^{-2/\varepsilon}}\right) \left(y + \frac{1 + e^{-2/\varepsilon} - 2e^{(y-1)/\varepsilon}}{1 - e^{-2/\varepsilon}}\right).$$

**Test 2. Rotating Flow:**  $\Omega = (-1,1)^{\times}(0,1), f = 0 \text{ and } \text{curl}\beta \neq 0,$ 

<sup>&</sup>lt;sup>5</sup> Each dof corresponds to a vertex in the graph; each nonzero entry to an edge.

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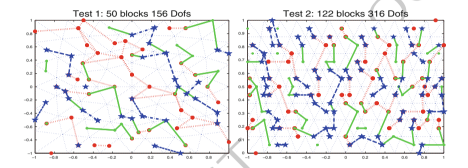
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$$\beta = \begin{bmatrix} 2y(1-x^2) \\ -2x(1-y^2) \end{bmatrix}^t \qquad g(x,y) = \begin{cases} 1 + \tanh(10(2x+1)) & x \le 0, \ y = 0, \\ 0 & \text{elsewhere } . \end{cases}$$

We stress that this test does not fit in the simple description given here, and special 158 care is required (see [3]). For the approximation, for each  $T \in \mathcal{T}_h$ , with barycenter 159  $(x_T, y_T)$ , we use the approximation

$$\beta|_T \approx \nabla \psi|_T$$
 with  $\psi|_T = 2y_T(1-x_T^2)x - 2x_T(1-2y_T^2)y$ ,

and so  $\psi$  is discontinuous. The coarsest grid has  $n_T = 224$  triangles. In Fig. 1 are



**Fig. 1.** Plot of the connected components (*blocks*) of  $\mathbb{B}^{\nu\nu}$  created during Tarjan's algorithm: Test 1 with  $\varepsilon = 10^{-5}$  (left); Test 2 with  $\varepsilon = 10^{-7}$  (right)

represented the plot of the strongly connected components of the graph depicting the 162 blocks for  $\mathbb{B}^{\nu\nu}$  created during Tarjan's algorithm, on the coarsest meshes; for Test 1 163 with  $\varepsilon = 10^{-5}$  (left figure) and for Test 2 with  $\varepsilon = 10^{-7}$  (right figure). We have used different line types (and colors) to distinguish strongly connected components 165 in the directed graph. In Table 1 we report the number of blocks  $N_h$  created during 166 Tarjan's algorithm; the maximum size of the largest such block  $(M_b)$ ; the average 167 block size  $(n_{av})$ ; and the number of block-Gauss-Seidel iterations. After Tarjan's 168 algorithm is used to re-order the matrix  $\mathbb{B}^{\nu\nu}$ , we use the block Gauss-Seidel algorithm (14) where each small block is solved exactly. In the tests that we report here and 170 also in all other similar tests that we have done (with similar advection dominance) 171 the number of block-Gauss-Seidel iterations and the size of the blocks is uniformly 172 bounded with respect to the number of dofs when the advection strongly dominates. 173 Thus, the computational cost for one block Gauss-Seidel iteration in the advection 174 dominated regime is the same as the cost of performing a fixed number of matrix 175 vector multiplications and the algorithm is optimal in such regime.

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		Т	est 1			Test 2						
$\varepsilon$ $J$		1	2	3	4	$\varepsilon$ $J$		1	2	3	4	t1.1
	$ N_b $	44	150	484	1182		$N_b$	31	1	1	1	t1.2
$10^{-3}$	$ M_b $	23	47	95	191	$10^{-3}$	$M_b$	211	1304	5296	21344	t1.3
10	$ n_{av} $	3.55	4.32	5.45	9.02		$ n_{av} $	10.19	1304	5296	21344	t1.4
	iters	7	19	43	166		iters	10	1	1	1	t1.5
	$N_b$	50	210	866	3474		$N_b$	122	468	1822	7106	t1.6
$10^{-5}$	$ M_b $	23	47	95	191	$10^{-5}$	$M_b$	4	4	7	37	t1.7
10	$ n_{av} $	3.12	3.08	3.05	3.07	10	$ n_{av} $	2.59	2.78	2.91	3.00	t1.8
	iters	4	4	4	14		iters	4	4	7	24	t1.9
	$N_b$	50	210	866	3522		$N_b$	122	468	1832	7247	t1.10
$10^{-7}$	$ M_b $	23	47	95	191	$10^{-7}$	$M_b$	4	4	4	6	t1.11
10	$ n_{av} $	3.12	3.08	3.05	3.03	10	$ n_{av} $	2.59	2.78	2.89	2.95	t1.12
	iters	4	4	4	4		iters	4	4	4	4	t1.13

**Table 1.** Number of blocks  $(N_h)$  created during the Tarjan's ordering algorithm, size of largest block  $(M_b)$ , average size of blocks  $(n_{av})$  and number of block-Gauss-Seidel iterations (iters) for Test 1 (left) and Test 2 (right).

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# A Nonoverlapping DD Preconditioner for a Weakly **Over-Penalized Symmetric Interior Penalty Method**

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1 Introduction 13

In this paper we present a nonoverlapping domain decomposition preconditioner for 14 a weakly over-penalized symmetric interior penalty method that is based on balancing domain decomposition by constraints (BDDC) methodology (cf. [2, 5, 7, 8]). The 16 full analysis of the preconditioner can be found in [4].

Let  $\Omega$  be a bounded polygonal domain in  $\mathbb{R}^2$  and  $f \in L_2(\Omega)$ . Consider the following model problem: Find  $u \in H_0^1(\Omega)$  such that

$$\int_{\Omega} \nabla u \cdot \nabla v \, dx = \int_{\Omega} f v \, dx \qquad \forall v \in H_0^1(\Omega). \tag{1}$$

Let  $\mathcal{T}_h$  be a quasi-uniform triangulation of  $\Omega$ , where the mesh parameter h mea- 21 sures the maximum diameter of the triangles in  $\mathcal{T}_h$ , and let

$$V_h = \{ v \in L_2(\Omega) : v | T \in P_1(T) \quad \forall T \in \mathcal{T}_h \}$$

be the discontinuous  $P_1$  finite element function space associated with  $\mathcal{T}_h$ . The model 23 problem (1) can be discretized by the following weakly over-penalized symmetric 24 interior penalty (WOPSIP) method (cf. [3, 9]): 25 Find  $u_h \in V_h$  such that 26

$$a_h(u_h, v) = \int_{\Omega} f v \, dx \qquad v \in V_h,$$

where 27

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$$a_h(v,w) = \sum_{T \in \mathcal{T}_h} \int_T \nabla v \cdot \nabla w \, dx + \sum_{e \in \mathcal{E}_h} \frac{1}{|e|^3} \int_e \Pi_e^0 \llbracket v \rrbracket \cdot \Pi_e^0 \llbracket w \rrbracket \, ds, \tag{2}$$

 $\mathscr{E}_h$  is the set of the edges of  $\mathscr{T}_h$ , |e| is the length of the edge e, [v] denotes the jump of 28 v across the edges, and  $\Pi_e^0$  is the orthogonal projection from  $[L_2(e)]^2$  onto  $[P_0(e)]^2$ . 29  $P_0(e)$  denotes the space of constant functions on the edge e.

For simplicity in presentation, we consider the Poisson model on conforming 31 meshes. But the results can be extended to heterogeneous elliptic problems on non- 32 conforming meshes (cf. [4]). We note that BDDC technique was used in [6] to couple 33 conforming finite element spaces from different subdomains that allows nonmatching meshes across subdomain boundaries, where condition number estimates independent of the coefficients were obtained for heterogeneous elliptic problems. The 36 main difference between [6] and this paper is that the finite element functions in this 37 paper can be discontinuous at the element boundaries.

The rest of the paper is organized as follows. In Sect. 2 we introduce a subspace 39 decomposition. We then design a BDDC preconditioner for the reduced problem in 40 Sect. 3. The condition number estimate is also presented. In Sect. 4 we report numerical results that illustrate the performance of the proposed preconditioner and confirm 42 the theoretical estimates.

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Throughout the paper we will use  $A \lesssim B$  and  $A \gtrsim B$  to represent the statements 44 that  $A \leq (\text{constant})B$  and  $A \geq (\text{constant})B$ , where the positive constant is independent of the mesh size, the subdomain size, and the number of subdomains. The statement 46  $A \approx B$  is equivalent to  $A \lesssim B$  and  $A \gtrsim B$ .

## 2 A Subspace Decomposition

In this section we propose an intermediate preconditioner for the WOPSIP method, 49 which is based on a subspace decomposition.

Let  $\Omega_1, \ldots, \Omega_J$  be a nonoverlapping partition of  $\Omega$  aligned with  $\mathcal{T}_h$  and  $\Gamma = 51$  $(\bigcup_{i=1}^J \partial \Omega_i) \setminus \partial \Omega$  be the interface of the subdomains. We assume that the subdomains are shape regular polygons (cf. [1, Sect. 7.5]). We denote the diameter of  $\Omega_j$  53 by  $H_i$  and define H to be  $\max_{1 \le j \le J} H_i$ .  $\mathcal{E}_{h,\Gamma}$  is the subset of  $\mathcal{E}_h$  containing the edges 54 on  $\Gamma$ .

First we decompose  $V_h$  into two subspaces as follows:

$$V_h = V_{hC} \oplus V_{hD}$$

where 57

 $V_{h,C} = \{v \in V_h : [v] = 0 \text{ at the midpoints of the edges on the boundaries } \}$ of the subdomains \}.

 $V_{h,D} = \{v \in V_h : \{\{v\}\}\} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = \{v \in V_h : \{\{v\}\}\}\} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = \{v \in V_h : \{\{v\}\}\}\} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = \{v \in V_h : \{\{v\}\}\}\} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = \{v \in V_h : \{\{v\}\}\}\} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \text{ and } V_{h,D} = 0 \text{ at the midpoints of the edges in } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoints of } \mathcal{E}_{h,D} = 0 \text{ at the midpoint$ v = 0 at the midpoints of the edges in  $\Omega \setminus \Gamma$  .

Here  $\{v\}$  denotes the average of v from the two sides of an edge in  $\mathcal{E}_{h,\Gamma}$ . Let  $A_h: V_h \longrightarrow V_h'$  be the symmetric positive-definite (SPD) operator defined by 59

$$\langle A_h v, w \rangle = a_h(v, w) \quad \forall v, w \in V_h,$$

where  $\langle \cdot, \cdot \rangle$  is the canonical bilinear form between a vector space and its dual. Similarly, we define  $A_{h,D}: V_{h,D} \longrightarrow V'_{h,D}$  and  $A_{h,C}: V_{h,C} \longrightarrow V'_{h,C}$  by

$$\langle A_{h,D}v, w \rangle = a_h(v, w) \qquad \forall v, w \in V_{h,D},$$

$$\langle A_{h,C}v, w \rangle = a_h(v, w) \qquad \forall v, w \in V_{h,C}.$$
(3)

$$\langle A_{h,C}v,w\rangle = a_h(v,w) \qquad \forall v,w \in V_{h,C}.$$
 (4)

Given any  $v \in V_h$ , we have a unique decomposition  $v = v_D + v_C$  where  $v_D \in V_{h,D}$  62 and  $v_C \in V_{h,C}$ . Then based on the definitions of the subspaces  $V_{h,D}$  and  $V_{h,C}$ , it can be 63 shown that

$$\langle A_h v, v \rangle \approx \langle A_{h,D} v_D, v_D \rangle + \langle A_{h,C} v_C, v_C \rangle \qquad \forall v \in V_h.$$
 (5)

Remark 1. Since functions in  $V_{h,C}$  are continuous at the midpoints of the edges in 65  $\mathcal{E}_{h,\Gamma}$ , we have

$$a_h(v,w) = \sum_{j=1}^{J} a_{h,j}(v_j, w_j) \qquad \forall v, w \in V_{h,C},$$
 (6)

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where  $v_j = v|_{\Omega_i}$ ,  $w_j = w|_{\Omega_i}$  and

$$a_{h,j}(v_j, w_j) = \sum_{\substack{T \in \mathscr{T}_h \\ T \subset \Omega_j}} \int_T \nabla v_j \cdot \nabla w_j \, dx + \sum_{\substack{e \in \mathscr{E}_h \\ e \subset \Omega_j}} \frac{1}{|e|^3} \int_e \Pi_e^0 \llbracket v_j \rrbracket \cdot \Pi_e^0 \llbracket w_j \rrbracket \, ds. \tag{7}$$

Note that the second sum on the right-hand side of (7) is over the edges interior to  $\Omega_i$  68 and therefore  $a_{h,j}(\cdot,\cdot)$  is a localized bilinear form. The introduction of the subspace 69 decomposition where the bilinear form can be localized as shown in (6) and (7) is 70 the key ingredient in designing our preconditioner in Sect. 3. 71

Next we decompose  $V_{h,C}$  into two subspaces  $V_{h,C}(\Omega \setminus \Gamma)$  and  $V_{h,C}(\Gamma)$  defined as 72 follows: 73

$$V_{h,C}(\Omega \setminus \Gamma) = \{ v \in V_{h,C} : v = 0 \text{ at all the midpoints of the edges in } \mathcal{E}_{h,\Gamma} \},$$
  
$$V_{h,C}(\Gamma) = \{ v \in V_{h,C} : a_h(v,w) = 0 \quad \forall w \in V_{h,C}(\Omega \setminus \Gamma) \}.$$

The space  $V_{h,C}(\Gamma)$  is the space of discrete harmonic functions, which are uniquely 74 determined by their values at the midpoints of the edges in  $\mathcal{E}_{h,\Gamma}$ . 75

Let the SPD operators  $A_{h,\Omega\setminus\Gamma}:V_{h,C}(\Omega\setminus\Gamma)\longrightarrow V_{h,C}(\Omega\setminus\Gamma)'$  and  $S_h:V_{h,C}(\Gamma)\longrightarrow$ 76  $V_{h,C}(\Gamma)'$  be defined by 77

$$\langle A_{h,\Omega \setminus \Gamma} v, w \rangle = a_h(v, w) \qquad \forall v, w \in V_{h,C}(\Omega \setminus \Gamma),$$
  
 $\langle S_h v, w \rangle = a_h(v, w) \qquad \forall v, w \in V_{h,C}(\Gamma).$ 

Note that given any  $v_C \in V_{h,C}$ , we have a unique decomposition  $v_C = v_{C,\Omega \setminus \Gamma} + v_{C,\Gamma}$  78 where  $v_{C,\Omega \setminus \Gamma} \in V_{h,C}(\Omega \setminus \Gamma)$  and  $v_{C,\Gamma} \in V_{h,C}(\Gamma)$ . It follows from the definitions of 79  $V_{h,C}(\Omega \setminus \Gamma)$  and  $V_{h,C}(\Gamma)$  that

$$\langle A_{h,C}v_C, v_C \rangle = \langle A_{h,\Omega \setminus \Gamma}v_{C,\Omega \setminus \Gamma}, v_{C,\Omega \setminus \Gamma} \rangle + \langle S_hv_{C,\Gamma}, v_{C,\Gamma} \rangle \qquad \forall v_C \in V_{h,C}. \tag{8}$$

Based on the relations (5) and (8), we define a preconditioner  $B_1: V_h' \longrightarrow V_h$  for 81  $A_h$  by

 $B_1 = I_D A_{h,D}^{-1} I_D^t + I_{h,\Omega \setminus \Gamma} A_{h,\Omega \setminus \Gamma}^{-1} I_{h,\Omega \setminus \Gamma}^t + I_{\Gamma} S_h^{-1} I_{\Gamma}^t,$ 

where  $I_D: V_{h,D} \longrightarrow V_h$ ,  $I_{h,\Omega \setminus \Gamma}: V_{h,C}(\Omega \setminus \Gamma) \longrightarrow V_h$ , and  $I_\Gamma: V_{h,C}(\Gamma) \longrightarrow V_h$  are so natural injections.

It follows from (5) and (8) that

$$\kappa(B_1 A_h) = \frac{\lambda_{\max}(B_1 A_h)}{\lambda_{\min}(B_1 A_h)} \approx 1. \tag{9}$$

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Remark 2. Let us observe the properties of the preconditioner  $B_1$  from the implementational point of view. First it is easy to implement the solve  $A_{h,D}^{-1}$  because  $A_{h,D}$  87 is a block diagonal matrix with small blocks. Next in view of (6) and (7), the solve 88  $A_{h,\Omega\setminus\Gamma}^{-1}$  can be implemented by solving independent subdomain problems in parallel. On the other hand, noting that  $S_h$  is a global solve, we need to design a good 90 preconditioner for  $S_h$  in order to obtain a good parallel preconditioner for  $A_h$ .

### 3 A BDDC Preconditioner

In this section we propose a preconditioner for the Schur complement operator  $S_h$  93 based on the BDDC methodology. 94

Let  $V_{h,j}$  be the space of discontinuous  $P_1$  finite element functions on  $\Omega_j$  that 95 vanish at the midpoints of the edges on  $\partial \Omega_j \cap \partial \Omega$ , and  $V_h(\Omega_j)$  be the subspace of 96  $V_{h,j}$  whose members vanish at the midpoints of the edges on  $\partial \Omega_j$ . We denote by  $\mathscr{H}_j$  97 the space of local discrete harmonic functions defined by

$$\mathcal{H}_j = \left\{ v \in V_{h,j} : a_{h,j}(v,w) = 0 \quad \forall w \in V_h(\Omega_j) \right\}.$$

The space  $\mathcal{H}_m$  is defined by gluing the spaces  $\mathcal{H}_j$  together along the interface 99  $\Gamma$  through enforcing the continuity of the mean values on the common edges of 100 subdomains:

$$\mathcal{H}_m = \{ v \in L_2(\Omega) : v_j = v |_{\Omega_j} \in \mathcal{H}_j \text{ for } 1 \le j \le J$$

$$\text{and } \int_{\partial \Omega_i \cap \partial \Omega_k} v_j \, ds = \int_{\partial \Omega_i \cap \partial \Omega_k} v_k \, ds \text{ for } 1 \le j, k \le J \},$$

and we equip  $\mathcal{H}_m$  with the bilinear form

$$a_h^m(v, w) = \sum_{1 \le i \le I} a_{h,j}(v_j, w_j).$$

Let  $\mathscr{E}_H$  be the set of the edges of the subdomains  $\Omega_1, \dots, \Omega_J$ . The BDDC preconditioner is based on a decomposition of  $\mathcal{H}_m$  into orthogonal subspaces with respect to  $a_h^m(\cdot,\cdot)$ : 105

$$\mathscr{H}_m = \mathring{\mathscr{H}} \oplus \mathscr{H}_0, \tag{10}$$

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where

$$\mathring{\mathscr{H}} = \left\{ v \in \mathscr{H}_m : \int_E v \, ds = 0 \quad \forall E \in \mathscr{E}_H \right\}$$

and

$$\mathcal{H}_0 = \left\{ v \in \mathcal{H}_m : a_h^m(v, w) = 0 \quad \forall w \in \mathring{\mathcal{H}} \right\}. \tag{11}$$

Then we equip  $\mathcal{H}_0$  and the localized subspaces  $\mathring{\mathcal{H}}_j$   $(1 \leq j \leq J)$  of  $\mathring{\mathcal{H}}$ : 108

$$\mathring{\mathscr{H}}_{j}=\left\{v\in\mathscr{H}_{j}:\int_{E}vds=0\text{ for all the edges }E\text{ of }\Omega_{j}\right\},$$

with the SPD operators  $S_0: \mathscr{H}_0 \longrightarrow \mathscr{H}'_0$  and  $S_j: \mathring{\mathscr{H}}_j \longrightarrow \mathring{\mathscr{H}}'_j$  defined by

$$\langle S_0 v, w \rangle = a_h^m(v, w) \qquad \forall v, w \in \mathcal{H}_0,$$
 (12)

$$\langle S_0 v, w \rangle = a_h^m(v, w) \qquad \forall v, w \in \mathcal{H}_0,$$

$$\langle S_j v, w \rangle = a_{h,j}(v, w) \qquad \forall v, w \in \hat{\mathcal{H}}_j.$$
(12)

Note that  $V_{h,C}(\Gamma)$  is a subspace of  $\mathscr{H}_m$  and there exists a projection  $P_{\Gamma}:\mathscr{H}_m\to$  $V_{h,C}(\Gamma)$  defined by averaging: 112

$$(P_{\Gamma}v)(m_e) = \{\{v\}\}\} (m_e) \quad \forall e \in \mathcal{E}_{h,\Gamma},$$

where  $m_e$  is the midpoint of e. The operator  $P_{\Gamma}$  connects the BDDC preconditioner based on  $\mathcal{H}_m$  to the Schur complement operator  $S_h$  on  $V_{h,C}(\Gamma)$ .

We can now define the BDDC preconditioner  $B_{BDDC}: V_{h,C}(\Gamma)' \longrightarrow V_{h,C}(\Gamma)$  for 115 the Schur complement operator  $S_h: V_{h,C}(\Gamma) \longrightarrow V_{h,C}(\Gamma)'$  as follows: 116

$$B_{BDDC} = (P_{\Gamma}I_0) S_0^{-1} (P_{\Gamma}I_0)^t + \sum_{i=1}^{J} (P_{\Gamma}\mathbb{E}_j) S_j^{-1} (P_{\Gamma}\mathbb{E}_j)^t,$$

where  $I_0$  is the natural injection of  $\mathscr{H}_0$  into  $\mathscr{H}_m$  and  $\mathbb{E}_j: \mathring{\mathscr{H}}_j \longrightarrow \mathring{\mathscr{H}}$  is the trivial extension defined by 118

$$\mathbb{E}_{j}\mathring{v}_{j} = \begin{cases} \mathring{v}_{j} & \text{on } \Omega_{j} \\ 0 & \text{on } \Omega \setminus \Omega_{j} \end{cases} \quad \forall \mathring{v}_{j} \in \mathring{\mathscr{H}}_{j}.$$

We then obtain the preconditioner  $B_2: V_h' \longrightarrow V_h$  for  $A_h$  by replacing the global 119 solve  $S_h^{-1}$  in (2) with the preconditioner  $B_{BDDC}$ : 120

$$B_2 = I_D A_{h,D}^{-1} I_D^t + I_{h,\Omega \setminus \Gamma} A_{h,\Omega \setminus \Gamma}^{-1} I_{h,\Omega \setminus \Gamma}^t + I_{\Gamma} B_{BDDC} I_{\Gamma}^t.$$

We can analyze the condition number of  $B_{BDDC}S_h$  by the theory of additive 121 Schwarz preconditioners (cf. [1, 10, 11], and the references therein). The proof of 122 the following result can be found in [4]. 123

**Lemma 1.** We have the following bounds for the eigenvalues of  $B_{BDDC}S_h$ 

$$\lambda_{\min}(B_{BDDC}S_h) \ge 1,$$

$$\lambda_{\max}(B_{BDDC}S_h) \lesssim \left(1 + \ln\frac{H}{h}\right)^2.$$

Combining (5), (8) and Lemma 1, we have the following estimate of the condition 125 number of the preconditioned system  $B_2A_h$ .

**Theorem 1.** There exists a positive constant C, independent of h, H and J, such that

$$\kappa(B_2 A_h) = \frac{\lambda_{\max}(B_2 A_h)}{\lambda_{\min}(B_2 A_h)} \le C \left(1 + \ln \frac{H}{h}\right)^2. \tag{128}$$

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### 4 Numerical Results

In this section we present some numerical results that illustrate the performance of  $^{130}$  the preconditioners  $B_1$  and  $B_2$ .

We consider the model problem (1) on the unit square  $(0,1)^2$  with the exact solution  $u(x,y)=y(1-y)\sin(\pi x)$ . We use a uniform triangulation  $\mathcal{T}_h$  of isosceles right triangles, where the mesh parameter h represents the length of the horizontal/vertical edges. The domain  $\Omega$  is divided into J nonoverlapping squares aligned with  $\mathcal{T}_h$  and the length of the horizontal/vertical edges of the squares is denoted by H. The discrete problem obtained by the WOPSIP method is solved by the preconditioned conjugate gradient method. The iteration is stopped when the relative residual is less than  $10^{-6}$ .

Numerical results for the preconditioners  $B_1$  and  $B_2$  are presented in Table 1, 140 which confirm the theoretical estimates in (9) and Theorem 1.

**Table 1.** Results for the preconditioners  $B_1$  and  $B_2$  with  $J = 4^2$ 

h	H/h	h	$B_1A_h$		$B_2A_h$			
	/ .	κ	$\lambda_{\min}$	$\lambda_{\max}$	κ	$\lambda_{\min}$	$\lambda_{\max}$	
			8.2624e-1					
			9.1258e-1					
$2^{-5}$	8	1.0919	9.5608e-1	1.0439	2.3215	9.5673e-1	2.2211	
$2^{-6}$	16	1.0433	9.7880e-1	1.0212	3.0490	9.7994e-1	2.9879	

We present in Table 2 the iteration counts and total time to solution for a parallel mplementation of our preconditioner. For comparison, results on a single processor of the same machine without preconditioning are also presented for J=1. The three operations  $A_{h,D}^{-1}, A_{h,\Omega\setminus\Gamma}^{-1}$ , and  $B_{BDDC}$  are performed one after the other, sequentially, 145

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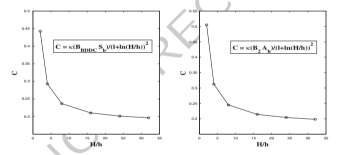
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but each of these operators is evaluated in parallel on the decomposed domain with 146 one subdomain per processor. Iteration counts are consistent with our theory and 147 confirm again that the method is scalable, and the running times show good parallel 148 speedup for large problems.

**Table 2.** Parallel performance of the preconditioner  $B_2$ 

		J=1	J =	$=4^2, H=2^{-2}$	J	$=8^2, H=2^{-3}$ $J=16^2, H=2^{-4}$	t2.1
	Its	Wall clock tim	ne Its V	Wall clock time	Its	Wall clock time Its Wall clock time	t2.2
$2^{-6}$	235	0.46	7	0.37	7	0.5 5 1.14	t2.3
_	450	3.75	8	2.22	8	1.06 6 1.96	t2.4
	884		9	20.12	8	4.35 6 2.71	t2.5
$2^{-9}$	1786	319.0	8	126.15	8	27.15 7 7.81	t2.6

The numbers  $\kappa(B_2A_h)/(1+\ln(H/h))^2$  and  $\kappa(B_{BDDC}S_h)/(1+\ln(H/h))^2$  are 150 plotted against H/h in Fig. 1. As H/h increases these two numbers settle down to 151 around 0.2, which indicates that the estimates in Lemma 1 and Theorem 1 are sharp. 152



**Fig. 1.** Left figure: the behavior of  $C = \kappa (B_{BDDC}S_h)/(1 + \ln(H/h))^2$  for the BDDC preconditioner; right figure: the behavior of  $C = \kappa (B_2 A_h) / (1 + \ln(H/h))^2$  for the preconditioner

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# Sharp Condition Number Estimates for the Symmetric 2 2-Lagrange Multiplier Method

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Summary. Domain decomposition methods are used to find the numerical solution of large 9 boundary value problems in parallel. In optimized domain decomposition methods, one solves 10 a Robin subproblem on each subdomain, where the Robin parameter a must be tuned (or optimized) for good performance. We show that the 2-Lagrange multiplier method can be analyzed 12 using matrix analytical techniques and we produce sharp condition number estimates.

1 Introduction 14

Consider the model problem

$$-\Delta u = f \text{ in } \Omega \text{ and } u = 0 \text{ on } \partial \Omega, \tag{1}$$

where  $\Omega$  is the domain, f is a given forcing and  $u \in H_0^1(\Omega)$  is the unknown solution. 16 In the present paper, we describe a symmetric 2-Lagrange multiplier (S2LM) domain 17 decomposition method to solve elliptic problems such as (1). When we discretize (1) 18 using e.g. piecewise linear finite elements, we obtain a linear system of the form

$$A\mathbf{u} = \mathbf{f},\tag{2}$$

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where  $\mathbf{u} \in \mathbb{R}^n$  is the finite element coefficient vector of the approximation to the 20 solution u of (1).

We now consider the domain decomposition [9]  $\Omega = \Gamma \cup \Omega_1 \cup ... \cup \Omega_p$ , where 22  $\Omega_1, \dots, \Omega_p$  are the (open, disjoint) "subdomains" and  $\Gamma = \Omega \cap \bigcup_{k=1}^p \partial \Omega_k$  is the "ar-23 tificial interface". We introduce the "local problems"

$$\begin{cases} -\Delta u_k = f & \text{in } \Omega_k, \quad \text{(PDE)} \\ u_k = 0 & \text{on } \partial \Omega_k \cap \partial \Omega, \quad \text{(natural b.c.)} \\ (a + D_v)u_k = \lambda_k & \text{on } \partial \Omega_k \cap \Gamma, \quad \text{(artificial b.c.)} \end{cases}$$
 (3)

where a > 0 is the Robin tuning parameter and k = 1, ..., p and  $D_v$  denotes the 25 directional derivative in the outwards pointing normal  $\nu$  of  $\partial \Omega_k$ . The interface  $\Gamma$  is 26

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artificial in that it is not a natural part of the "physical problem" (1) but instead is 27 introduced purely for the purpose of calculation.

We again discretize the systems (3) using a finite element method. The Robin b.c. 29 in (3) gives rise to a mass matrix on the interface  $\Gamma \cap \partial \Omega_k$ , which we lump. If the 30 grid is uniform, this mass matrix is aI (we absorb any h factors into the a coefficient) 31 - we make this simplification for the remainder of the present paper.

$$\begin{bmatrix}
A_{IIk} & A_{I\Gamma k} \\
A_{\Gamma Ik} & A_{\Gamma \Gamma k} + aI
\end{bmatrix}
\underbrace{\begin{bmatrix}
\mathbf{u}_{lk} \\
\mathbf{u}_{\Gamma k}
\end{bmatrix}}_{\mathbf{u}_{rk}} = \underbrace{\begin{bmatrix}
\mathbf{f}_{lk} \\
\mathbf{f}_{\Gamma k}
\end{bmatrix}}_{\mathbf{f}_{k}} + \begin{bmatrix}
0 \\
\boldsymbol{\lambda}_{k}
\end{bmatrix}.$$
(4)

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Here, we have used the suggestive subscripts I for interior nodes and  $\Gamma$  for the artificial interface nodes.

The FETI-2LM algorithm was introduced in [4] for cases without cross-points, 35 while the general case including cross points was introduced and analyzed in [7]. 36 The method consists of finding the value of  $\lambda = [\lambda_1^T, \dots, \lambda_n^T]^T$  which yields solutions 37  $\mathbf{u}_1, \dots, \mathbf{u}_p$  to (4) in such a way that  $\mathbf{u}_1, \dots, \mathbf{u}_p$  meet continuously across  $\Gamma$  and glue 38 together into the unique solution  $\mathbf{u}$  of (2).

The main result of the present paper is a new estimate of the condition number 40 of FETI-2LM algorithms using matrix analytical techniques. This new idea produces 41 sharp condition number estimates with much more straightforward proof techniques 42 than the techniques used in [7] (where the estimates are not sharp). As a result, the 43 present paper is a logical follow-up to [7].

The present paper focuses on 1-level algorithms which are known not to scale. 45 Scalable algorithms are considered in [8] and [3].

Our paper is organized as follows. In Sect. 2, we give the symmetric 2-Lagrange 47 multiplier method for general domains with cross points. In Sect. 3, we give spectral 48 estimates including our main result, Theorem 1, on the condition number of the symmetric 2-Lagrange multiplier system. In Sect. 4, we verify this Theorem with some 50 numerical experiments.

### 2 The Symmetric 2-Lagrange Multiplier Method

We now describe the 2-Lagrange multiplier method that we analyze in the present 53 paper. Consider the local problems (4) and eliminate the interior degrees of freedom 54 to obtain the relation 55

$$a\begin{bmatrix} \mathbf{u}_{G} \\ \vdots \\ \mathbf{u}_{\Gamma p} \end{bmatrix} = \begin{bmatrix} a(S_{1} + aI)^{-1} \\ & \ddots \\ & a(S_{p} + aI)^{-1} \end{bmatrix} \begin{pmatrix} \mathbf{g} \\ \vdots \\ \mathbf{g}_{p} \end{pmatrix} + \begin{bmatrix} \lambda_{1} \\ \vdots \\ \lambda_{p} \end{bmatrix} \end{pmatrix}, \quad (5)$$

where

$$S_k = A_{\Gamma\Gamma k} - A_{\Gamma Ik} A_{IIk}^{-1} A_{I\Gamma k}$$
 and  $\mathbf{g}_k = \mathbf{f}_{\Gamma k} - A_{\Gamma Ik} A_{IIk}^{-1} \mathbf{f}_{Ik}$  57

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are the "Dirichlet-to-Neumann maps" and "accumulated right-hand-sides" and where 58  $\mathbf{u}_{\Gamma j}$  denotes those degrees of freedom of the local solution  $\mathbf{u}_j$  associated with the artificial interface  $\Gamma$ .

The matrices  $S_k$  are symmetric and semidefinite. Since  $Q = a(S+aI)^{-1}$ , we find 61 that the spectrum  $\sigma(Q)$  is contained in the set  $[\varepsilon, 1-\varepsilon] \cup \{1\}$  for some  $\varepsilon > 0$ . The 62 eigenvalue 1 of Q comes from the kernel of S and hence the kernel of Q-I is spanned 63 by the indicating functions of the subdomains that "float".

### 2.1 Relations Between (4) and (2) and Continuity

We define the boolean restriction matrix  $R_k$  by selecting rows of the  $n \times n$  identity 66 matrix corresponding to those vertices of  $\Omega$  that are in  $\bar{\Omega}_k \cap \Omega$ . As a result, from 67 a finite element coefficient vector  $\mathbf{v}$  corresponding to a finite element function  $v \in 68$   $H_0^1(\Omega)$ , we can define a finite element coefficient vector  $\mathbf{v}_k = R_k \mathbf{v}$ , which corresponds 69 to a finite element function  $v \in H^1(\Omega_k) \cap H_0^1(\Omega)$ , which is obtained by restricting v 70 to  $\Omega_k$ .

The identity  $\int_{\Omega} = \sum_{k=1}^{p} \int_{\Omega_k}$  induces the following relations between (4) and (2): 72

$$A = \sum_{k=1}^{p} R_k^T \overbrace{\begin{bmatrix} A_{IIk} & A_{I\Gamma k} \\ A_{\Gamma Ik} & A_{\Gamma \Gamma k} \end{bmatrix}}^{A_{Nk}} R_k \quad \text{and} \quad \mathbf{f} = \sum_{k=1}^{p} R_k^T \mathbf{f}_k.$$
 (6)

Each interface vertex  $\mathbf{x}_i \in \Gamma$  is adjacent to  $m_i \geq 2$  subdomains. As a result, the 73 "many-sided trace"  $\mathbf{u}_G$  defined by (5) contains  $m_i$  entries corresponding to  $\mathbf{x}_i$ , one per 74 subdomain adjacent to  $\mathbf{x}_i$ . We define the orthogonal projection matrix K which averages function values for each interface vertex  $\mathbf{x}_i$ . A many-sided trace  $\mathbf{u}_G$  corresponds 76 to local functions  $\mathbf{u}_1, \ldots, \mathbf{u}_p$  that meet continuously across  $\Gamma$  if and only if 77

$$K\mathbf{u}_G = \mathbf{u}_G. \tag{7}$$

**2.2** A Problem in  $\lambda$ 

The **symmetric 2-Lagrange multiplier** (S2LM) system is given by

$$(Q - K)\lambda = -Q\mathbf{g}. (8)$$

We further let E be the orthogonal projection onto the kernel of Q-I.

**Lemma 1.** Assume that 
$$||EK|| < 1$$
. The problem (2) is equivalent to (8).

*Proof.* In order to solve (2) using local problems (4), one should find Robin boundary values  $\lambda_1, \ldots, \lambda_p$  which result in local solutions  $\mathbf{u}_1, \ldots, \mathbf{u}_p$  that meet continusously across  $\Gamma$ . As a result, we impose the condition (7), which we multiply by 84 a > 0 and convert to an expression in  $\lambda$  using (5) to obtain  $Ka(S+aI)^{-1}(\lambda + \mathbf{g}) = 85$   $a(S+aI)^{-1}(\lambda + \mathbf{g})$  or

$$(I - K)Q\lambda = (K - I)Q\mathbf{g} \tag{9}$$

With this continuity condition, there is clearly a unique  $\mathbf{u}$  which restricts to the  $\mathbf{u}_i$ :

$$\mathbf{u}_{i} = R_{i}\mathbf{u}, \quad j = 1, \dots, p. \tag{10}$$

Imposing continuity is not sufficient, we must also ensure that the "fluxes" match. 88 Indeed, if we impose on the solution  $\bf u$  of (10) that the Eq. (2) should hold, one 89 obtains

$$\mathbf{f} = A\mathbf{u} \stackrel{(6)}{=} \sum_{j=1}^{p} R_j^T A_{Nj} R_j \mathbf{u} \stackrel{(10)}{=} \sum_{j=1}^{p} R_j^T A_{Nj} \mathbf{u}_j$$
(11)

$$\stackrel{(4),(6)}{=} \mathbf{f} + \sum_{j=1}^{p} R_{j}^{T} \begin{pmatrix} 0 \\ \boldsymbol{\lambda}_{j} - a \mathbf{u}_{\Gamma j} \end{pmatrix}$$
 (12)

Canceling the **f** terms on each side and multiplying by K, we obtain  $K\lambda - Ka\mathbf{u}_G = 0$ . 91 Using (5), we obtain

$$K(Q-I)\lambda = -KQ\mathbf{g}. (13)$$

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We add (9) and (13) to obtain (8).

To see that the solution of (8) is unique, observe that the ranges of E and K intersect trivially by the hypothesis that ||EK|| < 1. As a result, the eigenspace of Q of eigenvalue 1 intersects trivially with the range of K and Q - K is nonsingular.  $\square$ 

We will further discuss the choice of the parameter a in Sect. 3.1.

### 3 Spectral Estimates

If we use GMRES or MINRES on the symmetric indefinite system (8), the residual 96 norm can be estimated as a function of the condition number of Q - K, cf. [2]. In 97 order to estimate the condition number of Q - K, we begin by giving a canonical 98 form for the pair of projections E and K. 99

**Lemma 2.** Let E and K be orthogonal projections. There is a choice of orthonormal 100 basis that block diagonalizes E and K simultaneously and such that the blocks  $E_k$  101 and  $K_k$  of E and K satisfy 102

$$E_k \in \left\{0, 1, \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix}\right\} \quad and \quad K_k \in \left\{0, 1, \begin{bmatrix} c_k^2 & c_k s_k \\ c_k s_k & s_k^2 \end{bmatrix}\right\},\tag{14}$$

where  $c_k = \cos \theta_k > 0$ ,  $s_k = \sin \theta_k > 0$  and  $\theta_k \in (0, \pi/2)$  is a "principal angle" 103 relating E and K.

The canonical form (14) can be obtained from the CS decomposition [1] by start-  $_{105}$  ing from E = diag(I,0) and picking orthonormal bases for the range and kernel of  $_{106}$  K. Due to space constraints, we omit this argument.

We also give a technical lemma which describes the spectrum of a sum of certain symmetric matrices. 108

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**Lemma 3.** Let X, Y be symmetric matrices of dimensions  $m \times m$ . Let  $0 < y_{\min} < y_{\max}$ and assume that  $|\sigma(Y)| \subset [y_{\min}, y_{\max}]$ . Denote by  $\rho(X)$  the spectral radius of X and 111 assume that  $\rho(X) < y_{\min}$ . Then, 112

$$|\sigma(X+Y)| \subset [y_{\min} - \rho(X), y_{\max} + \rho(X)]. \tag{15}$$

*Proof.* This follows from a Theorem of Weyl [5, Theorem 4.3.1, pp. 181−182]. □

### **3.1** Condition Number of Q - K

We now come to our main result.

**Theorem 1.** Let  $\varepsilon > 0$ . Assume that  $\sigma(Q) \subset [\varepsilon, 1 - \varepsilon] \cup \{1\}$ . Let E, K be orthogonal 115 projections and assume that ||EK|| < 1. Then we have the sharp estimates 116

$$|\sigma(Q - K)| \subset \left[\frac{\varepsilon + \sqrt{(1+\varepsilon)^2 - 4\|EK\|^2 \varepsilon} - 1}{2}, 1\right], \quad and$$

$$\kappa(Q - K) \leq \frac{2}{\varepsilon + \sqrt{(1+\varepsilon)^2 - 4\|EK\|^2 \varepsilon} - 1} = O((1 - \|EK\|)^{-1} \varepsilon^{-1}).$$
(17)

$$\kappa(Q - K) \le \frac{2}{\varepsilon + \sqrt{(1 + \varepsilon)^2 - 4\|EK\|^2 \varepsilon - 1}} = O((1 - \|EK\|)^{-1} \varepsilon^{-1}). \tag{17}$$

*Proof.* Let  $X = Q - \frac{1}{2}I - \varepsilon E$  and  $Y = \frac{1}{2}I + \varepsilon E - K$ . Then, Q - K = X + Y and we are in a position to use Lemma 3. We now estimate the spectral properties of X and Y.

**Spectral properties of X:** Recall that E projects onto the eigenspace of Q with 119 eigenvalue 1. As a result, after some orthonormal change of basis, we find that Q = 120 $\operatorname{diag}(Q_0, I)$  and  $E = \operatorname{diag}(0, I)$  and hence 121

$$\rho(X) \le \frac{1}{2} - \varepsilon. \tag{18}$$

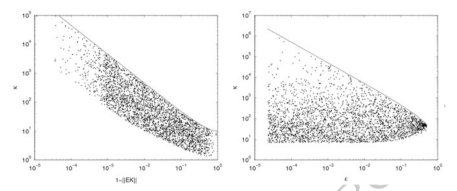
**Spectral properties of Y:** Lemma 2 shows that E and K block diagonalize si- 122 multaneously and Y is also block diagonal in the same basis. Using (14), we find that the kth block  $Y_k$  of Y is given by 124

$$Y_{k} = \begin{cases} \frac{1}{2} & \text{if } E_{k} = K_{k} = 0, \\ -\frac{1}{2} & \text{if } E_{k} = 0, K_{k} = 1, \\ \frac{1}{2} + \varepsilon & \text{if } E_{k} = 1, K_{k} = 0, \\ \begin{bmatrix} \frac{1}{2} + \varepsilon - c_{k}^{2} - c_{k} s_{k} \\ -c_{k} s_{k} & \frac{1}{2} - s_{k}^{2} \end{bmatrix} & \text{otherwise;} \end{cases}$$

$$(19)$$

where the case  $E_k = K_k = 1$  is excluded by the hypothesis that ||EK|| < 1. As a result, 125 the eigenvalues of  $Y_k$  are in the set  $\{\pm \frac{1}{2}, \frac{1}{2} + \varepsilon, \lambda_{\pm}(c_k^2)\}$ , where 126

$$\lambda_{\pm}(c_k^2) = \frac{\varepsilon \pm \sqrt{(1+\varepsilon)^2 - 4c_k^2 \varepsilon}}{2}.$$
 (20)



**Fig. 1.** Comparing random Q - K (points) versus the estimate (17) (solid). Left:  $\varepsilon = 0.1$ , varying ||EK||, 3,000 repetitions. Right: ||EK|| = 0.99, varying  $\varepsilon$ , 3,000 repetitions

Note that  $||EK|| = \sqrt{\rho(EKE)} = \max_k c_k$  and that the functions  $\lambda_{\pm}(c_k^2)$  are monotonic in  $c_k^2$ . Hence, we find the following bounds for the modulus of an eigenvalue of 128 Y:

$$|\sigma(Y)| \subset \left[ \frac{\sqrt{(1+\varepsilon)^2 - 4\|EK\|^2 \varepsilon} - \varepsilon}{2}, \frac{1}{2} + \varepsilon \right]. \tag{21}$$

Combining (15), (18), and (21) gives (16).

The examples  $Q = \text{diag}(1, 1 - \varepsilon)$  and  $K = \begin{bmatrix} c^2 & c\sqrt{1 - c^2} \\ c\sqrt{1 - c^2} & 1 - c^2 \end{bmatrix}$  for c = 0 and c = ||EK|| give the extreme eigenvalues of (21) and hence our estimates are sharp.

In view of Theorem 1, the Robin parameter a should be chosen so as to make 131  $\varepsilon$  as large as possible. This occurs precisely when a is the geometric mean of the 132 extremal positive eigenvalues of S. More details can be found in [7].

### 4 Numerical Verification

We verify numerically the validity of Theorem 1 by generating random  $5 \times 5$  matrices 135 Q and E as follows. We set  $Q = \operatorname{diag}(\varepsilon, q, 1 - \varepsilon, 1, 1)$  where q is chosen randomly 136 between  $\varepsilon$  and  $1 - \varepsilon$ . We generate randomly a 2-dimensional space and set K to be the 137 orthogonal projection onto that space. We compare the resulting condition number 138  $\kappa = \kappa(Q - K)$  against (17), cf. Fig. 1.

We observe that our estimates are correct and sharp for such "generic" random 140 matrices, although some "lucky" random matrices produce much milder condition 141 numbers than our estimates.

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5 Conclusions 143

We have analyzed a domain decomposition method with optimized Robin boundary 144 conditions. Our estimates rely on new matrix analytical techniques and are sharp. By 145 further estimating the quantities ||EK|| and  $\varepsilon$  (cf. [7]) our estimates are consistent 146 with and generalize the estimates calculated using Fourier transforms in the opti- 147 mized Schwarz literature (e.g. [6]). An upcoming paper [8] will further analyze the 148 weak scaling property of a 2-level algorithm and large-scale implementations are 149 being developed. There are also several remaining open problems, such as the analvsis of FETI-2LM for nonsymmetric and/or nonlinear problems and the analysis of 151 substructuring preconditioners.

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# **Time Domain Maxwell Equations Solved with Schwarz Waveform Relaxation Methods**

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1 Introduction

It is very natural to solve time dependent problems with Domain Decomposition 10 Methods by using an implicit scheme for the time variable and then applying a classical iterative domain decomposition method at each time step. This is however not 12 what the Schwarz Waveform Relaxation (SWR) methods do. The SWR methods 13 are a combination of the Schwarz Domain Decomposition methods, see [10], and 14 the Waveform Relaxation algorithm, see [7]. Combined, one obtains a new method 15 which decomposes the domain into subdomains on which time dependent problems 16 are solved. Iterations are then introduced, where communication between subdomains is done at artificial interfaces along the whole time window.

This new approach has been introduced by Bjørhus [1] for hyperbolic problems 19 with Dirichlet boundary conditions and was analyzed for the heat equation by Gan- 20 der and Stuart [5]. Giladi and Keller [6] analyzed this same approach applied to the 21 advection diffusion equation with constant coefficients. For the wave equation and 22 SWR see [3] in which they treat the one-dimensional case with overlapping sub- 23 domains and for the n-dimensional case [4], again with overlap. In this paper, we 24 analyze for the first time the SWR algorithm applied to the time domain Maxwell 25 equations.

# 2 Maxwell Equations and the Schwarz Waveform Relaxation Algorithm

The global domain  $\Omega$  is decomposed into non overlapping subdomains  $\tilde{\Omega}_i$ . We de- 29 note by  $\Omega_i$  the domain  $\Omega_i$  enlarged by a band of width  $\delta$  inside of  $\Omega$ . The part of 30  $\partial\Omega_i$  in  $\tilde{\Omega}_j$  is denoted  $\Gamma_{ij}$ , i.e.  $\Gamma_{ij}:=\partial\Omega_i\cap\tilde{\Omega}_j$ . If  $\Omega_i$  possesses a part of the boundary of the global domain  $\Omega$ , we denote it by  $\Gamma_{i0} := \partial \Omega_i \cap \partial \Omega$ . The SWR algorithm 32 with characteristic transmission conditions for the time domain Maxwell equations 33 is given by

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$$\begin{cases}
-\varepsilon \partial_{t} \mathbf{E}^{i,n} + \nabla \times \mathbf{H}^{i,n} - \sigma \mathbf{E}^{i,n} = \mathbf{J}, & \Omega_{i} \times (0,T), \\
\mu \partial_{t} \mathbf{H}^{i,n} + \nabla \times \mathbf{E}^{i,n} = 0, & \Omega_{i} \times (0,T), \\
\mathscr{B}_{\mathbf{n}_{i}}(\mathbf{E}^{i,n}, \mathbf{H}^{i,n}) = 0, & \Gamma_{i0} \times (0,T), \\
(\mathbf{E}^{i,n}, \mathbf{H}^{i,n})(\mathbf{x},0) = (\mathbf{E}_{0}, \mathbf{H}_{0}), & \Omega_{i}, \\
\mathscr{B}_{\mathbf{n}_{i}}(\mathbf{E}^{i,n}, \mathbf{H}^{i,n}) = \mathscr{B}_{\mathbf{n}_{i}}(\mathbf{E}^{j,n-1}, \mathbf{H}^{j,n-1}), \Gamma_{ij} \times (0,T),
\end{cases}$$
(1)

where  $\varepsilon$  is the electric permittivity,  $\mu$  the magnetic permeability and  $\sigma$  the conductivity. The indices i and j, always different, range over the indices of all subdomains, i.e. 36  $i, j \in \{1, 2, \dots, I\}$  with  $i \neq j$  and I being the number of subdomains. In the algorithm 37  $\mathbf{n}_i$  is the unit outward normal vector to  $\Omega_i$ . The impedance

$$\mathscr{B}_{\mathbf{n}}(\mathbf{E},\mathbf{H}) := \frac{\mathbf{E}}{Z} \times \mathbf{n} + \mathbf{n} \times (\mathbf{H} \times \mathbf{n}),$$
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plays the role of the Dirichlet value for this hyperbolic system [2] and corresponds 40 to the inward characteristic variables of the Maxwell equations. The last line of (1), 41 which is called the *characteristic transmission condition*, establishes how the subdo- 42 mains communicate with each other. 43

## 3 Convergence in a Finite Number of Steps

From now on, we restrict our analysis to the specific situation where  $\Omega = \mathbb{R}^3$  which 45 is subdivided into two subdomains

$$\Omega_1 = (-\infty, L] \times \mathbb{R}^2, \quad \Omega_2 = [0, +\infty) \times \mathbb{R}^2.$$
 (2)

The artificial boundaries are therefore given by  $\Gamma_{12}=\{L\} imes\mathbb{R}^2$  and  $\Gamma_{21}=\{0\} imes\mathbb{R}^2$  47 with an overlap of width L. We also choose the coefficients  $\varepsilon$ ,  $\mu$  and  $\sigma$  to be constant. 48

Maxwell equations describe the motion of electromagnetic waves which prop- 49 agate at finite speed, namely the speed of light in the vacuum. This fact has been 50 proven for a broad class of hyperbolic systems, see for instance [8]; the Maxwell 51 equations are simply one such example. The speed of propagation is given by 52  $c := 1/\sqrt{\varepsilon \mu}$ , which is constant. 53

Remark 1. The next result also holds when the coefficients are non constant and with 54 a domain  $\Omega$  decomposed into many subdomains  $\Omega_i$  having a more complicated ge- 55 ometry and non constant overlap width.

**Proposition 1 (Convergence in a finite number of steps).** The SWR algorithm (1) 57 for two subdomains defined in (2) with overlap L converges as soon as the number 58 of iterations n satisfies 59

$$n > \frac{Tc}{L},$$
 60

where T is the length of the time interval and  $c = 1/\sqrt{\varepsilon\mu}$  is the speed of propagation. 61

*Proof.* The Maxwell equations are linear and thus allow us to restrict our attention to 62 the error equations, i.e. (1) where  $\mathbf{J} = 0$  and  $(\mathbf{E}_0, \mathbf{H}_0) = 0$ . We prove in the following 63 that for  $t < t_n := n \frac{L}{c}$ , 64

$$Supp(\mathbf{E}^{i,n+1}, \mathbf{H}^{i,n+1})(t) = \emptyset, \quad t < t_n.$$
(3)

The error of the Maxwell equations is non-zero at iteration one only because the 65 initial guesses  $(\mathbf{E}^{i,0},\mathbf{H}^{i,0})$  are non-zero on the artificial boundaries  $\Gamma_{ij}$ . The speed 66 of propagation is finite and thus the error propagates from the artificial boundaries 67 inside the domain  $\Omega_i$ . For the first iteration we have that

$$\operatorname{Supp}(\mathbf{E}^{i,1}, \mathbf{H}^{i,1})(t) \subset \{\mathbf{x} \in \Omega_i | \operatorname{dist}(x, \Gamma_{ij}) < tc, j \neq i, j \in \{1, 2\}\},$$

since after a time t, the electromagnetic wave can only have propagated on a distance 70 tc from the artificial boundaries. The overlap is of width L, hence  $(\mathbf{E}^{1,1},\mathbf{H}^{1,1})(0,y,z,t)$  71 and  $(\mathbf{E}^{2,1},\mathbf{H}^{2,1})(L,y,z,t)$  are zero unless tc>L, i.e. unless the time is greater or equal 72 to  $t_1:=\frac{L}{c}$ . 73

For the next iteration we have that the trace of  $(\mathbf{E}^{1,1}, \mathbf{H}^{1,1})$  at  $\Gamma_{21}$  and  $(\mathbf{E}^{2,1}, \mathbf{H}^{2,1})$  74 at  $\Gamma_{12}$  are zero for times  $t < t_1$ , i.e.  $B_{\mathbf{n}_i}(\mathbf{E}^{j,n-1}, \mathbf{H}^{j,n-1}) = 0$  at  $\Gamma_{ij}$  for n = 2 and  $t < t_1$ . 75 Therefore, when solving for  $(\mathbf{E}^{i,2}, \mathbf{H}^{i,2})$  we see that for  $t < t_1$ , we have zero boundary 76 conditions and zero initial condition, hence

$$(\mathbf{E}^{i,2}, \mathbf{H}^{i,2})(\mathbf{x},t) = 0$$
, for  $t < t_1$ .

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For times  $t > t_1$ , we have a similar result as for the first iteration, namely

Supp(
$$\mathbf{E}^{i,2}, \mathbf{H}^{i,2}$$
) $(t) \subset \{ \mathbf{x} \in \Omega_i | \operatorname{dist}(x, \Gamma_{ij}) < (t - t_1)c, j \neq i, j \in \{1, 2\} \}.$  80

We define  $t_2 := \frac{L}{c} + t_1 = 2t_1$ , such that Supp $(\mathbf{E}^{i,2}, \mathbf{H}^{i,2})(t) = 0$  on  $\Gamma_{ji}$  for  $t < t_2$ . And 81 so forth for the following iterations, which proves (3).

Hence, if T, the length of the time window, is finite and  $t_n := n\frac{L}{c} > T$ , the solution 83  $(\mathbf{E}^{i,n+1}, \mathbf{H}^{i,n+1})$  is zero and the algorithm has converged.

# 4 Convergence of the SWR Algorithm

Under the same setting (2) as in previous section, we prove that the SWR algorithm 86 (1) also has a contraction factor. 87

**Theorem 1.** The convergence factor of the classical Schwarz Waveform Relaxation 88 algorithm (1) in the frequency domain with domain decomposition (2) is given by 89

$$\rho(s, k_y, k_z, L, \sigma) = \left| \frac{\sqrt{|\mathbf{k}|^2 + \mu s^2 \varepsilon + \mu s \sigma} - s \sqrt{\mu \varepsilon}}{\sqrt{|\mathbf{k}|^2 + \mu s^2 \varepsilon + \mu s \sigma} + s \sqrt{\mu \varepsilon}} e^{-L\sqrt{|\mathbf{k}|^2 + \mu s^2 \varepsilon + \mu s \sigma}} \right|,$$

where s is the Laplace variable,  $\Re(s) \ge 0$ , and  $|\mathbf{k}|^2 = k_y^2 + k_z^2$  is the sum of the squares 91 of the Fourier frequencies in the y and z directions.

*Proof.* We consider the error equations for which **J** and the initial condition are zero. 93 We first apply the Laplace transform to (1) which transforms the time t into a complex frequency s with  $\Re(s) \ge 0$  and transforms the derivative with respect to t into a 95 multiplication by s. Then we apply a Fourier transform in the y and z directions and 96 obtain, 97

$$\frac{\partial}{\partial x} \begin{bmatrix} \check{E}_2 \\ \check{E}_3 \\ \check{H}_2 \\ \check{H}_3 \end{bmatrix} + \begin{bmatrix} 0 & 0 & -\frac{k_y k_z}{\varepsilon s + \sigma} & \frac{k_y^2}{\varepsilon s + \sigma} + \mu s \\ 0 & 0 & -\frac{k_z^2}{\varepsilon s + \sigma} - \mu s & \frac{k_y k_z}{\varepsilon s + \sigma} \\ \frac{k_y k_z}{\mu s} & -\frac{k_y^2}{\mu s} - (\varepsilon s + \sigma) & 0 & 0 \\ \frac{k_z^2}{\mu s} + \varepsilon s + \sigma & -\frac{k_y k_z}{\mu s} & 0 & 0 \end{bmatrix} \begin{bmatrix} \check{E}_2 \\ \check{E}_3 \\ \check{H}_2 \\ \check{H}_3 \end{bmatrix} = 0 \quad (4)$$

For components  $\check{E}_1$  and  $\check{H}_1$ , we have two algebraic equations

$$-\varepsilon s \check{E}_1 + ik_y \check{H}_3 - ik_z \check{H}_2 - \sigma \check{E}_1 = 0,$$
  

$$\mu s \check{H}_1 + ik_y \check{E}_3 - ik_z \check{E}_2 = 0.$$
<sup>99</sup>

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The solution of (4) is given by a linear combination of the eigenvectors times an exponential of the corresponding eigenvalue,

$$(\check{E}_{2}^{1,n}, \check{E}_{3}^{1,n}, \check{H}_{2}^{1,n}, \check{H}_{3}^{1,n})^{T} = (\alpha_{1}^{n}\mathbf{v}_{1} + \alpha_{2}^{n}\mathbf{v}_{2})e^{-\lambda(x-L)} + (\alpha_{3}^{n}\mathbf{v}_{3} + \alpha_{4}^{n}\mathbf{v}_{4})e^{\lambda(x-L)},$$

$$(\check{E}_{2}^{2,n}, \check{E}_{3}^{2,n}, \check{H}_{2}^{2,n}, \check{H}_{3}^{2,n})^{T} = (\beta_{1}^{n}\mathbf{v}_{1} + \beta_{2}^{n}\mathbf{v}_{2})e^{-\lambda x} + (\beta_{3}^{n}\mathbf{v}_{3} + \beta_{4}^{n}\mathbf{v}_{4})e^{\lambda x}.$$

$$(5)$$

where  $\lambda = \sqrt{|k|^2 + \mu s^2 \varepsilon + \mu s \sigma}$  and the eigenvalues are  $\lambda_{1,2} = -\lambda$  and  $\lambda_{3,4} = \lambda$ . 102 The corresponding eigenvectors are

$$\mathbf{v}_{1} = \begin{pmatrix} \frac{k_{y}k_{z}}{\lambda(\varepsilon s + \sigma)} \\ \frac{k_{z}^{2} + \mu s^{2} \varepsilon + \mu s \sigma}{\lambda(\varepsilon s + \sigma)} \\ 1 \\ 0 \end{pmatrix}, \mathbf{v}_{2} = \begin{pmatrix} -\frac{k_{y}^{2} + \mu s^{2} \varepsilon + \mu s \sigma}{\lambda(\varepsilon s + \sigma)} \\ -\frac{k_{y}k_{z}}{\lambda(\varepsilon s + \sigma)} \\ 0 \\ 1 \end{pmatrix}, \mathbf{v}_{3} = \begin{pmatrix} -\frac{k_{y}k_{z}}{\lambda(\varepsilon s + \sigma)} \\ -\frac{k_{z}^{2} + \mu s^{2} \varepsilon + \mu s \sigma}{\lambda(\varepsilon s + \sigma)} \\ -\frac{k_{z}^{2} + \mu s^{2} \varepsilon + \mu s \sigma}{\lambda(\varepsilon s + \sigma)} \\ 1 \\ 0 \end{pmatrix}, \mathbf{v}_{4} = \begin{pmatrix} \frac{k_{y}^{2} + \mu s^{2} \varepsilon + \mu s \sigma}{\lambda(\varepsilon s + \sigma)} \\ \frac{k_{y}k_{z}}{\lambda(\varepsilon s + \sigma)} \\ 0 \\ 1 \end{pmatrix}.$$

$$(6)$$

The speed of propagation is finite. The wave of the error equations propagates starting from the interfaces. Therefore, no wave is coming from the infinite boundary and then the growing exponential term of (5) is not present in the solution, i.e.  $\alpha_1 = \alpha_2 = \beta_3 = \beta_4 = 0$ . Hence,

$$(\check{E}_{2}^{1,n}, \check{E}_{3}^{1,n}, \check{H}_{2}^{1,n}, \check{H}_{3}^{1,n})^{T} = (\alpha_{3}^{n}\mathbf{v}_{3} + \alpha_{4}^{n}\mathbf{v}_{4})e^{\lambda(x-L)}, (\check{E}_{2}^{2,n}, \check{E}_{3}^{2,n}, \check{H}_{2}^{2,n}, \check{H}_{3}^{2,n})^{T} = (\beta_{1}^{n}\mathbf{v}_{1} + \beta_{2}^{n}\mathbf{v}_{2})e^{-\lambda x}.$$
 (7)

To determine the values of  $\alpha_i$  and  $\beta_i$ , we need to use the transmission conditions. 108 They are, for the first subdomain,  $\mathscr{B}_{\mathbf{n}}(\check{\mathbf{E}}^{1,n},\check{\mathbf{H}}^{1,n}) = \mathscr{B}_{\mathbf{n}}(\check{\mathbf{E}}^{2,n-1},\check{\mathbf{H}}^{2,n-1})$  with  $\mathbf{n} = 109 (1,0,0)^T$ , i.e.

 $\begin{bmatrix} \frac{1}{Z} \check{E}_{3}^{1,n} + \check{H}_{2}^{1,n} \\ -\frac{1}{Z} \check{E}_{2}^{1,n} + \check{H}_{3}^{1,n} \end{bmatrix} = \begin{bmatrix} \frac{1}{Z} \check{E}_{3}^{2,n-1} + \check{H}_{2}^{2,n-1} \\ -\frac{1}{Z} \check{E}_{2}^{2,n-1} + \check{H}_{3}^{2,n-1} \end{bmatrix}$ 111

We substitute the values of the electric and magnetic fields by their values given in 112 (7). This gives an equation relating  $\boldsymbol{\alpha}^n = (\alpha_3^n, \alpha_4^n)^T$  and  $\boldsymbol{\beta}^n = (\beta_1^n, \beta_2^n)^T$ , 113

$$A_1 \boldsymbol{\alpha}^n = A_2 e^{-\lambda L} \boldsymbol{\beta}^{n-1}, \tag{8}$$

114

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where matrices  $A_1$  and  $A_2$  are given by

$$A_{1} = \begin{bmatrix} -(k_{z}^{2} + \mu s^{2} \varepsilon + \mu s \sigma) + Z\lambda(\varepsilon s + \sigma) & k_{y}k_{z} \\ k_{y}k_{z} & -(k_{y}^{2} + \mu s^{2} \varepsilon + \mu s \sigma) + Z\lambda(\varepsilon s + \sigma) \end{bmatrix},$$

$$A_{2} = \begin{bmatrix} k_{z}^{2} + \mu s^{2} \varepsilon + \mu s \sigma + Z\lambda(\varepsilon s + \sigma) & -k_{y}k_{z} \\ -k_{y}k_{z} & k_{y}^{2} + \mu s^{2} \varepsilon + \mu s \sigma + Z\lambda(\varepsilon s + \sigma) \end{bmatrix}.$$

$$(9)$$

We do the same computations for the second subdomain for which we have the transmission conditions  $\mathscr{B}_{-n}(\hat{\mathbf{E}}^{2,n},\hat{\mathbf{H}}^{2,n}) = \mathscr{B}_{-n}(\hat{\mathbf{E}}^{1,n-1},\hat{\mathbf{H}}^{1,n-1})$ , and obtain

$$A_1 \boldsymbol{\beta}^n = A_2 e^{-\lambda L} \boldsymbol{\alpha}^{n-1}. \tag{10}$$

We isolate  $\alpha^n$  and  $\beta^n$  in (8) and (10) and iterate one more time to obtain

$$\boldsymbol{\alpha}^{n} = (A_{1}^{-1}A_{2})^{2}e^{-2\lambda L}\boldsymbol{\alpha}^{n-2}, \ \boldsymbol{\beta}^{n} = (A_{1}^{-1}A_{2})^{2}e^{-2\lambda L}\boldsymbol{\beta}^{n-2}.$$
 (11)

The parameters  $\alpha^n$  and  $\beta^n$  characterize completely the solution of (4), therefore the effective contraction factor after two iterations is given by the spectral radius of  $(A_1^{-1}A_2)^2e^{-2\lambda L}$ . This matrix has eigenvalues

$$v_1 := \left(\frac{\lambda - s\sqrt{\varepsilon\mu}}{\lambda + s\sqrt{\varepsilon\mu}}\right)^2 e^{-2\lambda L}, \quad v_2 := \left(\frac{\lambda - s\sqrt{\varepsilon\mu} - Z\sigma}{\lambda + s\sqrt{\varepsilon\mu} + Z\sigma}\right)^2 e^{-2\lambda L}.$$
 121

The largest eigenvalue in modulus is given by the first one which concludes the proof. 122

**Corollary 1.** The SWR algorithm (1) with non-zero conductivity,  $\sigma > 0$ , converges 123 in the  $L^2$  norm, i.e. if we denote by  $e^{i,n} := (E_2^{i,n}, E_3^{i,n}, H_2^{i,n}, H_3^{i,n})$ , then 124

$$||e^{i,n}(\Gamma_{ij},t)||_2 \longrightarrow 0 \quad (n \to +\infty),$$

where  $\Gamma_{ij}$  is defined in (2) and  $||\cdot||_2$  denotes the norm in  $L^2(0,T;L^2(\mathbb{R}^2))$ .

*Proof.* We use the notation  $\check{e}^{i,n} = (\check{E}_2^{i,n}, \check{E}_3^{i,n}, \check{H}_2^{i,n}, \check{H}_3^{i,n})$  for the solution in the Fourier 127 Laplace variables. From relations (11) with the notation  $R := A_1^{-1}A_2e^{-\lambda L}$  and iterating 2n times we obtain

$$\boldsymbol{\alpha}^{2n} = R^{2n} \boldsymbol{\alpha}^0, \quad \boldsymbol{\beta}^{2n} = R^{2n} \boldsymbol{\beta}^0.$$

The matrix R has eigenvalues  $v_1$  and  $v_2$  and therefore can be diagonalized using 131 the matrix of eigenvectors S, i.e.  $D = S^{-1}RS$ . The following argument, for the first subdomain  $\Omega_1$ , is similar also for the second one. 133

We define  $\gamma^n := S^{-1} \alpha^n$  for all n = 0, 1, ..., and from (7) we can reconstruct the solution of  $\check{e}^{1,2n}$  from the initial iterate, 135

$$\check{e}^{1,2n}(x,k_y,k_z,s) = e^{\lambda(x-L)}[\mathbf{v}_3 \ \mathbf{v}_4]R^{2n}\boldsymbol{\alpha}^0 = e^{\lambda(x-L)}[\mathbf{v}_3 \ \mathbf{v}_4]SS^{-1}R^{2n}S\boldsymbol{\gamma}^0 
= e^{\lambda(x-L)}[\mathbf{v}_3 \ \mathbf{v}_4]SD^{2n}\boldsymbol{\gamma}^0.$$
136

The diagonal matrix is of the form  $D = \text{diag}(v_1, v_2)$ , hence we obtain a new form for the solution evaluated at x = L, 138

$$\check{e}^{1,2n}(L,k_y,k_z,s) = v_1^{2n} \gamma_1^0 \mathbf{w}_1 + v_2^{2n} \gamma_2^0 \mathbf{w}_2, \tag{12}$$

where  $[\mathbf{w}_1 \ \mathbf{w}_2] := [\mathbf{v}_3 \ \mathbf{v}_4] S$ .

Finally Theorem 7.23 of [9] shows that the limit  $e^{i,n}(L,k_v,k_z,s)$  when  $s=\xi+$  $i\omega \rightarrow i\omega$  is the Fourier transform of  $e^{i,n}$  in the y, z and t variables. Therefore the Plancherel theorem applies and 142

$$||e^{i,n}(L,y,z,t)||_2 = ||e^{i,n}(L,k_y,k_z,i\omega)||_2,$$
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which implies by (12)

$$||e^{i,n}(L,y,z,t)||_2 = ||v_1^{2n}\gamma_1^0\mathbf{w}_1 + v_2^{2n}\gamma_2^0\mathbf{w}_2||_2$$
 145

By the dominated convergence theorem we can insert the limit, when n goes to 146 infinity, into the norm and, since  $\lim_{n\to\infty} v_i$  is almost everywhere zero for i=1,2,147it concludes the proof. 148

# **5 Numerical Experiments**

For this section we restrict the geometry of the global domain to  $\Omega = [0,1]^3$  and to 150 subdomains 151

$$\Omega_1 = [0, \frac{1}{2} + 2\Delta x] \times [0, 1] \times [0, 1], \quad \Omega_2 = [\frac{1}{2}, 1] \times [0, 1] \times [0, 1], \quad \text{152}$$

where  $\Delta x$  is the spatial mesh size in the direction x. We consider a time window 153 of length T=1. The parameters  $\varepsilon$ ,  $\mu$  and  $\sigma$  are constant and equal to one. On the physical domain we set boundary conditions for perfectly conducting medium.

The discretization is done with the Yee scheme which is explicit in time. We 156 set a global grid on the whole domain  $\Omega$  having 24 grid points in each direction x, 157 y and z. The overlap is of 2 mesh points. The number of grid points for the time 158 variable is N = 144 which guarantees that the CFL condition is satisfied. Since the

domain is bounded, only a finite number of discrete frequencies are possible. Since 160 the domain is of width one, the minimum frequency in space is given by  $k_{min} = \pi$  and the maximum by  $k_{max} = \frac{\pi}{\Delta y}$ . Equivalently for the time frequencies we have  $\omega_{max} =$  $\frac{\pi}{\Delta t}$ . Since there is no finite value imposed, we take  $\omega_{min} = \frac{\pi}{2T} = \frac{\pi}{2}$ . The discrete frequencies are therefore given by 164

$$k_y, k_z \in \{\pi, 2\pi, \dots, \frac{\pi}{\Delta y}\}, \quad \omega \in \{\frac{\pi}{2}, \pi, \dots, \frac{\pi}{\Delta t}\}.$$

From Corollary 1 we have that

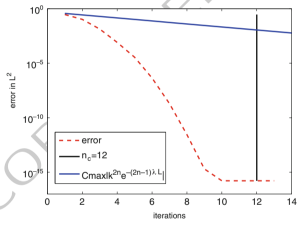
$$||e^{i,n}(L,y,z,t)||_2 \le C \max_{(k_y,k_z,\omega)} |v_1|^n,$$
 (13)

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169

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where the constant C is the maximum over all frequencies of  $||\gamma_1^0 \mathbf{w}_1 + \frac{v_2}{v_1} \gamma_2^0 \mathbf{w}_2||_2$ . 167 We also expect the solution to converge in a finite number of iterations as shown in 168 Fig. 1.



**Fig. 1.** The plain blue line is the upper bound in (13), and the dashed line is the error  $||E_2^{1,n}||$  in the  $L^2$  norm evaluated at the interface x = b with respect to the iterations. The error converges before the relation of Proposition 1 is satisfied (vertical line)

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# Comparison of a One and Two Parameter Family of Transmission Conditions for Maxwell's Equations with Damping

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1 Introduction 13

Transmission conditions between subdomains have a substantial influence on the 14 convergence of iterative domain decomposition algorithms. For Maxwell's equa- 15 tions, transmission conditions which lead to rapidly converging algorithms have been 16 developed both for the curl-curl formulation of Maxwell's equation, see [1-3], and 17 also for first order formulations, see [6, 7]. These methods have well found their 18 way into applications, see for example [9] and the references therein. It turns out 19 that good transmission conditions are approximations of transparent boundary con- 20 ditions. For each form of approximation chosen, one can try to find the best remain- 21 ing free parameters in the approximation by solving a min-max problem. Usually 22 allowing more free parameters leads to a substantially better solution of the min- 23 max problem, and thus to a much better algorithm. For a particular one parameter 24 family of transmission conditions analyzed in [4], we investigate in this paper a two 25 parameter counterpart. The analysis, which is substantially more complicated than 26 in the one parameter case, reveals that in one particular asymptotic regime there is 27 only negligible improvement possible using two parameters, compared to the one 28 parameter results. This analysis settles an important open question for this family 29 of transmission conditions, and also suggests a direction for systematically reducing 30 the number of parameters in other optimized transmission conditions.

# 2 Schwarz Methods for Maxwell's Equations

We consider in this paper a boundary value problem associated to three time- 33 harmonic Maxwell equations with an impedance condition on the boundary of the 34

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computational domain  $\Omega$ ,

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$$-i\omega\varepsilon\mathbf{E} + \operatorname{curl}\mathbf{H} - \sigma\mathbf{E} = \mathbf{J}, i\omega\mu\mathbf{H} + \operatorname{curl}\mathbf{E} = \mathbf{0}, \Omega$$

$$\mathscr{B}_{\mathbf{n}}(\mathbf{E}, \mathbf{H}) := \mathbf{n} \times \frac{\mathbf{E}}{Z} + \mathbf{n} \times (\mathbf{H} \times \mathbf{n}) = \mathbf{s}, \partial\Omega.$$
(1)

with  $\mathbf{E}, \mathbf{H}$  being the unknown electric and magnetic fields and  $\varepsilon, \mu, \sigma$  being respectively the electric permittivity, magnetic permeability and the conductivity of the propagation medium and  $\mathbf{n}$  the outward normal to  $\partial \Omega$ .

A family of Schwarz methods for (1) with a possibly non-overlapping decomposition 39 of the domain  $\Omega$  into  $\Omega_1$  and  $\Omega_2$ , with interfaces  $\Gamma_{12} := \partial \Omega_1 \cap \Omega_2$  and  $\Gamma_{21} := \partial \Omega_2 \cap \Omega_2$  40  $\Omega_1$ , is given by

$$\begin{split} -i\omega\varepsilon\mathbf{E}^{1,n} + & \operatorname{curl}\mathbf{H}^{1,n} - \sigma\mathbf{E}^{1,n} = \mathbf{J} & \text{in } \Omega_{1}, \\ & i\omega\mu\mathbf{H}^{1,n} + \operatorname{curl}\mathbf{E}^{1,n} = \mathbf{0} & \text{in } \Omega_{1}, \\ & (\mathcal{B}_{\mathbf{n}_{1}} + \mathcal{S}_{1}\mathcal{B}_{\mathbf{n}_{2}})(\mathbf{E}^{1,n}, \mathbf{H}^{1,n}) = (\mathcal{B}_{\mathbf{n}_{1}} + \mathcal{S}_{1}\mathcal{B}_{\mathbf{n}_{2}})(\mathbf{E}^{2,n-1}, \mathbf{H}^{2,n-1}) \text{ on } \Gamma_{12}, \\ & -i\omega\varepsilon\mathbf{E}^{2,n} + \operatorname{curl}\mathbf{H}^{2,n} - \sigma\mathbf{E}^{2,n} = \mathbf{J} & \text{in } \Omega_{2}, \\ & i\omega\mu\mathbf{H}^{2,n} + \operatorname{curl}\mathbf{E}^{2,n} = \mathbf{0} & \text{in } \Omega_{2}, \\ & (\mathcal{B}_{\mathbf{n}_{2}} + \mathcal{S}_{2}\mathcal{B}_{\mathbf{n}_{1}})(\mathbf{E}^{2,n}, \mathbf{H}^{2,n}) = (\mathcal{B}_{\mathbf{n}_{2}} + \mathcal{S}_{2}\mathcal{B}_{\mathbf{n}_{1}})(\mathbf{E}^{1,n-1}, \mathbf{H}^{1,n-1}) \text{ on } \Gamma_{21}, \end{split}$$

where  $\mathscr{S}_j,\ j=1,2$  are tangential operators. For the case of constant coefficients 42 and the domain  $\Omega=\mathbb{R}^2$ , with the Silver-Müller radiation condition  $\lim_{r\to\infty} r$  43  $(\mathbf{H}\times\mathbf{n}-\mathbf{E})=0$  and the two subdomains  $\Omega_1=(0,\infty)\times\mathbb{R},\ \Omega_2=(-\infty,L)\times\mathbb{R},\ L\geq 0,$  44 the following convergence result was obtained in [4] using Fourier analysis:

**Theorem 1.** For  $\sigma > 0$ , if  $\mathcal{S}_i$ , j = 1, 2 have the constant Fourier symbol

$$\sigma_{j} = \mathscr{F}(\mathscr{L}_{j}) = -\frac{s - i\tilde{\omega}}{s + i\tilde{\omega}}, \quad \tilde{\omega} = \omega\sqrt{\varepsilon\mu}, \qquad s \in \mathbb{C},$$
 (3)

then the optimized Schwarz method (2), has the convergence factor

$$\rho(k,\tilde{\omega},Z,\sigma,L,s) = \left| \left( \frac{\sqrt{k^2 - \tilde{\omega}^2 + i\tilde{\omega}\sigma Z} - s}{\sqrt{k^2 - \tilde{\omega}^2 + i\tilde{\omega}\sigma Z} + s} \right) e^{-\sqrt{k^2 - \tilde{\omega}^2 + i\tilde{\omega}\sigma Z} L} \right|. \tag{4}$$

In order to obtain the most efficient algorithm, we choose  $\sigma_j$ , j=1,2 such that  $\rho$  is 48 minimal over the range of numerical frequencies  $k \in K = [k_{\min}, k_{\max}]$ , e.g.  $k_{\min} = 0$  49 and  $k_{\max} = \frac{C}{h}$  with h the mesh size and C a constant. We look for s of the form s = p + iq, such that (p,q) is solution of the min-max problem

$$\rho^* := \min_{p,q \ge 0} \left( \max_{k \in K} \rho(k, \tilde{\omega}, Z, \sigma, L, p + iq)) \right). \tag{5}$$

In [4] we have solved this min-max problem for the case p=q without overlap, and 52 we have obtained the following result: 53

**Theorem 2.** For  $\sigma > 0$  and L = 0, the solution of the min-max problem (5) with p = q 54 is for h small given by

$$p^* = \frac{(\omega \sigma \mu)^{\frac{1}{4}} \sqrt{C}}{2^{\frac{1}{4}} \sqrt{h}} \quad and \quad \rho_1^* = 1 - \frac{2^{\frac{3}{4}} (\omega \sigma \mu)^{\frac{1}{4}} \sqrt{h}}{\sqrt{C}} + O(h). \tag{6}$$

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For the overlapping case, we obtained in [8]:

**Theorem 3.** For  $\sigma > 0$  and L = h, a local minimum of the min-max problem (5) with 57 p = q is for h small given by

$$p^* = \frac{(2\omega\sigma\mu)^{\frac{1}{3}}}{2h^{\frac{1}{3}}} \quad and \quad \rho_{1L}^* = 1 - 2^{\frac{7}{6}} (\omega\sigma\mu)^{\frac{1}{6}} h^{\frac{1}{3}} + O(h^{\frac{2}{3}}). \tag{7}$$

# 3 Analysis of the Two Parameter Family of Transmission Conditions 59

As before, we set  $k_{\min}=0$ ,  $k_{\max}=\frac{C}{h}$  and denote by  $(p^*,q^*)$  a local minimum of (5). 61 We first consider the non-overlapping case.

**Theorem 4.** For  $\sigma > 0$  and L = 0, a local minimum  $(p^*, q^*)$  of (5) is for h small 63 given by

$$p^* = \frac{3^{\frac{3}{8}}(\omega\sigma\mu)^{\frac{1}{4}}\sqrt{C}}{2^{\frac{3}{4}}\sqrt{h}}, q^* = \frac{3^{\frac{7}{8}}(2\omega\sigma\mu)^{\frac{1}{4}}\sqrt{C}}{6\sqrt{h}}, \rho_2^* = 1 - \frac{3^{\frac{3}{8}}(2\omega\sigma\mu)^{\frac{1}{4}}\sqrt{h}}{\sqrt{C}} + O(h).$$
(8)

*Proof.* By solving the min-max problem (5) numerically for different parameter values and different mesh sizes h, we observe that the solution of (5) equioscillates once, 66 i.e.  $(p^*, q^*)$  is solution of

$$\rho(\bar{k}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*) = \rho(k_{\text{max}}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*), \tag{9}$$

where  $\bar{k}$  is an interior local maximum of  $\rho$ . We also observe the asymptotic behavior 68

$$\bar{k} \sim \bar{C}, \quad p^* \sim C_p h^{-\frac{1}{2}}, \quad q^* \sim C_q h^{-\frac{1}{2}}.$$
 69

In order to determine the constants  $\bar{C}$ ,  $C_p$  and  $C_q$ , it is necessary to have three equations. The first is (9), the second describes the interior local maximum of  $\rho$  in k,

$$\frac{\partial \rho}{\partial k}(\bar{k}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*)) = 0,$$
 72

and the third is the necessary condition for a local minimum of the min-max problem, 73

$$\begin{split} \frac{d\rho}{dq}(k_{\max},\tilde{\boldsymbol{\omega}},\boldsymbol{\sigma},Z,0,p^*+iq^*) &= \\ \frac{\partial\rho}{\partial q}(k_{\max},\tilde{\boldsymbol{\omega}},\boldsymbol{\sigma},Z,0,p^*+iq^*) + \frac{\partial\rho}{\partial p}(k_{\max},\tilde{\boldsymbol{\omega}},\boldsymbol{\sigma},Z,0,p^*+iq^*) \frac{\partial p}{\partial q} &= 0. \end{split}$$

Since  $\frac{d\rho}{dq}(k_{\max},\tilde{\omega},\sigma,Z,0,p^*+iq^*)=\frac{d\rho}{dq}(\bar{k},\tilde{\omega},\sigma,Z,0,p^*+iq^*)$  a similar expansion 75 together with the previous one, gives

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$$\frac{\partial p}{\partial q} = -\frac{\frac{\partial \rho}{\partial q}(k_{\max}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*) - \frac{\partial \rho}{\partial q}(\bar{k}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*)}{\frac{\partial \rho}{\partial p}(k_{\max}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*) - \frac{\partial \rho}{\partial p}(\bar{k}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*)},$$
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and thus asymptotically, the three equations lead to the system

$$\begin{split} (\sqrt{A_1} + \bar{C}^2 - \tilde{\omega}^2)(AC_p + BC_q) - 2\sqrt{A_1}BC_q &= 0, \\ 2C_p(C_p^2 + C_q^2) - C(BC_p + AC_q) &= 0, \\ A(C_q^2 - C_p^2) + 2C_pC_qB &= 0, \end{split}$$

where  $A=\sqrt{2\sqrt{A_1}-A_2}$ ,  $B=\sqrt{2\sqrt{A_1}+A_2}$ ,  $A_1=\bar{C}^4-2(\bar{C}\tilde{\omega})^2+\tilde{\omega}^4+(\tilde{\omega}\sigma Z)^2$  and 79  $A_2=2(\bar{C}^2-\tilde{\omega}^2)$ . The solution of this system is

$$\bar{C} = \frac{\sqrt{\tilde{\omega}\left(-Z\sigma\sqrt{3}+3\tilde{\omega}\right)}}{\sqrt{3}}, \quad C_p = \frac{3^{\frac{3}{8}}(\tilde{\omega}\sigma Z)^{\frac{1}{4}}\sqrt{C}}{2^{\frac{3}{4}}}, \quad C_q = \frac{3^{\frac{7}{8}}(2\tilde{\omega}\sigma Z)^{\frac{1}{4}}\sqrt{C}}{6}, \qquad \text{81}$$

from which (8) follows. It remains to show that  $(p^*, q^*)$  is a local minimum, i.e. for 82 any variation  $(\delta p, \delta q)$  and  $k \in \{\bar{k}, k_{\text{max}}\}$ , we must have

$$\rho(k, \tilde{\omega}, \sigma, Z, 0, p^* + \delta p + i(q^* + \delta q)) \ge \rho(k, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*).$$

By the Taylor formula, it suffices to prove that there is no variation  $(\delta p, \delta q)$  such 85 that for  $k \in \{\bar{k}, k_{\max}\}$ 

$$\delta p \frac{\partial \rho}{\partial p}(k, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*) + \delta q \frac{\partial \rho}{\partial q}(k, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*) < 0. \tag{10}$$

We prove this by contradiction, and it is necessary to obtain the next higher order 87 terms in the expansions of  $p^*$ ,  $q^*$  and  $\bar{k}$ . After a lengthy computation, we find that 88 asymptotically

$$\bar{k} \sim \bar{C} + \tilde{C}h, \quad p^* \sim C_p h^{-\frac{1}{2}} + \tilde{C}_p h^{\frac{3}{2}}, \quad q^* \sim C_q h^{-\frac{1}{2}} + \tilde{C}_q h^{\frac{1}{2}}.$$

The computation of these new three constants allows us to obtain the partial derivatives of  $\rho$  91

$$\frac{\partial \rho}{\partial p}(\bar{k}) \sim \frac{2}{C}h, \quad \frac{\partial \rho}{\partial q}(\bar{k}) \sim -\frac{3^{\frac{1}{4}}(2\omega\sigma\mu)^{\frac{1}{2}}}{C^{2}}h^{2},$$

$$\frac{\partial \rho}{\partial p}(k_{\text{max}}) \sim -\frac{2}{C}h, \quad \frac{\partial \rho}{\partial q}(k_{\text{max}}) \sim \frac{3^{\frac{1}{4}}(2\omega\sigma\mu)^{\frac{1}{2}}}{C^{2}}h^{2}.$$
93

Introducing these results into (10), we get  $\delta p \frac{2}{C}h - \delta q \frac{3^{\frac{1}{4}}(2\omega\sigma\mu)^{\frac{1}{2}}}{C^2}h^2 < 0$  and  $-\delta p \frac{2}{C}h + 94\delta q \frac{3^{\frac{1}{4}}(2\omega\sigma\mu)^{\frac{1}{2}}}{C^2}h^2 < 0$ , clearly a contradiction, and thus  $(p^*,q^*)$  is a local minimum.

We see that for h small, both the one parameter and two parameter transmission 96 conditions can be written as  $\rho_1^*=1-\alpha_1\sqrt{h}+O(h)$  and  $\rho_2^*=1-\alpha_2\sqrt{h}+O(h)$ . The 97 ratio  $\frac{\alpha_2}{\alpha_1}$  is equal to  $3^{\frac{3}{8}}/\sqrt{2}\approx 1.067$ , which shows that the convergence factors are 98 almost equal. Hence the hypothesis p=q, used in [4] to simplify the analysis, is 99 justified.

We treat now the overlapping case of (5), with an overlap of one mesh size.

**Theorem 5.** For  $\sigma > 0$  and L = h, a local minimum  $(p^*, q^*)$  of (5) is for h small 102 given by

$$p^* = \frac{3^{\frac{1}{2}} (\omega \sigma \mu)^{\frac{1}{3}}}{2^{\frac{4}{3}} h^{\frac{1}{3}}}, \quad q^* = \frac{(\omega \sigma \mu)^{\frac{1}{3}}}{2^{\frac{4}{3}} h^{\frac{1}{3}}}, \quad \rho_{2L}^* = 1 - 2^{\frac{5}{6}} 3^{\frac{3}{8}} (\omega \sigma \mu)^{\frac{1}{6}} h^{\frac{1}{3}} + O(h^{\frac{2}{3}}). \quad (11)$$

*Proof.* As in the proof of Theorem 4, we first observe numerically that the solution of (5) equioscillates once, i.e.  $(p^*, q^*)$  is solution of

$$\rho(\bar{k}_1, \tilde{\omega}, \sigma, Z, h, p^* + iq^*) = \rho(\bar{k}_2, \tilde{\omega}, \sigma, Z, h, p^* + iq^*),$$

where  $\bar{k}_1$  and  $\bar{k}_2$  are interior local maxima of  $\rho$ , and we obtain asymptotically for h 107 small

$$\bar{k}_1 \sim C_{b_1}, \bar{k}_2 \sim C_{b_2} h^{-\frac{2}{3}}, p^* \sim C_p h^{-\frac{1}{3}} \text{ and } q^* \sim C_q h^{-\frac{1}{3}}.$$

It remains to find  $C_{b_1}$ ,  $C_{b_2}$ ,  $C_p$  and  $C_q$ . Proceeding as before, we obtain four equations from the necessary conditions of a minimum, with solution

$$C_p = rac{3^{rac{1}{2}}(2\omega\sigma\mu)^{rac{1}{2}}}{2}, C_q = rac{C_p}{\sqrt{3}}, C_{b_1} = rac{\sqrt{ ilde{\omega}\left(-Z\sigma\sqrt{3}+3 ilde{\omega}
ight)}}{\sqrt{3}}, C_{b_2} = \sqrt{2C_p},$$
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which leads to (11). To prove that  $(p^*, q^*)$  is a local minimum, proceeding as before, we obtain after a lengthy computation the higher order expansion 114

$$ar{k}_1 \sim C_{b_1} + ilde{C}_{b_1} h^{rac{2}{3}}, ar{k}_2 \sim C_{b_2} h^{-rac{2}{3}} + ilde{C}_{b_2}, p^* \sim C_p h^{-rac{1}{3}} + ilde{C}_p h^{rac{1}{3}}, q^* \sim C_q h^{-rac{1}{3}} + ilde{C}_q h^{rac{1}{3}}.$$
 115

The computation of these four new constants allows us then to obtain the partial derivatives of  $\rho$ ,

$$\frac{\frac{\partial \rho}{\partial p}(\bar{k}_1)}{\frac{\partial \rho}{\partial p}(\bar{k}_1)} \sim \frac{8 \cdot 2^{\frac{1}{6}} h^{\frac{2}{3}}}{3^{\frac{1}{4}} (\omega \sigma \mu)^{\frac{1}{6}}}, \frac{\partial \rho}{\partial q}(\bar{k}_1) \sim -\frac{2 \cdot 2^{\frac{5}{6}} (\omega \sigma \mu)^{\frac{1}{6}} h^{\frac{4}{3}}}{3^{\frac{1}{4}}}, 
\frac{\partial \rho}{\partial p}(\bar{k}_2) \sim -\frac{4 \cdot 2^{\frac{1}{6}} h^{\frac{2}{3}}}{3^{\frac{1}{4}} (\omega \sigma \mu)^{\frac{1}{6}}}, \frac{\partial \rho}{\partial q}(\bar{k}_2) \sim \frac{2^{\frac{5}{6}} (\omega \sigma \mu)^{\frac{1}{6}} h^{\frac{4}{3}}}{3^{\frac{1}{4}}}.$$
(12)

In order to reach a contradiction, we assume again there exists, by the Taylor theorem, a variation  $(\delta p, \delta q)$  such that  $\delta p \frac{\partial \rho}{\partial p}(k, \tilde{\omega}, \sigma, Z, h, 119)$   $p^* + iq^*) + \delta q \frac{\partial \rho}{\partial q}(k, \tilde{\omega}, \sigma, Z, h, p^* + iq^*) < 0$ , for  $k \in \{\bar{k}_1, k_2\}$ . Using (12), we get 120  $8 \frac{2^{\frac{1}{6}h^{\frac{2}{3}}}}{3^{\frac{1}{4}}(\omega\sigma\mu)^{\frac{1}{6}}} \delta p - 2 \frac{2^{\frac{5}{6}}(\omega\sigma\mu)^{\frac{1}{6}h^{\frac{4}{3}}}}{3^{\frac{1}{4}}} \delta q < 0$  and  $-4 \frac{2^{\frac{1}{6}h^{\frac{2}{3}}}}{3^{\frac{1}{4}}(\omega\sigma\mu)^{\frac{1}{6}}} \delta p + \frac{2^{\frac{5}{6}}(\omega\sigma\mu)^{\frac{1}{6}h^{\frac{4}{3}}}}{3^{\frac{1}{4}}} \delta q < 0$ , 121 clearly a contradiction, and thus  $(p^*, q^*)$  is a local minimum.

We also observe in this case that for h small, both convergence factors can be written 123 as  $\rho_{1L}^*=1-\alpha_{1L}\ h^{\frac{1}{3}}+O(h^{\frac{2}{3}})$  and  $\rho_{2L}^*=1-\alpha_{2L}h^{\frac{1}{3}}+O(h^{\frac{2}{3}})$ , and the ratio  $\frac{\alpha_{2L}}{\alpha_{1L}}$  is 124 equal to  $3^{\frac{1}{4}}/2^{\frac{1}{3}}\approx 1.044$ , hence both convergence factors are almost equal. We show 125 an example of these convergence factors in Fig. 1.

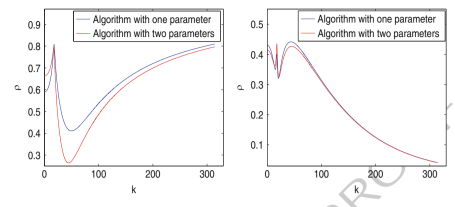


Fig. 1. Convergence factor comparison of algorithms with one and two parameters for  $\omega = 2\pi$ ,  $\sigma = 2$  and  $\mu = \varepsilon = 1$ , for the non-overlapping case, L = 0, on the *left*, and the overlapping case,  $L = h = \frac{1}{100}$ , on the *right* 

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### 4 Numerical Results

We present now a numerical test in order to compare the performance of both the 128 one and two parameter algorithms. We compute the propagation of a plane wave in 129 a heterogeneous medium. The domain is  $\Omega = (-1, 1)^2$ . The relative permittivity and 130 the conductivity of the background media is  $\varepsilon_1 = 1.0$  and  $\sigma_1 = 1.8$ , while that of the square material inclusion is  $\varepsilon_2 = 8.0$  and  $\sigma_2 = 7.5$ , see the left picture of Fig. 2. 132 The magnetic permeability  $\mu$  is constant in  $\Omega$  and we impose on the boundary an 133 incident field  $(H_r^{inc}, H_v^{inc}, E_z^{inc})$ . The domain  $\Omega$  is decomposed into two subdomains  $\Omega_1 = (-1,L) \times (-1,1)$  and  $\Omega_2 = (0,1) \times (-1,1)$ ; L is the overlapping size and is equal to the mesh size. We use, in each subdomain, a discontinuous Galerkin method 136 (DG) with a uniform polynomial approximation of order one, two and three, denoted 137 by DG-P1, DG-P2 and DG-P3, see [5]. The results are shown in Fig. 3, and are in 138 good agreement with our analytical results.

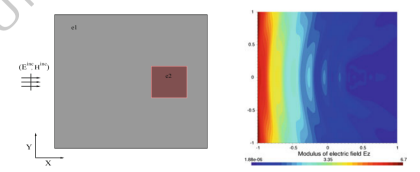


Fig. 2. Configuration of our test problem on the *left*, and the numerical solution on the *right* 

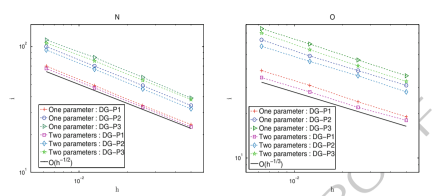


Fig. 3. Number of iterations against the mesh size h, to attain a relative residual reduction of  $10^{-8}$ 

5 Conclusion 140

We compared in this paper a one and a two parameter family of transmission 141 conditions for optimized Schwarz methods applied to Maxwell's equations. Our 142 asymptotic analysis reveals that the addition of a second parameter does not lead 143 to a significant improvement of the algorithm, and it is therefore justified to consider 144 only the simpler case of a one parameter family of transmission conditions. These 145 results are also confirmed by our numerical experiments.

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### **AUTHOR QUERIES**

- AQ1. Please provide opening parenthesis for "... $\sigma$ , L, p+iq)" in Eq. 5 AQ2. Please provide opening parenthesis for "... $(\bar{k}, \tilde{\omega}, \sigma, Z, 0, p^* + iq^*)$ ".

# Hybrid Domain Decomposition Solvers for the Helmholtz and the Time Harmonic Maxwell's **Equation**

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Summary. We present hybrid finite element methods for the Helmholtz equation and the time 10 harmonic Maxwell equations, which allow us to reduce the unknowns to degrees of freedom 11 supported only on the element facets and to use efficient iterative solvers for the resulting 12 system of equations. For solving this system, additive and multiplicative Schwarz preconditioners with local smoothers and a domain decomposition preconditioner with an exact subdomain solver are presented. Good convergence properties of these preconditioners are shown 15 by numerical experiments.

1 Introduction 17

When solving the Helmholtz equation with a standard finite element method (FEM), 18 due to the oscillatory behaviour of the solution and the pollution error [8] a large 19 number of degrees of freedom (DoFs) is needed to resolve the wave, especially for 20 high wave numbers. To overcome this difficulty, many methods have been developed 21 during the last years. Apart from hp FEM [8], Galerkin Least Square Methods [7] or 22 Discontinuous Galerkin Methods [6], some methods make use of problem adapted 23 functions like plane waves. The most popular among them are the Partition of Unity 24 Method [9], the Discontinuous Enrichment Approach [5] or the UWVF [2, 10]. All 25 these techniques end up with large, complex valued, indefinite, possible symmetric 26 linear systems. Although some advances have been made [3, 4], efficient precondi- 27 tioners for wave type problems are still a big challenge.

In the present work the hybrid FEM from [11] is used for the Helmholtz equation 29 and extended to the Maxwell case. This method allows us to use efficient iterative 30 methods for solving the resulting linear system of equations. Following hybridiza- 31 tion techniques from [1], the tangential continuity of the flux field is broken across 32 element interfaces. In order to impose continuity again, Lagrange multipliers sup- 33 ported only on the facets, which can be interpreted as the tangential component of the 34 unknown field, are introduced. Adding a second set of Lagrange multipliers, 35

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representing the tangential component of the flux field, allows us, due to local Robin 36 boundary conditions, to eliminate the volume DoFs. Because, after hybridization, 37 there is no coupling between volume basis functions of different elements, elimina- 38 tion of the volume DoFs can be done cheaply element by element, and the system 39 of equation is reduced onto the smaller set of Lagrange multipliers. For the reduced 40 system we present additive (AS) and multiplicative Schwarz (MS) block precondi- 41 tioners with blocks related to DoFs of one facet and element, respectively. Addi- 42 tionally a domain decomposition (DD) preconditioner, which directly solves for the 43 DoFs belonging to one subdomain, is investigated. This preconditioner is especially 44 advantageous for domains contains cavity like structures. Numerical tests show, that 45 a preconditioned CG iteration has good convergence properties combined with these 46 preconditioners.

### 2 Hybridization of the Wave Equations

In the sequence, we will stick to the following settings. As computational domain we 49 consider a Lipschitz polyhedron  $\Omega \subset \mathbb{R}^d$  with d=2,3 and the boundary  $\Gamma=\partial\Omega$ . 50 In the scalar case, we search for a function  $u:\Omega\to\mathbb{C}$  and a vector valued field 51  $\mathbf{v}: \Omega \to \mathbb{C}^d$ , which fulfills the Helmholtz equation in mixed form

$$\operatorname{grad} u = i\omega \mathbf{v}$$
 and  $\operatorname{div} \mathbf{v} = i\omega u$  in  $\Omega$ 

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with absorbing boundary conditions  $\mathbf{v} \cdot \mathbf{n} + u = g$  on  $\Gamma$ , where  $\omega$  is the angular frequency and **n** the outer normal vector. From [9] we know, that the solution u exists 55 and is unique.

In the vectorial case, i.e. the harmonic Maxwell's equations, we search for a 57 vector valued function  $\mathbf{E}: \Omega \to \mathbb{C}^3$  and a flux field  $\mathbf{H}: \Omega \to \mathbb{C}^3$ , which solves

$$\operatorname{curl} \mathbf{H} + i\omega \mathbf{E} = 0$$
 and  $\operatorname{curl} \mathbf{E} - i\omega \mathbf{H} = 0$  in  $\Omega$  59

under the boundary condition  $-n \times H + E_{\parallel} = g$  on  $\Gamma$  , where  $E_{\parallel}$  represents the tangential component of **E**, i.e.  $\mathbf{n} \times \mathbf{E} \times \mathbf{n}$ .

When deriving the hybrid formulation, we use a regular finite element mesh  $\mathcal{T}$  with 62 elements T, and the set of facets is called  $\mathscr{F}$ . The vector  $\mathbf{n}_T$  is the outer normal 63 vector of the element T, and  $\mathbf{n}_F$  represents the normal vector onto a facet F. Furthermore, we denote a volume integral as  $(u, v)_T := \int_T uv \, d\mathbf{x}$ , and a surface integral as 65  $\langle u, v \rangle_{\partial T} := \int_{\partial T} uv \, ds.$ 

### 2.1 The Mixed Hybrid Formulation for the Helmholtz Equation

The mixed hybrid formulation for the Helmholtz equation was already introduced in 68 [11]. For completeness, we repeat the problem formulation: Find  $(u, \mathbf{v}, u^F, v^F) \in L^2(\Omega) \times H(\text{div}, T) \times L^2(\mathscr{F}) \times L^2(\mathscr{F}) =: X \times \tilde{Y} \times X^F \times Y^F$ , such 70

Find 
$$(u, \mathbf{v}, u^r, v^r) \in L^-(\Omega^2) \times H(\operatorname{div}, I) \times L^-(\mathscr{F}) \times L^-(\mathscr{F}) =: X \times I \times X^r \times I^r$$
, such that for all  $(\sigma, \mathbf{w}, \sigma^F, w^F) \in X \times \widetilde{Y} \times X^F \times Y^F$ 

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$$\begin{split} &\sum_{T \in \mathscr{T}} \left( \left( i \boldsymbol{\omega} \boldsymbol{u}, \boldsymbol{\sigma} \right)_T - \left( i \boldsymbol{\omega} \mathbf{v}, \mathbf{w} \right)_T - \left( \operatorname{div} \mathbf{v}, \boldsymbol{\sigma} \right)_T - \left( \boldsymbol{u}, \operatorname{div} \mathbf{w} \right)_T + \left\langle \boldsymbol{u}^F, \mathbf{n}_T \cdot \mathbf{w} \right\rangle_{\partial T} \right. \\ &+ \left\langle \mathbf{n}_T \cdot \mathbf{v}, \boldsymbol{\sigma}^F \right\rangle_{\partial T} + \left\langle \mathbf{n}_F \cdot \mathbf{v} - \boldsymbol{v}^F, \mathbf{n}_F \cdot \mathbf{w} - \boldsymbol{w}^F \right\rangle_{\partial T} \right) + \left\langle \boldsymbol{u}^F, \boldsymbol{\sigma}^F \right\rangle_{\Gamma} = \left\langle g, \boldsymbol{\sigma}^F \right\rangle_{\Gamma}. \end{split}$$

### 2.2 The Mixed Hybrid Formulation for the Maxwell Problem

We will now concentrate on the derivation of the mixed hybrid formulation for the 73 vectorial wave equation. We start from the mixed system of equations from above, 74 multiply the first equation with a test function  $\mathbf{e} \in U := (L^2(\Omega))^3$  and the second one 75 with a function  $\mathbf{h} \in V := H(\text{curl}, \Omega)$  and integrate over the domain  $\Omega$ . Performing 76 integration by parts elementwise leads to

$$\sum_{T \in \mathcal{T}} \left( \left( \operatorname{curl} \mathbf{H}, \mathbf{e} \right)_T + \left( i \omega \mathbf{E}, \mathbf{e} \right)_T \right) = 0 \qquad \forall \mathbf{e} \in U$$

$$\sum_{T \in \mathcal{T}} \left( \left( \mathbf{E}, \operatorname{curl} \mathbf{h} \right)_T - \left( i \omega \mathbf{H}, \mathbf{h} \right)_T - \left\langle \mathbf{E}, \mathbf{n}_T \times \mathbf{h} \right\rangle_{\partial T} \right) = 0 \qquad \forall \mathbf{h} \in V$$

Note that for a tangential continuous field  $\mathbf{E}$ , i.e.  $\mathbf{n} \times \mathbf{E} \times \mathbf{n}$  is continuous on element 78 interfaces, the boundary integrals for inner facets cancel due to the tangential continuity of  $\mathbf{h}$ , and inserting the absorbing boundary condition into the boundary facet 80 integrals leads to the standard mixed finite element formulation for our problem. 81

Next, the tangential continuity of the flux field  $\mathbf{H}$  is broken across element interfaces, thus we search for  $\mathbf{H} \in \widetilde{V} := \left\{ \mathbf{v} \in (L^2(\Omega))^3 : \mathbf{v}|_T \in H(\operatorname{curl},T) \ \forall T \in \mathscr{T} \right\}$ . In 83 order to reinforce continuity, Lagrange multipliers  $\mathbf{E}^F$ , which are only supported on 84 the element facets, i.e. they are from the space  $U^F := (L^2(\mathscr{F}))^3$ , are introduced. The 85 continuity of the tangential fluxes is reached via an additional equation, which forces 86 the jump of  $[\mathbf{n} \times \mathbf{H}] := \mathbf{n}_{T_1} \times \mathbf{H}|_{T_1} + \mathbf{n}_{T_2} \times \mathbf{H}|_{T_2}$  for inner facets  $F \in \mathscr{F}_I$  with adjacent 87 elements  $T_1$  and  $T_2$  to zero, thus

$$\sum_{F \in \mathscr{F}_I} \left\langle [\mathbf{n} \times \mathbf{H}], \mathbf{e} \right\rangle_F = \sum_{T \in \mathscr{T}} \left( \left\langle \mathbf{n}_T \times \mathbf{H}, \mathbf{e} \right\rangle_{\partial T} - \left\langle \mathbf{n}_T \times \mathbf{H}, \mathbf{e} \right\rangle_{\partial T \cap \Gamma} \right) = 0, \quad \forall \mathbf{e} \in U^F. \quad \text{89}$$

The resulting system of equations for  $(\mathbf{E}, \mathbf{H}, \mathbf{E}^F) \in U \times \tilde{V} \times U^F$  reads as

$$\begin{split} \sum_{T \in \mathscr{T}} \left( \left( \operatorname{curl} \mathbf{H}, \mathbf{e} \right)_T + \left( i \omega \mathbf{E}, \mathbf{e} \right)_T \right) &= 0 & \forall \mathbf{e} \in U \\ \sum_{T \in \mathscr{T}} \left( \left( \mathbf{E}, \operatorname{curl} \mathbf{h} \right)_T - \left( i \omega \mathbf{H}, \mathbf{h} \right)_T - \left\langle \mathbf{E}^F, \mathbf{n}_T \times \mathbf{h} \right\rangle_{\partial T} \right) &= 0 & \forall \mathbf{h} \in \tilde{V} \\ - \sum_{T \in \mathscr{T}} \left\langle \mathbf{n}_T \times \mathbf{H}, \mathbf{e}^F \right\rangle_{\partial T} + \left\langle \mathbf{E}^F, \mathbf{e}^F \right\rangle_{\Gamma} &= \left\langle \mathbf{g}, \mathbf{e}^F \right\rangle_{\Gamma} & \forall \mathbf{e}^F \in U^F. \end{split}$$

In this system of equations, the Lagrange parameter  $\mathbf{E}^F$  plays the role of the tangential component of  $\mathbf{E}$ , evaluated on the facets. Because there is no coupling between 92 volume DoFs belonging to different elements, it is possible to eliminate the volume 93 unknowns  $\mathbf{E}$  and  $\mathbf{H}$ , cheaply by static condensation (compare [1]). The resulting system of equations needs now to be solved only for the Lagrange multipliers. 95

In order to eliminate the inner DoFs, one has to solve the first two equations 96 of the system from above for some function  $\mathbf{E}^F$  element by element. But this is 97 equivalent to solving a Dirichlet problem, and uniqueness of the solution can not 98 be guaranteed. This drawback can be compensated by adding a new facet unknown 99  $\mathbf{H}^F \in V^F := (L^2(\mathscr{F}))^3$  representing  $\mathbf{n}_F \times \mathbf{H}$  on the facets via a consistent stabilization 100 term  $\sum_{T} \langle \mathbf{n}_{F} \times \mathbf{H} - \mathbf{H}^{F}, \mathbf{n}_{F} \times \mathbf{h} - \mathbf{h}^{F} \rangle_{\partial T}$ . We obtain

$$\sum_{T \in \mathscr{T}} \left( \left( \operatorname{curl} \mathbf{H}, \mathbf{e} \right)_T + \left( i \omega \mathbf{E}, \mathbf{e} \right)_T \right) = 0 \qquad \forall \mathbf{e} \in U$$

$$\sum_{T \in \mathscr{T}} \left( \left( \mathbf{E}, \operatorname{curl} \mathbf{h} \right)_T - \left( i \omega \mathbf{H}, \mathbf{h} \right)_T - \left\langle \mathbf{E}^F, \mathbf{n}_T \times \mathbf{h} \right\rangle_{\partial T}$$
(1)

$$-\langle \mathbf{n}_T \times \mathbf{H}, \mathbf{n}_T \times \mathbf{h} \rangle_{\partial T} + \langle \mathbf{H}^F, \mathbf{n}_F \times \mathbf{h} \rangle_{\partial T} = 0 \qquad \forall \mathbf{h} \in \tilde{V}$$
 (2)

$$\sum_{T \in \mathcal{T}} \left( \left\langle \mathbf{n}_F \times \mathbf{H}, \mathbf{h}^F \right\rangle_{\partial T} - \left\langle \mathbf{H}^F, \mathbf{h}^F \right\rangle_{\partial T} \right) = 0 \qquad \forall \mathbf{h}^F \in V^F \quad (3)$$

$$-\langle \mathbf{n}_{T} \times \mathbf{H}, \mathbf{n}_{T} \times \mathbf{h} \rangle_{\partial T} + \langle \mathbf{H}^{F}, \mathbf{n}_{F} \times \mathbf{h} \rangle_{\partial T} ) = 0 \qquad \forall \mathbf{h} \in \tilde{V}$$
(2)  

$$\sum_{T \in \mathscr{T}} \left( \langle \mathbf{n}_{F} \times \mathbf{H}, \mathbf{h}^{F} \rangle_{\partial T} - \langle \mathbf{H}^{F}, \mathbf{h}^{F} \rangle_{\partial T} \right) = 0 \qquad \forall \mathbf{h}^{F} \in V^{F}$$
(3)  

$$-\sum_{T \in \mathscr{T}} \langle \mathbf{n}_{T} \times \mathbf{H}, \mathbf{e}^{F} \rangle_{\partial T} + \langle \mathbf{E}^{F}, \mathbf{e}^{F} \rangle_{\Gamma} = \langle \mathbf{g}, \mathbf{e}^{F} \rangle_{\Gamma} \quad \forall \mathbf{e}^{F} \in U^{F}.$$
(4)

Now, by static condensation the time harmonic Maxwell's equation with absorbing 102 boundary conditions has to be solved on the element level, where uniqueness is guar- 103 anteed, and the resulting system contains only the facet unknowns  $\mathbf{E}^F$  and  $\mathbf{H}^F$ . Thus 104 we search for a function  $\mathbf{w} \in W := U^F \times V^F$  such that

$$s(\mathbf{w}, \mathbf{v}) = f(\mathbf{v}) \qquad \forall \mathbf{v} \in W,$$

113

where the Schur complement bilinearform s and the linearform f are obtained 107 from (1) to (4) by eliminating the unknowns E and H. Elimination of the inner DoFs 108 can be also seen as calculating for a given incoming impedance trace  $\mathbf{E}^F - \mathbf{H}^F$  the 109 resulting outgoing impedance trace  $\mathbf{E}^{F} + \mathbf{H}^{F}$  on the element level. By exchanging the Dirichlet and Neumann traces  $\mathbf{E}^F$ ,  $\mathbf{H}^F$  by incoming and outgoing impedance traces, 111 one obtains an equivalent formulation which fits well into the context of the UWVF 112

3 Iterative Solvers 114

In this section, we focus on solving the system of equations. As already mentioned, 115 the volume DoFs can be eliminated cheaply element by element, and the resulting 116 system of equation just has to be solved for the much smaller number of facet DoFs. 117 Because volume DoFs of one element couple apart from themselves only to facet 118 DoFs of the surrounding facets, the Schur complement matrix S obtained by static 119 condensation is sparse, and it just has nonzero entries between facet DoFs belong- 120 ing to facets of the same element. Due to the hybrid formulation, efficient iterative 121 solvers can be used for the reduced system of equations.

Because the Schur complement matrix is complex symmetric, a preconditioned 123 CG-iteration together with an AS or MS block preconditioner,  $M_{AS}$  and  $M_{MS}$  is used, 124 although convergence for complex symmetric matrices is not guaranteed. The iteration matrices of these two preconditioners are given as

$$I - M_{AS}^{-1}S = I - \sum_{i=1}^{n} P_i,$$
127

126

155

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$$I - M_{MS}^{-1}S = \left(\prod_{i=n}^{1} (I - P_i)\right) \left(\prod_{i=1}^{n} (I - P_i)\right),$$
 128

where  $P_i$  is the matrix representation of the variational projector  $\mathscr{P}_i: W \to W_i \subset W$ with respect to the bilinearform s. In the scalar case  $W = X^F \times Y^F$ . We will use two different choices of subspaces  $W_i$ , functions supported on the facet  $F_i$  or on facets, which are boundary facets of the element  $T_i$ . Note that the first strategy leads to 133 nonoverlapping blocks, while the blocks of the second choice overlap.

Apart from an AS or MS Preconditioner, a DD preconditioner compareable to 135 [12] was used, which is based on a partitioning of the domain  $\Omega$  into N subdomains 136  $\Omega_i$ . The iteration matrix of this preconditioner can be described by

$$I - M_{DD}^{-1}S = \left(\prod_{i=n}^{1} (I - P_{I,i})\right) \left(I - \sum_{i=1}^{N} P_{\Omega_{i}}\right) \left(\prod_{i=1}^{n} (I - P_{I,i})\right), \tag{138}$$

where  $P_{\Omega_i}$  and  $P_{I,i}$  are matrices corresponding to variational projection operators 139 which project to the spaces  $W_{\Omega_i}$  and  $W_{I,i}$ . The space  $W_{\Omega_i}$  contains functions which are supported only on facets in the interior of the subdomain  $\Omega_i$ , while the space  $W_{I,i}$  141 is choosen such that it contains functions which are only supported on facets of an 142 element  $T_i$  such that  $\partial T_i \cap \partial \Omega_i \neq \emptyset$ . Again a nonoverlapping option is to collect the 143 functions supported on a facet  $F_i$  which is located on  $\Gamma$  or the subdomain interfaces 144 in  $W_{I,i}$ . Thus, in each preconditioner step a forward block Gauss Seidel iteration is 145 carried out, followed by a direct inversion of each subdomain block and a backward 146 block Gauss Seidel step. Note that solving directly for the unknowns in a subdomain 147 is equivalent to solve a problem with robin boundary conditions on the subdomain, 148 and uniqueness and existence are guaranteed.

One big advantage of the DD preconditioner is, that it can cope with problems 150 containing cavity like structures. For such problems other preconditioners suffer 151 from internal reflections, which leads to high iteration numbers. If the whole cavity 152 is contained in one single subdomain  $\Omega_i$ , the DD preconditioner inverts the whole 153 matrix block related to the cavity, and internal reflections are treated exactly. Thus they do not influence the iteration number.

### 4 Numerical Results

In order to demonstrate the dependence of the number of iterations on polynomial 157 order, wavelength and meshsize h for the presented preconditioners, we choose a 158 simple two dimensional model problem with a wave of Gaussian amplitude and 159 wavelength  $\lambda$  propagating through a unit square domain (compare Fig. 1). For a 160 meshsize  $h=\lambda=0.1$  the lefthand plot shows the number of iterations for different polynomial orders. For the three preconditioners, the DoFs of an element were collected in one block. In addition, for the DD preconditioner, the computational domain was divided into nine subdomains. If the polynomial order is large enough to resolve the wave, i.e. larger than four, the number of iterations stays constant or is only slightly growing with growing polynomial order, while the number of facet unknowns grows linearly in 2D.

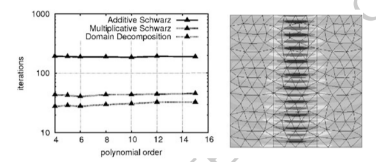


Fig. 1. Iterations depending on the polynomial order (left) for the 2D model problem (right)

**Table 1.** Iterations depending on wavelength and mesh size for the MS/DD Preconditioner (p = 6).

3).								
λ	0.64	0.32	0.16	0.08	0.04	0.02	0.01	
h = 0.16	35/40	35/38	32/33	31/31				
h = 0.08	52/42	48/38	50/36	47/33	50/38			,
h = 0.04	88/55	76/47	74/43	76/39	65/35	97/59		,
h = 0.02	147/75	129/55	113/48	117/44	118/42	115/38	199/82	,
h = 0.01	246/107	236/80	226/60	203/53	228/49	271/50	291/45	,

Next we investigate the dependence on h and  $\lambda$  for a fixed polynomial order of 6. The results are presented in Table 1. For  $\lambda$  smaller than  $\frac{h}{2}$ , which corresponds 169 to less than three unknowns per wavelength, the solution can not be resolved, and 170 the solvers show large iteration numbers. Fixing h, the iteration number is mini-171 mal at about  $h \approx \lambda$ , i.e. at about six unknowns per wavelength, and it increases for 172 growing wavelength. For h=0.16 every subdomain consists of only a small number 173 of elements, and an inversion of the DoFs subdomain by subdomain is compare-174 able to an inversion element by element. Therefore the two preconditioners show 175 about the same performance. If h decreases, it is more and more advantageous to 176 collect the unknowns in subdomain blocks. While the iteration number almost doubles for the MS preconditioner if the mesh size is divided by 2, the increase is much 178

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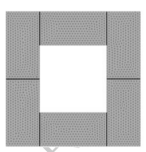
less for the DD preconditioner. Table 2 shows, that the DD preconditioner also per- 179 forms better than the MS preconditioner with respect to time, although one iteration 180 is more expensive.

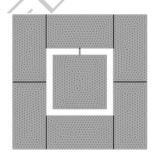
**Table 2.** Iteration times for  $\lambda =$ 0.08 and a polynomial order of 6.

h	DoFs	MS	DD
0.16	69980	0.35	0.37
0.08	217900	1.73	1.33
0.04	701228	9.30	5.15
0.02	2518524	53.5	22.4
0.01	9857920	367	111

Table 3. Iteration numbers and computational times for the cavity and the square.

	ca	vity	square		
	its.	time(s)	its.	time(s)	
DD (element)	35	40.4	34	31.2	
DD (facet)	64	69.7	61	59.7	
MS (element)		1720	102	88.9	
AS (element)	$> 10^5$	> 1h	575	186	





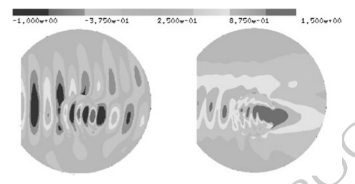
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**Fig. 2.** A resonator (*right*) is compared with the domain without cavity (*left*)

Now we compare the preconditioners for a resonator and the domain without 182 cavity (compare Fig. 2). From the top of the square an incident wave with  $\lambda = 0.01$  183 is prescribed. The DD-preconditioner uses, depending on the presence of the cavity 184 six and seven subdomains, respectively, where all cavity DoFs, including the cavity 185 boundary are collected in one single block. Table 3 shows the iteration numbers 186 and computational times for different preconditioners and for the two examples. For 187 the domain without cavity the performance of the preconditioners is compareable. 188 When the cavity is added, reflections inside the cavity lead to an enormous increase 189 in iteration numbers and computational times for the AS and the MS preconditioner. 190 Because of direct inversion of the cavity DoFs, the DD preconditioner does not suffer 191 from internal reflections and the iteration number stays almost constant, which leads 192 together with a larger number of unknowns to a moderate increase in computational 193 time.

We finish the numerical results section with an example from optics. A small 195 sphere with radius 0.3 and refractive index 2 is placed (not exactly in the center) in 196 a spherical computational domain with radius 1 and background refractive index 1. 197

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**Fig. 3.** Real part of  $E_v$  (*left*) and |E| (*right*) evaluated at a cross section parallel to the xy plane

We prescribe an incident wave from the left with a Gaussian amplitude and wavelength 0.35, such that the diameter of the computational domain is approximately six 199 wavelength in free space. In order to resolve the wave we used 3,256 elements with 200 a polynomial order of 6, which results in 1.66 millions of unknowns. The solution 201 was obtained by 258 cg-iterations with a Block AS preconditioner (Fig. 3). 202

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# Multiscale Domain Decomposition Preconditioners for 2 **Anisotropic High-Contrast Problems**

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1 Summary

In this paper, we study robust two-level domain decomposition preconditioners for 10 highly anisotropic multiscale problems. We present a construction of coarse spaces 11 that emploies initial multiscale basis functions and discuss techniques to achieve 12 smaller dimensional coarse spaces without sacrificing the robustness of the preconditioner. We also present numerical results and consider possible extensions of these 14 approaches where the dimension of the coarse space can be reduced further.

2 Introduction 16

Anisotropy in the diffusion arises in many applications in geosciences and engi- 17 neering. In flows porous media, high anisotropy can be due to the presence of frac- 18 tures that may have preferred high-conductivity directions. Because of high varia- 19 tions among the matrix and fracture conductivities, the permeability can have high 20 anisotropy at the fine-scale. This is the case when fracture network conducts only in 21 some preferred directions (e.g., in one direction in 2D problems and one or two di- 22 rections in 3D problems). This preferred direction is the direction of high anisotropy 23 and it can have heterogeneous spatial variations. For example, the presence of frac- 24 ture pockets can create highly anisotropic isolated regions, while fracture corridors 25 can form long highly anisotropic channels that span a rich hierarchy of scales. It is a 26 challenging task to design robust preconditioners for such problems (e.g., [4]) or to 27 solve them on a coarse grid (e.g., [2]).

In this paper, we discuss robust preconditioners for highly anisotropic multiscale 29 diffusion problems. We assume that the high-anisotropy is also highly heterogeneous 30 over the problem domain and these spatial variations cannot be captured within a 31 coarse block. In the paper, robust two-level domain decomposition preconditioners 32 are constructed by designing coarse spaces that contain essential features of the finescale solution. The construction of the coarse spaces is based on recently introduced 34

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methods [1, 3]. We show that, for anisotropic problems, the coarse spaces can have 35 a large dimension because fine-scale features within high-anisotropy regions need 36 to be represented on a coarse grid. In this paper, we propose a number of remedies 37 for this problem. Note that the proposed methods differ from existing methods for 38 anisotropic problems [4].

The coarse spaces used in two-level domain decomposition preconditioners are 40 constructed based on local spectral problems with a pre-computed scalar weight 41 function. The computation of the weight function uses an initial coarse space where 42 one basis function per coarse node is defined. We show that the local eigenvalue 43 problem can contain many small eigenvalues, which are asymptotically vanishing as 44 the contrast increases. One needs to include all eigenvectors that correspond to these 45 small, asymptotically vanishing, eigenvalues. Because the number of these small 46 eigenvalues defines the dimension of the coarse space, it is important to choose a 47 weight function such that the dimension of the coarse space is as small as possible. 48 If we consider the initial space as the span of piecewise (bi)linear functions, then the 49 dimension of the coarse space can be very large. In particular, the coarse space contains all fine-scale functions with respect to the slow variable (defined as the variable 51 representing the direction of slow conductivity) within high-anisotropy regions. On 52 the other hand, using multiscale basis functions [2] in the initial space allows cap- 53 turing the effects of high-conductivity inclusions (cf. [1, 3]) that are isolated within 54 coarse grid blocks. As a result, the coarse space contains all fine-scale functions 55 with respect to slow variables within high-anisotropy channels. This can lead to a 56 substantial dimension reduction; however, unlike to the isotropic high-conductivity 57 case, the dimension of the coarse space can still be very large as discussed in the 58 paper. Numerical results are presented. We also discuss techniques that allow us to 59 use smaller dimensional coarse spaces at the expenses of solving several lower di- 60 mensional problems in the channels of high-anisotropy.

## 3 Problem Setting and Domain Decomposition Framework

Let  $D \subset \mathbb{R}^2$  (or  $\mathbb{R}^3$ ) be a polygonal domain which is the union of a disjoint polygonal 63 subregions  $\{D_i\}_{i=1}^N$ . We seek  $u \in H_0^1(D)$ 

$$a(u,v) := \int_D \kappa(x) \nabla u \cdot \nabla v dx = \int_D f v dx, \text{ where } \kappa(x) = \begin{pmatrix} \eta(x) & 0 \\ 0 & 1 \end{pmatrix}. \tag{1}$$

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Here  $\eta(x)$  is a heterogeneous field with high contrast,  $\eta(x) \ge 1$ . More general cases 65 where the direction of anisotropy can change continuously in space will be considered elsewhere. Next, we introduce some notations following [1].

We assume that  $\{D_i\}_{i=1}^N$  form a quasiuniform triangulation of D and denote H=68  $\max_i \operatorname{diam}(D_i)$ . Let  $\mathscr{T}^h$  be a fine triangulation which refine  $\{D_i\}_{i=1}^N$ . We denote by 69  $V^h(D)$  the usual finite element discretization of piecewise linear continuous functions 70 with respect to the fine triangulation  $\mathcal{T}^h$ . Denote also by  $V_0^h(D)$  the subset of  $V^h(D)$ with vanishing values on  $\partial D$ . Similar notations,  $V^h(\Omega)$  and  $V^h_0(\Omega)$ , are used for 72 subdomains  $\Omega \subset D$ . 73

The Galerkin finite element approximation of (1) is to find  $u \in V_0^h(D)$  with 74  $a(u,v) = \int_D fv$  for all  $v \in V_0^h(D)$ , or in matrix form

$$Au = b, (2)$$

where for all  $u, v \in V^h(D)$  (considered as vectors) we have  $v^T A u = a(u, v)$  and  $v^T b = 76$  $\int_D f v$ . We assume that  $\kappa$  is piecewise constant coefficient in  $\mathcal{T}^h$  with value  $\kappa = \kappa_e =$  $(\eta_e, 0; 0, 1)$  on each fine triangulation element  $e \in \mathcal{T}^h$ .

We denote by  $\{D_i^{\prime}\}_{i=1}^{N}$  the overlapping decomposition obtained from the original nonoverlapping decomposition  $\{D_i\}_{i=1}^N$  by enlarging each subdomain  $D_i$  to  $D_i'=80$  $D_i \cup \{x \in D, \operatorname{dist}(x, D_i) < \delta_i\}, \quad i = 1, \dots, N$ , where dist is some distance function and 81 let  $\delta = \max_{1 \le i \le N} \delta_i$ . Let  $V_0^h(D_i')$  be the set of finite element functions with support 82 in  $D_i'$ . We also denote by  $R_i^T: V_0^h(D_i') \to V^h(D)$  the extension by zero operator. 83 We use a partition of unity  $\{\xi_i\}_{i=1}^N$  subordinated to the covering  $\{D_i'\}_{i=1}^N$  such 84

that

$$\sum_{i=1}^{N} \xi_i = 1, \quad \xi_i \in V^h(D), \quad 0 \le \xi_i \le 1 \quad \text{ and } \quad \operatorname{Supp}(\xi_i) \subset D_i', \ i = 1, \dots, N, \quad (3)$$

where  $\operatorname{Supp}(\xi_i)$  stands for the support of the function  $\xi_i$ . This partition of unity is 86 used to truncate global functions to local conforming functions, an essential property 87 in the construction of a stable splitting of the space.

Given a coarse triangulation  $\mathscr{T}^H$ , we introduce  $N_c$  coarse basis functions  $\{\Phi_i\}_{i=1}^{N_c}$ . 89 We define the coarse space by  $V_0^H = \operatorname{span}\{\Phi_i\}_{i=1}^{N_c}$ , and the coarse matrix  $A_0 = R_0AR_0^T$  where  $R_0^T = [\Phi_1, \dots, \Phi_{N_c}]$ . We use a two level additive preconditioner of the form

$$B^{-1} = R_0^T A_0^{-1} R_0 + \sum_{i=1}^N R_i^T A_i^{-1} R_i = R_0^T A_0^{-1} R_0 + B_{1L}^{-1}, \tag{4}$$

where  $B_{1L}^{-1} = \sum_{i=1}^{N} R_i^T A_i^{-1} R_i$  and the local matrices are defined by  $vA_i w = a(v, w)$  for 92 all  $v, w \in V_0^h(D_i')$ ,  $i = 1, \dots, N$  (see [5]).

We denote by  $\{y_i\}_{i=1}^{N_v}$  the vertices of the coarse mesh  $\mathcal{T}^H$  and define 94

$$\omega_i = \bigcup \{ K \in \mathcal{T}^H; \ y_i \in \overline{K} \}, \ \omega_K = \bigcup \{ \omega_i; \ y_i \in \overline{K} \}.$$
 (5)

Additionally, we use a partition of unity  $\{\chi_i\}_{i=1}^{N_v}$  subordinated to the covering 95  $\{\omega_i\}_{i}^{N_v}$  such that

$$\sum_{i=1}^{N_{\nu}} \chi_i = 1, \quad \chi_i \in V^h(D), \quad 0 \le \chi_i \le 1 \quad \text{and} \quad \text{Supp}(\chi_i) \subset \omega_i, i = 1, \dots, N_{\nu}. \quad (6)$$

## **Coarse Space Construction and Dimension Reduction**

In this section we define a local spectral multiscale coarse space using eigenvectors of 98 high-anisotropy eigenvalue problems. First we introduce the notation for eigenvalue 99 problems following [1]. For  $i=1,\ldots,N_{\nu}$ , define the matrix  $A^{\omega_i}$  and the *modified* 100 *mass matrix* of same dimension  $M^{\omega_i}$  by

$$v^T A^{\omega_i} w = \int_{\omega_i} \kappa \nabla v \cdot \nabla w dx \text{ and } v^T M^{\omega_i} w = \int_{\omega_i} \widetilde{\kappa} v w dx \ \forall v, w \in \widetilde{V}^h(\omega_i),$$
 (7)

where  $\widetilde{V}^h(\omega_i) = \{v \in V^h(\omega_i) : v = 0 \text{ on } \partial \omega_i \cap \partial D\}$ . Here  $\widetilde{\kappa}$  is an scalar weight derived from the high-anisotropy coefficient matrix  $\kappa = [\kappa_{ij}]$  and contains the relevant information we need for the construction of the coarse basis functions. Several possible the choices for  $\widetilde{\kappa}$  can be considered. Here  $\widetilde{\kappa}$  is defined by

$$\widetilde{\kappa} = \max \left\{ \sum_{i=1}^{N} \kappa \nabla \xi_i \cdot \nabla \xi_i, \sum_{j=1}^{N_{\nu}} \kappa \nabla \chi_j \cdot \nabla \chi_j \right\}, \tag{8}$$

where  $\{\xi_j\}_{j=1}^N$  and  $\{\chi_i\}_{i=1}^{N_v}$  are the partition of unity introduced in (3) and (6), respectively. From now on, we assume that the overlapping decomposition is constructed from the coarse mesh and then  $\xi_i=\chi_i$  and  $D_i'=\omega_i$  for all  $i=1,\ldots,N=N_v$ , 108 and  $\delta \asymp H$ . We consider the finite dimensional symmetric eigenvalue problems 109  $A^{\omega_i}\psi=\widetilde{\lambda}M^{\omega_i}\psi$ , with  $A^{\omega_i}$  and  $M^{\omega_i}$  defined by (7) and (8),  $i=1,\ldots,N$ . Denote its 110 eigenvalues and eigenvectors by  $\{\widetilde{\lambda}_\ell^{\omega_i}\}$  and  $\{\psi_\ell^{\omega_i}\}$ , respectively. Note that the eigenvectors  $\{\psi_\ell^{\omega_i}\}$  form an orthonormal basis of  $\widetilde{V}^h(\omega_i)$  with respect to the  $M^{\omega_i}$  inner 112 product. Assume that  $\widetilde{\lambda}_1^{\omega_i} \le \widetilde{\lambda}_2^{\omega_i} \le \cdots \le \widetilde{\lambda}_\ell^{\omega_i} \le \cdots$ , and note that  $\widetilde{\lambda}_1^{\omega_i} = 0$  for all 113 interior subdomains. In particular,  $\psi_\ell^{\omega_i}$  denotes the  $\ell$ -th eigenvector of the matrix 114 associated to the neighborhood of  $y_i$ ,  $i=1,\ldots,N_v$ .

Let  $\{\chi_i\}_{i=1}^{N_v}$  be a partition of unity (3). Define the coarse basis functions

$$\Phi_{i,\ell} = I^h(\chi_i \psi_\ell^{\omega_i}) \quad \text{ for } 1 \le \ell \le L_i \text{ and } 1 \le i \le N_v,$$
(9)

where  $I^h$  is the fine-scale nodal value interpolation and  $L_i$  is an integer number for each  $i = 1, ..., N_v$ . Denote by  $V_0^H$  the *spectral multiscale* space

$$V_0^H = \text{span}\{\Phi_{i,\ell} : 1 \le \ell \le L_i \text{ and } 1 \le i \le N_v\}.$$
 (10)

The idea is to use only eigenvectors of contrast dependent eigenvalues. Next, we discuss how the choice of  $\widetilde{\kappa}$  affects the eigenvalues. If we choose  $\chi_i$  to be piecewise linear functions on the coarse grid, then, it is easy to see that we have  $\widetilde{\kappa}(x_1,x_2) = \sum_i \eta(x_1,x_2) |\partial_{x_1}\chi_i(x_1,x_2)|^2 + |\partial_{x_2}\chi_i(x_1,x_2)|^2$  and  $\widetilde{\kappa}$  will have similar behavior as  $\eta(x)$ . In this case, one can show that the number of small eigenvalues is the same as the fine degrees of freedom in the form of discrete functions that depend on  $x_2$  within high-anisotropy inclusions and channels. Indeed, if we consider the associated Rayleigh quotient,  $R(\nu) = \frac{\nu^T A^{\omega_i} \nu}{\nu^T M^{\omega_i} \nu}$ , we have

$$R(v) = \frac{\int_{\omega_i} \kappa \nabla v \cdot \nabla v}{\int_{\omega_i} \widetilde{\kappa} v^2} = \frac{\int_{\omega_i} \eta(x_1, x_2) |\partial_{x_1} v(x_1, x_2)|^2 + |\partial_{x_2} v(x_1, x_2)|^2}{\int_{\omega_i} (\sum_i \eta(x_1, x_2) |\partial_{x_1} \chi_i(x_1, x_2)|^2 + |\partial_{x_2} \chi_i(x_1, x_2)|^2) v(x_1, x_2)^2}.$$
 127

Then, for functions that depends only on  $x_2$  inside the region R where  $\eta$  is high, 128 the numerator reduces to  $\int_{\omega_i \setminus R} (|\partial_{x_1} v(x_1, x_2)|^2 + |\partial_{x_2} v(x_1, x_2)|^2) + \int_R |\partial_{x_2} v(x_1, x_2)|^2$  129

(which is independent of the high value of  $\eta(x)$  in R) and the quotient will go to zero 130 as the value of  $\eta$  in R goes to infinity. Including all fine grid functions of  $x_2$  into the coarse space can lead to a high dimensional coarse spaces. Note that the dimension 132 of the coarse space will be much higher than the case with scalar coefficient  $\kappa$  where the number of small eigenvalues is equal to the number of isolated inclusions and 134 channels within a coarse block; see [1, 3]. To reduce the dimension of the coarse 135 space, we propose the use of multiscale basis functions.

We are interested in partition of unity functions that can reduce the number of 137 degrees of freedom associated with isolated high-anisotropy inclusions. This can be 138 achieved by minimizing high-conductivity components for the scalar function  $\tilde{\kappa}$ . In particular, by choosing multiscale finite element basis functions or energy minimiz- 140 ing basis functions (e.g., [6]), we can eliminate all isolated high-conductivity inclusions. This can be observed in our numerical experiments. We recall the definition of 142 the "standard" multiscale finite element basis functions that coincide with (the piece- 143 wise linear functions on the coarse grid)  $\chi_i^0$  on the boundaries of the coarse partition. 144 They are denoted by  $\chi_i^{ms}$  and satisfy:

$$-\operatorname{div}(\kappa \nabla \chi_i^{ms}) = 0 \text{ in } K \in \omega_i, \quad \chi_i^{ms} = \chi_i^0 \text{ in } \partial K, \ \forall \ K \in \omega_i,$$
 (11)

where K is a coarse grid block within  $\omega_0$ , see [2] for more details and more general 146 multiscale basis functions constructions. In Fig. 1, we depict  $\eta(x)$  (left picture) and  $\widetilde{\kappa}$  147 (right picture) using multiscale basis functions on the coarse grid. One can observe 148 that isolated inclusions are removed in  $\kappa$ . The coarse space contains functions de- 149 pending only on  $x_2$  within long channels. The situation is more complicated if highanisotropy regions form complex channel patterns. For example, if high-anisotropy 151 region is vertical for the coefficients considered in our numerical example, then initial multiscale spaces can represent them and no additional degrees are needed. More 153 complex channel shapes will be studied elsewhere.

We note that for the proposed methods, in each  $\omega_i$ ,  $i=1,\ldots,N_{\nu}$ , we only need to specify the number of eigenvectors  $L_i$  based on the quantities  $\{1/\widetilde{\lambda}_l^{\omega_i}\}$ . These eigenvectors are used to construct the coarse space. In practice, one only needs to compute the first L<sub>i</sub> eigenvalues. Hierarchical approximation with several triangulations can 158 also be considered for the eigenvalues and eigenvectors.

Weighted  $L^2$  approximation and weighted  $H^1$  stability properties of the coarse 160 space  $V_0^H$  in (10) hold (as in [1, 3]). In order to describe better these properties of  $V_0^H$ , we need to introduce a relevant interpolation operator. Given  $v \in V^h(\omega_i)$ , set

$$I_{L_i}^{\omega_i} v = \sum_{\ell=1}^{L_i} \left( \int_{\omega_i} \widetilde{\kappa} v \psi_\ell^{\omega_i} dx \right) \psi_\ell^{\omega_i}, \quad i = 1, \dots, N_v,$$
 (12)

and define the coarse interpolation  $I_0: V^h(D) \to V_0^H$  by

$$I_{0}v = \sum_{i=1}^{N_{v}} \sum_{\ell=1}^{L_{i}} \left( \int_{\omega_{i}} \widetilde{\kappa} v \psi_{\ell}^{\omega_{i}} dx \right) I^{h}(\chi_{i} \psi_{\ell}^{\omega_{i}}) = \sum_{i=1}^{N_{v}} I^{h} \left( \chi_{i} (I_{L_{i}}^{\omega_{i}} v) \right), \tag{13}$$

where  $I^h$  is the fine-scale nodal value interpolation.

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**Lemma 1.** For each coarse element K we have

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$$\int_K \widetilde{\kappa}(v - I_0 v)^2 \preceq \widetilde{\lambda}_{K,L+1}^{-1} \int_{\omega_K} \kappa \nabla v \cdot \nabla v dx$$

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• 
$$\int_{K} \kappa \nabla I_{0} v \cdot \nabla I_{0} v dx \leq \max\{1, \widetilde{\lambda}_{K,L+1}^{-1}\} \int_{\omega_{K}} \kappa \nabla v \cdot \nabla v dx,$$

where 
$$\widetilde{\lambda}_{K,L+1} = \min_{y_i \in K} \widetilde{\lambda}_{L_i+1}^{\omega_i}$$
 and  $\omega_K$  is defined in (5).

Using Lemma 1, we can estimate the condition number of the preconditioned 169 operator  $B^{-1}A$  with  $B^{-1}$  defined in (4) using the coarse space  $V_0^H$  in (10). Following [1, 3], one has the following result.

**Theorem 1.** The condition number,  $cond(B^{-1}A)$ , of the preconditioned operator 172  $B^{-1}A$  with  $B^{-1}$  defined in (4) satisfies 173

$$cond(B^{-1}A) \leq 1 + \widetilde{\lambda}_{L+1}^{-1}, \quad where \quad \widetilde{\lambda}_{L+1} = \min_{1 \leq i \leq N_v} \widetilde{\lambda}_{L_i+1}^{\omega_i}.$$
 174

Recall that we assumed  $\xi_i = \chi_i$ ,  $i = 1, ..., N = N_v$ . It can be easily shown that if we choose  $L_i$  as the number of contrast dependent eigenvalues, then  $\lambda_{L+1}$  scales 176 as O(1), i.e., independent of the contrast. The dependency of the condition number 177 on  $\delta$  and H is controlled by the partition of unity  $\{\chi_i\}$ . The condition number is 178 independent of h and it is, in the general case of different partitions of unity,  $\{\chi_i\}$  179 and  $\{\xi_i\}$ , of order  $O(H^2/\delta^2)$ , see [3]. 180

### **5 Numerical Results**

In this section, we show representative 2D numerical results for the additive preconditioner (4) with the local spectral multiscale coarse space defined in (10). We take 183  $D = [0,1] \times [0,1]$  that is divided into  $10 \times 10$  equal square coarse blocks to construct 184 the coarse mesh. Inside each coarse block we use a fine-scale triangulation where 185 triangular elements constructed from  $10 \times 10$  squares are used.

We test our approach on a permeability field that contains inclusions and channels 187 on a background of conductivity one (see the left picture of Fig. 1 for  $\eta(x)$  in (1)). 188 We use multiscale finite element basis functions as the initial partition of unity. From 189 the right picture of Fig. 1 we see that the modified weight  $\widetilde{\kappa}$  does not contain any isolated inclusions and only contains long high-anisotropy channels connecting bound- 191 aries of coarse-grid blocks. This is automatically achieved from the choice of the 192 partition of unity functions. There are fewer small (asymptotically vanishing) eigen- 193 values when local eigenvalue problem is solved with the modified weight  $\widetilde{\kappa}$ . Thus, 194 a good choice of partition of unity functions  $\chi_i$  in (8) will ensure fewer new multiscale basis functions needed to achieve an optimal convergence with respect to the 196 contrast. Numerical results are presented in Table 1. We observe that using the pro- 197 posed coarse spaces, the number of iterations is independent of contrast. In Table 1 198 we also show the dimension of the coarse spaces. The dimension of the local spectral 199 coarse space is smaller if we use  $\tilde{\kappa}$  in (10) with multiscale basis functions instead of 200 piecewise linear basis functions.

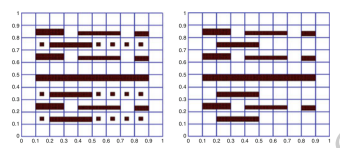


Fig. 1. Left: Coarse mesh and coefficient (we plot  $\eta(x)=10^6$  and recall that  $\eta(x)=1$  elsewhere). Right: Coefficient  $\widetilde{\kappa}$  in (8) using multiscale basis functions (we plot  $\widetilde{\kappa}(x) \geq 10^6$ ). See Table 1

η	LIN	MS	EMF	LSM (bilin. $\chi_i$ )	LSM (MS $\chi_i$ )
		122(1.51e+2)			55(26.9)
		258(1.28e+3)			28(5.82)
		483(1.26e+4)			29(6.02)
$10^{6}$	627(1.34e+5)	709(1.27e+5)	599(9.63e+4)	30(5.753)	29(6.04)
Dim	81=0.79%	81=0.79%	81=0.79%	732=7.19%	497=4.87%

**Table 1.** Number of iterations and estimated condition number for the PCG and various values of  $\eta$  with the coefficient depicted in Figure 1. We set the tolerance to 1e - 10, H = 1/10, h = 1/100, and  $\dim(V_h) = 10201$ . The notation MS stands for the (linear boundary condition) multiscale (MS) coarse space, EMF is the energy minimizing coarse space, see e.g., [6], and LSM is the local spectral multiscale coarse space defined in (10). We select the first L eigenvalues such that  $\tilde{\lambda}_L - \tilde{\lambda}_{L-1} > 0.05$  (which is and easy way to select the small eigenvalues- in this example, the value 0.05 was chose by trial-and-error).

## **6 Discussion on Coarse Space Dimension Reduction**

Now we discuss approaches to avoid the use of high-dimensional coarse spaces without sacrificing the efficiency of the preconditioner at the expense of solving problems in high-anisotropy channels. As was observed in the presented numerical tests, the strongly anisotropic channels cause a substantial increase of the size of the coarse space and the complexity of the method. To avoid this, we can replace the coarse solve  $R_0^T A_0^{-1} R_0$  in (4) by  $R_0^T \widetilde{A}_0^{-1} R_0 + R_{an}^T A_{an}^{-1} R_{an}$ . Here the matrix  $\widetilde{A}_0$  is a small dimensional coarse matrix. The matrix  $A_{an}$  is acting on the fine-mesh degrees restricted to subdomain of high-anisotropy channels  $\Omega_{an}$ . It is based on the original matrix A and is constructed locally (element-by-element) by preserving the strongest links (off-diagonal entries) of the element stiffness matrices in the channels. To illustrate this idea, which was developed in [4] for Crouzeix-Raviart elements, we write an element stiffness matrix  $A_e$  for  $e \subset \Omega_{an}$ :  $A_e = [b_e + c_e, -c_e, -b_e; -c_e, a_e + 214 c_e, -a_e; -b_e, -a_e, a_e + b_e]$ , where  $|a_e| \le b_e \le c_e$ . Then the matrix  $A_{an}$  is defined as assembly of the matrices  $B_e = [c_e, -c_e, 0; -c_e, c_e, 0; 0, 0, 0]$ ,  $e \subset \Omega_{an}$ . It is easy 216

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t1.1 t1.2 t1.3 t1.4 t1.5

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to see that  $A_{an}$  is a stiffness matrix corresponding to a diffusion problem defined 217 on a carcass of piecewise linear lines in  $\Omega_{an}$  following the directions of dominating 218 anisotropy.

In the case of apparent dominant anisotropy direction (i.e., when A<sub>an</sub> is block 220 diagonal with tridiagonal blocks), inverting  $A_{an}$  will involve solving block-diagonal 221 problems with tridiagonal blocks (in 2-D only). In this case optimal complexity is 222 achieved by using a sparse direct solver. In general, one may consider including 223 some of the degrees of freedom associated with high-anisotropy regions into the 224 coarse space while using  $A_{an}^{-1}$  to handle the others. Another possibility is to use an 225 auxiliary space of Crouzeix-Raviart elements combined with the technique from [4]. 226 These issues will be studied in our subsequent work. 227

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## A Robust FEM-BEM Solver for Time-Harmonic Eddy **Current Problems**

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Summary. This paper is devoted to the construction and analysis of robust solution techniques for time-harmonic eddy current problems in unbounded domains. We discretize the 10 time-harmonic eddy current equation by means of a symmetrically coupled finite and boundary 11 element method, taking care of the different physical behavior in conducting and non- 12 conducting subdomains, respectively. We construct and analyse a block-diagonal preconditioner for the system of coupled finite and boundary element equations that is robust with 14 respect to the space discretization parameter as well as all involved "bad" parameters like the 15 frequency, the conductivity and the reluctivity. Block-diagonal preconditioners can be used 16 for accelerating iterative solution methods such like the Minimal Residual Method.

1 Introduction 18

In many practical applications, the excitation is time-harmonic. Switching from the time domain to the frequency domain allows us to replace expensive time-integration 20 procedures by the solution of a system of partial differential equations for the am- 21 plitudes belonging to the sine- and to the cosine-excitation. Following this strat- 22 egy, [7, 13] and [4, 5] applied harmonic and multiharmonic approaches to parabolic 23 initial-boundary value problems and the eddy current problem, respectively. Indeed, 24 in [13], a preconditioned MinRes solver for the solution of the eddy current problem 25 in bounded domains was constructed that is robust with respect to both the discretiza- 26 tion parameter h and the frequency  $\omega$ . The key point of this parameter-robust solver 27 is the construction of a block-diagonal preconditioner, where standard **H(curl)** FEM 28 magneto-static problems have to be solved or preconditioned. The aim of this con- 29 tribution is to generalize these ideas to the case of unbounded domains in terms of 30 a coupled Finite Element (FEM) – Boundary Element (BEM) Method. In this case 31 we are also able to construct a block-diagonal preconditioner, where now standard 32 coupled FEM-BEM **H(curl)** problems, as arising in the magneto-static case, have 33 to be solved or preconditioned. We mention, that this preconditioning technique fits 34 into the framework of operator preconditioning, see, e.g. [1, 11, 16, 19].

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The paper is now organized as follows. We introduce the frequency domain 36 equations in Sect. 2. In the same section, we provide the symmetrically coupled 37 FEM-BEM discretization of these equations. In Sect. 3, we construct and analyse our 38 parameter-robust block-diagonal preconditioner used in a MinRes setting for solving 39 the resulting system of linear algebraic equations. Finally, we discuss the practical 40 realization of our preconditioner.

#### 2 Frequency Domain FEM-BEM

As a model problem, we consider the following eddy current problem:

$$\begin{cases} \sigma \frac{\partial \mathbf{u}}{\partial t} + \mathbf{curl} \left( v_1 \, \mathbf{curl} \, \mathbf{u} \right) = \mathbf{f} & \text{in } \Omega_1 \times (0, T), \\ \mathbf{curl} \left( \mathbf{curl} \, \mathbf{u} \right) = \mathbf{0} & \text{in } \Omega_2 \times (0, T), \\ \text{div } \mathbf{u} = \mathbf{0} & \text{in } \Omega_2 \times (0, T), \\ \mathbf{u} = \mathcal{O}(|\mathbf{x}|^{-1}) & \text{for } |\mathbf{x}| \to \infty, \\ \mathbf{curl} \mathbf{u} = \mathcal{O}(|\mathbf{x}|^{-1}) & \text{for } |\mathbf{x}| \to \infty, \\ \mathbf{u} = \mathbf{u}_0 & \text{on } \Omega_1 \times \{0\}, \\ \mathbf{u}_1 \times \mathbf{n} = \mathbf{u}_2 \times \mathbf{n} & \text{on } \Gamma \times (0, T), \\ v_1 \mathbf{curl} \, \mathbf{u}_1 \times \mathbf{n} = \mathbf{curl} \, \mathbf{u}_2 \times \mathbf{n} & \text{on } \Gamma \times (0, T), \end{cases}$$

$$(1)$$

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where the computational domain  $\Omega=\mathbb{R}^3$  is split into the two non-overlapping sub- 44 domains  $\Omega_1$  and  $\Omega_2$ . The conducting subdomain  $\Omega_1$  is assumed to be a simply 45 connected Lipschitz polyhedron, whereas the non-conducting subdomain  $\Omega_2$  is the 46 complement of  $\Omega_1$  in  $\mathbb{R}^3$ , i.e  $\mathbb{R}^3 \setminus \overline{\Omega}_1$ . Furthermore, we denote by  $\Gamma$  the interface be- 47 tween the two subdomains, i.e.  $\Gamma = \overline{\Omega}_1 \cap \overline{\Omega}_2$ . The exterior unit normal vector of  $\Omega_1$  48 on  $\Gamma$  is denoted by  $\mathbf{n}$ , i.e.  $\mathbf{n}$  points from  $\Omega_1$  to  $\Omega_2$ . The reluctivity  $v_1$  is supposed to be 49 independent of  $|\mathbf{curl} \mathbf{u}|$ , i.e. we assume the eddy current problem (1) to be linear. The 50 conductivity  $\sigma$  is zero in  $\Omega_2$ , and piecewise constant and uniformly positive in  $\Omega_1$ . 51

We assume, that the source  $\mathbf{f}$  is given by a time-harmonic excitation with the 52 frequency  $\omega > 0$  and amplitudes  $\mathbf{f}^c$  and  $\mathbf{f}^s$  in the conducting domain  $\Omega_1$ . Therefore, 53 the solution  $\mathbf{u}$  is time-harmonic as well, with the same base frequency  $\omega$ , i.e. 54

$$\mathbf{u}(\mathbf{x},t) = \mathbf{u}^{\mathbf{c}}(\mathbf{x})\cos(\omega t) + \mathbf{u}^{\mathbf{s}}(\mathbf{x})\sin(\omega t). \tag{2}$$

In fact, (2) is the real reformulation of a complex time-harmonic approach  $\mathbf{u}(\mathbf{x},t) = 55$   $\hat{\mathbf{u}}(\mathbf{x})e^{\mathrm{i}\omega t}$  with the complex-valued amplitude  $\hat{\mathbf{u}} = \mathbf{u^c} - \mathrm{i}\mathbf{u^s}$ . Using the time-harmonic 56 representation (2) of the solution, we can state the eddy current problem (1) in the 57 frequency domain as follows:

Find 
$$\mathbf{u} = (\mathbf{u^c}, \mathbf{u^s})$$
: 
$$\begin{cases} \omega \sigma \mathbf{u^s} + \mathbf{curl} (v_1 \mathbf{curl} \mathbf{u^c}) = \mathbf{f^c} \text{ in } \Omega_1, \\ \mathbf{curl} \mathbf{curl} \mathbf{u^c} = \mathbf{0} \text{ in } \Omega_2, \\ -\omega \sigma \mathbf{u^c} + \mathbf{curl} (v_1 \mathbf{curl} \mathbf{u^s}) = \mathbf{f^s} \text{ in } \Omega_1, \\ \mathbf{curl} \mathbf{curl} \mathbf{u^s} = \mathbf{0} \text{ in } \Omega_2, \end{cases}$$
(3)

with the corresponding decay and interface conditions from (1).

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Remark 1. In practice, the reluctivity  $v_1$  depends on the inductivity |**curlu**| in a non- 60 linear way in ferromagnetic materials. Having in mind applications to problems with 61 nonlinear reluctivity, we prefer to use the real reformulation (3) instead of a complex 62 approach. For overcoming the nonlinearity the preferable way is to apply Newton's 63 method due to its fast convergence. It turns out, that Newton's method cannot be ap- 64 plied to the nonlinear complex-valued system (see [4]), but it can be applied to the 65 reformulated real-valued system. Anyhow, the analysis of the linear problem also 66 helps to construct efficient solvers for the nonlinear problem.

Deriving the variational formulation and integrating by parts once more in the exte-68 rior domain yields: Find  $(\mathbf{u}^{\mathbf{c}}, \mathbf{u}^{\mathbf{s}}) \in \mathbf{H}(\mathbf{curl}, \Omega_1)^2$  such that

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$$\begin{cases} \omega(\sigma\mathbf{u}^{\mathbf{s}},\mathbf{v}^{\mathbf{c}})_{L_{2}(\Omega_{1})} + (\nu_{1}\mathbf{curl}\,\mathbf{u}^{\mathbf{c}},\mathbf{curl}\,\mathbf{v}^{\mathbf{c}})_{L_{2}(\Omega_{1})} - \langle \gamma_{\!\!N}\mathbf{u}^{\mathbf{c}},\gamma_{\!\!D}\mathbf{v}^{\mathbf{c}}\rangle_{\tau} = \langle \mathbf{f}^{\mathbf{c}},\mathbf{v}^{\mathbf{c}}\rangle, \\ -\omega(\sigma\mathbf{u}^{\mathbf{c}},\mathbf{v}^{\mathbf{s}})_{L_{2}(\Omega_{1})} + (\nu_{1}\mathbf{curl}\,\mathbf{u}^{\mathbf{s}},\mathbf{curl}\,\mathbf{v}^{\mathbf{s}})_{L_{2}(\Omega_{1})} - \langle \gamma_{\!\!N}\mathbf{u}^{\mathbf{s}},\gamma_{\!\!D}\mathbf{v}^{\mathbf{s}}\rangle_{\tau} = \langle \mathbf{f}^{\mathbf{s}},\mathbf{v}^{\mathbf{s}}\rangle, \end{cases}$$

for all  $(\mathbf{v}^{\mathbf{c}}, \mathbf{v}^{\mathbf{s}}) \in \mathbf{H}(\mathbf{curl}, \Omega_1)^2$ . Here  $\gamma_D$  and  $\gamma_N$  denote the Dirichlet trace  $\gamma_D := \mathbf{n} \times 70$  $(\mathbf{u} \times \mathbf{n})$  and the Neumann trace  $\gamma_N := \mathbf{curl} \, \mathbf{u} \times \mathbf{n}$  on the interface  $\Gamma$ .  $\langle \cdot, \cdot \rangle_{\tau}$  denotes the 71  $L_2(\Gamma)$ -based duality product. In order to deal with the expression on the interface 72  $\Gamma$ , we use the framework of the symmetric FEM-BEM coupling for eddy current 73 problems (see [10]). So, using the boundary integral operators A, B, C and N, as 74 defined in [10], we end up with the weak formulation of the time-harmonic eddy 75 current problem: Find  $(\mathbf{u^c}, \mathbf{u^s}) \in \mathbf{H}(\mathbf{curl}, \Omega_1)^2$  and  $(\lambda^c, \lambda^s) \in \mathbf{H}_{\parallel}^{-\frac{1}{2}}(\text{div}_{\Gamma}0, \Gamma)^2$  such 76 that

$$\begin{cases} \boldsymbol{\omega}(\boldsymbol{\sigma}\mathbf{u}^{\mathbf{s}}, \mathbf{v}^{\mathbf{c}})_{L_{2}(\Omega_{1})} + (\boldsymbol{v}_{1}\mathbf{curl}\mathbf{u}^{\mathbf{c}}, \mathbf{curl}\mathbf{v}^{\mathbf{c}})_{L_{2}(\Omega_{1})}, \\ -\langle \mathbf{N}(\gamma_{D}\mathbf{u}^{\mathbf{c}}), \gamma_{D}\mathbf{v}^{\mathbf{c}}\rangle_{\tau} + \langle \mathbf{B}(\lambda^{\mathbf{c}}), \gamma_{D}\mathbf{v}^{\mathbf{c}}\rangle_{\tau} = \langle \mathbf{f}^{\mathbf{c}}, \mathbf{v}^{\mathbf{c}}\rangle, \\ \langle \boldsymbol{\mu}^{\mathbf{c}}, (\mathbf{C} - \mathbf{Id})(\gamma_{D}\mathbf{u}^{\mathbf{c}})\rangle_{\tau} - \langle \boldsymbol{\mu}^{\mathbf{c}}, \mathbf{A}(\lambda^{\mathbf{c}})\rangle_{\tau} = 0, \\ -\boldsymbol{\omega}(\boldsymbol{\sigma}\mathbf{u}^{\mathbf{c}}, \mathbf{v}^{\mathbf{s}})_{L_{2}(\Omega_{1})} + (\boldsymbol{v}_{1}\mathbf{curl}\mathbf{u}^{\mathbf{s}}, \mathbf{curl}\mathbf{v}^{\mathbf{s}})_{L_{2}(\Omega_{1})}, \\ -\langle \mathbf{N}(\gamma_{D}\mathbf{u}^{\mathbf{s}}), \gamma_{D}\mathbf{v}^{\mathbf{s}}\rangle_{\tau} + \langle \mathbf{B}(\lambda^{\mathbf{s}}), \gamma_{D}\mathbf{v}^{\mathbf{s}}\rangle_{\tau} = \langle \mathbf{f}^{\mathbf{s}}, \mathbf{v}^{\mathbf{s}}\rangle, \\ \langle \boldsymbol{\mu}^{\mathbf{s}}, (\mathbf{C} - \mathbf{Id})(\gamma_{D}\mathbf{u}^{\mathbf{s}})\rangle_{\tau} - \langle \boldsymbol{\mu}^{\mathbf{s}}, \mathbf{A}(\lambda^{\mathbf{s}})\rangle_{\tau} = 0, \end{cases}$$
(4)

for all  $(\mathbf{v^c}, \mathbf{v^s}) \in \mathbf{H}(\mathbf{curl}, \Omega_1)^2$  and  $(\mu^c, \mu^s) \in \mathbf{H}_{\parallel}^{-\frac{1}{2}}(\operatorname{div}_{\Gamma}0, \Gamma)^2$ . This variational form 78 is the starting point of the discretization in space. Therefore, we use a regular triangulation  $\mathcal{T}_h$ , with mesh size h > 0, of the domain  $\Omega_1$  with tetrahedral elements.  $\mathcal{T}_h$  80 induces a mesh  $\mathcal{K}_h$  of triangles on the boundary  $\Gamma$ . On these meshes, we consider 81 Nédélec basis functions of order p yielding the conforming finite element subspace 82  $\mathcal{N}\mathcal{D}_p(\mathcal{T}_h)$  of  $\mathbf{H}(\mathbf{curl},\Omega_1)$ , see [17]. Further, we use the space of divergence free 83 Raviart-Thomas basis functions  $\mathscr{RT}^0_p(\mathscr{K}_h) := \{\lambda_h \in \mathscr{RT}_p(\mathscr{K}_h), \operatorname{div}_{\Gamma} \lambda_h = 0\}$  being 84 a conforming finite element subspace of  $\mathbf{H}_{\parallel}^{-\frac{1}{2}}(\mathrm{div}_{\Gamma}0,\Gamma)$ . Let  $\{\pmb{\varphi}_i\}$  denote the basis 85 of  $\mathcal{ND}_p(\mathcal{T}_h)$ , and let  $\{\psi_i\}$  denote the basis of  $\mathcal{RT}_p^0(\mathcal{K}_h)$ . Then the matrix entries 86 corresponding to the operators in (4) are given by the formulas

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$$\begin{split} (\mathbf{K})_{\mathit{ij}} &:= (\nu \; \mathbf{curl} \, \phi_i, \mathbf{curl} \, \phi_j)_{L_2(\Omega_1)} - \langle \mathbf{N}(\gamma_{\!\mathcal{D}} \phi_i), \gamma_{\!\mathcal{D}} \phi_j \rangle_{\tau}, \\ (\mathbf{M})_{\mathit{ij}} &:= \omega(\sigma \phi_i, \phi_j)_{L_2(\Omega_1)}, \\ (\mathbf{A})_{\mathit{ij}} &:= \left\langle \psi_i, \mathbf{A}(\psi_j) \right\rangle_{\tau}, \\ (\mathbf{B})_{\mathit{ij}} &:= \left\langle \psi_i, (\mathbf{C} - \mathbf{Id})(\gamma_{\!\mathcal{D}} \phi_j) \right\rangle_{\tau}. \end{split}$$

The entries of the right-hand side vector are given by the formulas  $(\mathbf{f}^c)_i := 88$   $(\mathbf{f}^c, \varphi_i)_{\mathbf{L}_2(\Omega_1)}$  and  $(\mathbf{f}^s)_i := (\mathbf{f}^s, \varphi_i)_{\mathbf{L}_2(\Omega_1)}$ . The resulting system  $\mathscr{A} \mathbf{x} = \mathbf{f}$  of the coupled 89 finite and boundary element equations has now the following structure: 90

$$\begin{pmatrix} \mathbf{M} & 0 & \mathbf{K} & \mathbf{B}^{T} \\ 0 & 0 & \mathbf{B} & -\mathbf{A} \\ \mathbf{K} & \mathbf{B}^{T} & -\mathbf{M} & 0 \\ \mathbf{B} & -\mathbf{A} & 0 & 0 \end{pmatrix} \begin{pmatrix} \mathbf{u}^{s} \\ \lambda^{s} \\ \mathbf{u}^{c} \\ \lambda^{c} \end{pmatrix} = \begin{pmatrix} \mathbf{f}^{c} \\ 0 \\ \mathbf{f}^{s} \\ 0 \end{pmatrix}.$$
(5)

In fact, the system matrix  $\mathscr{A}$  is symmetric and indefinite and obtains a double 91 saddle-point structure. Since  $\mathscr{A}$  is symmetric, the system can be solved by a Min-92 Res method, see, e.g., [18]. Anyhow, the convergence rate of any iterative method 93 deteriorates with respect to the meshsize h and the "bad" parameters  $\omega$ ,  $\nu$  and  $\sigma$ , 94 if applied to the unpreconditioned system (5). Therefore, preconditioning is a challenging topic. 96

### 3 A Parameter-Robust Preconditioning Technique

In this section, we investigate a preconditioning technique for double saddle-point 98 equations with the block-structure (5). Due to the symmetry and coercivity properties 99 of the underlying operators, the blocks fulfill the following properties:  $\mathbf{K} = \mathbf{K}^T \geq 0$ , 100  $\mathbf{M} = \mathbf{M}^T > 0$  and  $\mathbf{A} = \mathbf{A}^T > 0$ .

In [19] a parameter-robust block-diagonal preconditioner for the distributed optimal control of the Stokes equations is constructed. The structural similarities to that preconditioner gives us a hint how to choose the block-diagonal preconditioner in our case. Therefore, we propose the following preconditioner

$$\mathscr{C} = \operatorname{diag} \left( \mathscr{I}_{FEM}, \mathscr{I}_{BEM}, \mathscr{I}_{FEM}, \mathscr{I}_{BEM} \right),$$

where the diagonal blocks are given by  $\mathscr{I}_{FEM} = \mathbf{M} + \mathbf{K}$  and  $\mathscr{I}_{BEM} = \mathbf{A} + \mathbf{B} \mathscr{I}_{FEM}^{-1} \mathbf{B}^T$ . 106 Being aware that  $\mathscr{I}_{FEM}$  and  $\mathscr{I}_{BEM}$  are symmetric and positive definite, we conclude 107 that  $\mathscr{C}$  is also symmetric and positive definite. Therefore,  $\mathscr{C}$  induces the energy norm 108  $\|\mathbf{u}\|_{\mathscr{C}} = \sqrt{\mathbf{u}^T \mathscr{C} \mathbf{u}}$ . Using this special norm, we can apply the Theorem of Babuška-109 Aziz [3] to the variational problem: 110

Find 
$$\mathbf{x} \in \mathbb{R}^N$$
:  $\mathbf{w}^T \mathcal{A} \mathbf{x} = \mathbf{w}^T \mathbf{f}$ ,  $\forall \mathbf{w} \in \mathbb{R}^N$ .

The main result is now summarized in the following lemma.

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**Lemma 1.** The matrix  $\mathscr{A}$  satisfies the following norm equivalence inequalities:

$$\frac{1}{\sqrt{7}} \|\mathbf{x}\|_{\mathscr{C}} \leq \sup_{\mathbf{w} \neq 0} \frac{\mathbf{w}^T \mathscr{A} \mathbf{x}}{\|\mathbf{w}\|_{\mathscr{C}}} \leq 2 \|\mathbf{x}\|_{\mathscr{C}} \quad \forall \mathbf{x} \in \mathbb{R}^N.$$

*Proof.* Throughout the proof, we use the following notation:  $\mathbf{x} = (\mathbf{x_1}, \mathbf{x_2}, \mathbf{x_3}, \mathbf{x_4})^T$  113 and  $\mathbf{y} = (\mathbf{y_1}, \mathbf{y_2}, \mathbf{y_3}, \mathbf{y_4})^T$ . The upper bound follows by reapplication of Cauchy's inequality several time. The expressions corresponding to the Schur complement can 115 be derived in the following way: 116

$$\mathbf{y_1}^T \mathbf{B}^T \mathbf{x_4} = \mathbf{y_1} \mathcal{I}_{FEM}^{1/2} \mathcal{I}_{FEM}^{-1/2} \mathbf{B}^T \mathbf{x_4} \leq \|\mathcal{I}_{FEM}^{1/2} \mathbf{y_1}\|_{l_2} \|\mathcal{I}_{FEM}^{-1/2} \mathbf{B}^T \mathbf{x_4}\|_{l_2}.$$

Therefore, we end up with an upper bound with constant 2. 117

In order to compute the lower bound, we use a linear combination of special test 118 vectors. For the choice  $\mathbf{w_1} = (\mathbf{x_1}, \mathbf{x_2}, -\mathbf{x_3}, -\mathbf{x_4})^T$ , we obtain 119

$$\mathbf{w_1}^T \mathcal{A} \mathbf{x} = \mathbf{x_1}^T \mathbf{M} \mathbf{x_1} + \mathbf{x_3}^T \mathbf{M} \mathbf{x_3};$$
  
 $\mathbf{x_2}^T \mathbf{x_3}^T \mathbf{x_4} + \mathbf{x_3}^T \mathbf{x_4} + \mathbf{x_3}^T \mathbf{x_4} + \mathbf{x_4}^T \mathbf{x_4} + \mathbf{x_5}^T \mathbf{x_4} + \mathbf{x_5}^T \mathbf{x_5} + \mathbf{x_$ 

for  $\mathbf{w_2} = (\mathbf{x_3}, -\mathbf{x_4}, \mathbf{x_1}, -\mathbf{x_2})^T$ , we get

$$\mathbf{w_2}^T \mathcal{A} \mathbf{x} = \mathbf{x_1}^T \mathbf{K} \mathbf{x_1} + \mathbf{x_3}^T \mathbf{K} \mathbf{x_3} + \mathbf{x_2}^T \mathbf{A} \mathbf{x_2} + \mathbf{x_4}^T \mathbf{A} \mathbf{x_4};$$

for 
$$\mathbf{w_3} = ((\mathbf{x_4}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0}, (\mathbf{x_2}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0})^T$$
, we have

$$\begin{aligned} \mathbf{w_3}^T \mathscr{A} \mathbf{x} &= \mathbf{x_4}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_4} + \mathbf{x_2}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_2} \\ &+ \mathbf{x_4}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1} + \mathbf{x_4}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_3} \\ &+ \mathbf{x_2}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_1} - \mathbf{x_2}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3}; \end{aligned}$$

for 
$$\mathbf{w_4} = (-(\mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0}, -(\mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0})^T$$
, we get

for 
$$\mathbf{w_4} = (-(\mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0}, -(\mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0})^T$$
, we get 
$$\mathbf{w_4}^T \mathscr{A} \mathbf{x} = -\mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1} - \mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_3} - \mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_4} - \mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_1} - \mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_2} + \mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3};$$

and, finally, for the choice  $\mathbf{w_5} = (-(\mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0}, (\mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1})^T, \mathbf{0})^T$ , 123 we obtain

$$\begin{aligned} \mathbf{w_5}^T \mathscr{A} \mathbf{x} &= -\mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1} - \mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_3} \\ &- \mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_4} + \mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_1} \\ &+ \mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_2} - \mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3}. \end{aligned}$$

Therefore, we end up with the following expression

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$$\begin{split} &(\mathbf{w_1} + \mathbf{w_2} + \mathbf{w_3} + \mathbf{w_4} + \mathbf{w_5})^T \mathscr{A} \mathbf{x} = \mathbf{x_1}^T \mathbf{M} \mathbf{x_1} + \mathbf{x_3}^T \mathbf{M} \mathbf{x_3} \\ &+ \mathbf{x_1}^T \mathbf{K} \mathbf{x_1} + \mathbf{x_3}^T \mathbf{K} \mathbf{x_3} + \mathbf{x_2}^T \mathbf{A} \mathbf{x_2} + \mathbf{x_4}^T \mathbf{A} \mathbf{x_4} \\ &+ \mathbf{x_4}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_4} + \mathbf{x_2}^T \mathbf{B} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{B}^T \mathbf{x_2} \\ &- \mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_3} - \mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_1} \\ &- \mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3} - \mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1} \\ &- 2 \mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1} + 2 \mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3}. \end{split}$$

For estimating the non-symmetric terms, we use the following result:

$$\begin{aligned} -2\mathbf{x_3}^T\mathbf{K}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{M}\mathbf{x_1} &\geq -2\|(\mathbf{K}+\mathbf{M})^{-1/2}\mathbf{K}\mathbf{x_3}\|_{l_2}\|(\mathbf{K}+\mathbf{M})^{-1/2}\mathbf{M}\mathbf{x_1}\|_{l_2} \\ &\geq -\|(\mathbf{K}+\mathbf{M})^{-1/2}\mathbf{K}\mathbf{x_3}\|_{l_2}^2 - \|(\mathbf{K}+\mathbf{M})^{-1/2}\mathbf{M}\mathbf{x_1}\|_{l_2}^2 \\ &= -\mathbf{x_3}^T\mathbf{K}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{K}\mathbf{x_3} - \mathbf{x_1}^T\mathbf{M}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{M}\mathbf{x_1}. \end{aligned}$$

Analogously, we obtain

$$2\mathbf{x_1}^T\mathbf{K}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{M}\mathbf{x_3} \ge -\mathbf{x_1}^T\mathbf{K}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{K}\mathbf{x_1} - \mathbf{x_3}^T\mathbf{M}(\mathbf{K}+\mathbf{M})^{-1}\mathbf{M}\mathbf{x_3}.$$

Hence, putting all terms together, we have

$$\begin{aligned} &(\mathbf{w_1} + \mathbf{w_2} + \mathbf{w_3} + \mathbf{w_4} + \mathbf{w_5})^T \mathscr{A} \mathbf{x} = \mathbf{x}^T \mathscr{C} \mathbf{x} \\ &- 2\mathbf{x_3}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_3} - 2\mathbf{x_1}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_1} \\ &- 2\mathbf{x_3}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_3} - 2\mathbf{x_1}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_1}. \end{aligned}$$

In order to get rid of the four remaining terms, we use, for i = 1,3,

$$\mathbf{x_i}^T \mathbf{K} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{K} \mathbf{x_i} \le \mathbf{x_i}^T \mathbf{K} \mathbf{x_i}$$
 and  $\mathbf{x_i}^T \mathbf{M} (\mathbf{K} + \mathbf{M})^{-1} \mathbf{M} \mathbf{x_i} \le \mathbf{x_i}^T \mathbf{M} \mathbf{x_i}$ .

Hence by adding w<sub>1</sub> and w<sub>2</sub> twice more, we end up with the desired result

$$\underbrace{(3\mathbf{w}_1 + 3\mathbf{w}_2 + \mathbf{w}_3 + \mathbf{w}_4 + \mathbf{w}_5)^T}_{:=\mathbf{w}^T} \mathscr{A}\mathbf{x} \ge \mathbf{x}^T \mathscr{C}\mathbf{x} + \mathbf{x}_2^T \mathbf{A}\mathbf{x}_2 + \mathbf{x}_4^T \mathbf{A}\mathbf{x}_4 \ge \mathbf{x}^T \mathscr{C}\mathbf{x}.$$

The next step is to compute (and estimate) the  $\mathscr C$  norm of the special test vector. 131 Straightforward estimations yield

$$\|\mathbf{w}\|_{\mathscr{C}}^2 = \|3\mathbf{w}_1 + 3\mathbf{w}_2 + \mathbf{w}_3 + \mathbf{w}_4 + \mathbf{w}_5\|_{\mathscr{C}}^2 \le 7\|\mathbf{x}\|_{\mathscr{C}}^2.$$

This completes the proof.

Now, from Lemma 1, we obtain that the condition number of the preconditioned 134 system can be estimated by the constant  $c=2\sqrt{7}$  that is obviously independent of 135 the meshsize h and all involved parameters  $\omega$ ,  $\nu$  and  $\sigma$ , i.e. 136

$$\kappa_{\mathscr{C}}(\mathscr{C}^{-1}\mathscr{A}) := \|\mathscr{C}^{-1}\mathscr{A}\|_{\mathscr{C}}\|\mathscr{A}^{-1}\mathscr{C}\|_{\mathscr{C}} \le 2\sqrt{7}. \tag{6}$$

The condition number defines the convergence behaviour of the MinRes method 137 applied to the preconditioned system (see e.g. [9]), as stated in the following theorem: 138

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**Theorem 1 (Robust solver).** The MinRes method applied to the preconditioned system  $\mathcal{C}^{-1}A\mathbf{u} = \mathcal{C}^{-1}\mathbf{f}$  converges. At the 2m-th iteration, the preconditioned residual 141  $\mathbf{r}^{\mathbf{m}} = \mathscr{C}^{-1}\mathbf{f} - \mathscr{C}^{-1}\mathscr{A}\mathbf{u}^{\mathbf{m}}$  is bounded as

$$\left\|\mathbf{r^{2m}}\right\|_{\mathscr{C}} \le \frac{2q^m}{1+q^{2m}} \left\|\mathbf{r^0}\right\|_{\mathscr{C}}, \quad where \quad q = \frac{2\sqrt{7}-1}{2\sqrt{7}+1}. \tag{7}$$

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#### 4 Conclusion, Outlook and Acknowledgments

The method developed in this work shows great potential for solving time-harmonic 144 eddy current problems in an unbounded domain in a robust way. The solution of a 145 fully coupled 4 × 4 block-system can be reduced to the solution of a block-diagonal 146 matrix, where each block corresponds to standard problems. We mention, that by 147 analogous procedure, we can state another robust block-diagonal preconditioner  $\mathscr{C}=148$ diag  $(\tilde{\mathscr{I}}_{FEM}, \tilde{\mathscr{I}}_{BEM}, \tilde{\mathscr{I}}_{FEM}, \tilde{\mathscr{I}}_{BEM})$ , with  $\tilde{\mathscr{I}}_{FEM} = \mathbf{M} + \mathbf{K} + \mathbf{B}^T \tilde{\mathscr{I}}_{BEM}^{-1} \mathbf{B}$  and  $\tilde{\mathscr{I}}_{BEM} = \mathbf{M} + \mathbf{K} + \mathbf{B}^T \tilde{\mathscr{I}}_{BEM}^{-1} \mathbf{B}$ A, leading to a condition number bound of 4, see e.g. [15].

Of course this block-diagonal preconditioner is only a theoretical one, since the 151 exact solution of the diagonal blocks corresponding to a standard FEM discretized 152 stationary problem and the Schur-complement of a standard FEM-BEM discretized 153 stationary problem are still prohibitively expensive. Nevertheless, as for the FEM 154 discretized version in [13], this theoretical preconditioner allows us replace the solution of a time-dependent problem by the solution of a sequence of time-independent 156 problems in a robust way, i.e. independent of the space and time discretization pa- 157 rameters h and  $\omega$  and all additional "bad" parameters. Therefore, the issue of finding 158 robust solvers for the fully coupled time-harmonic system matrix A can be reduced 159 to finding robust solvers for the blocks  $\mathscr{I}_{FEM}$  and  $\mathscr{I}_{BEM}$ , or  $\mathscr{I}_{FEM}$  and  $\mathscr{I}_{BEM}$ . By 160 replacing these diagonal blocks by standard preconditioners, it is straight-forward 161 to derive mesh-independent convergence rates, see, e.g., [8]. Unfortunately, the con- 162 struction of fully robust preconditioners for the diagonal blocks is not straight for- 163 ward and has to be studied. Candidates are H matrix, multigrid multigrid and do- 164 main decomposition preconditioners, see, e.g. [2, 6] and [12], respectively.

The preconditioned MinRes solver presented in this paper can also be generalized 166 to eddy current optimal control problems studied in [14] for the pure FEM case in 167 bounded domains.

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# **Domain Decomposition Methods for Auxiliary Linear Problems of an Elliptic Variational Inequality**

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Summary. Elliptic variational inequalities with multiple bodies are considered. It is assumed 7 that an active set method is used to handle the nonlinearity of the inequality constraint, which 8 results in auxiliary linear problems. We describe two domain decomposition methods for solving such linear problems, namely, the FETI-FETI (finite element tearing and interconnecting) 10 and hybrid methods, which are combinations of already existing domain decomposition methods. 12

Estimates of the condition numbers of both methods are provided. The FETI-FETI method 13 has a condition number which depends linearly on the number of subdomains across each body 14 and polylogarithmically on the number of element across each subdomain. The hybrid method 15 is a scalable alternative to the FETI-FETI method, and has a condition number with two polylogarithmic factors depending on the number of elements across each subdomain and across 17 each body. We present numerical results confirming these theoretical findings.

1 Introduction 19

Consider the following inequality constrained minimization problem,

$$\min \quad \sum_{i=1}^{N} \left( \frac{1}{2} \int_{\Omega_{i}} \rho(x) |\nabla u^{i}(x)|^{2} dx - \int_{\Omega_{i}} f(x) u^{i}(x) dx \right),$$
where  $u^{i} \in H^{1}(\Omega_{i}), u^{i} = 0 \text{ on } \Gamma_{u}^{i}, i = 1, \dots, N,$ 

$$u^{i} - u^{j} \leq 0 \text{ on } \partial \Omega_{i} \cap \partial \Omega_{j}, \forall i < j,$$
(1)

with variable coefficients and multiple bodies  $\Omega_i \subset \mathbb{R}^2$  with their boundaries and 21 the Dirichlet boundaries denoted by  $\partial \Omega_i$  and  $\Gamma_u^i$ , respectively, for  $i=1,\cdots,N$ . The 22 bodies are decomposed into subdomains,

$$\Omega_i = igcup_{j=1}^{N_i} \Omega_{i,j}, \quad i=1,\cdots,N.$$
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Here, bodies mean separate physical entities; for instance, two rubber balls in contact 25 with each other are considered two bodies. Subdomains, on the other hand, is artifi- 26 cially introduced for convenience; a rubber ball can consist of as many subdomains 27

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as the modeler wants. We assume that the coefficient  $\rho$  varies moderately within 28 each body,  $\Omega_i$ ,  $i=1,\cdots,N$ . The diameters of  $\Omega_i$  and  $\Omega_{i,j}$  are denoted by  $H_i$  and  $H_{i,j}$ , 29 respectively. The smallest diameters of any element in  $\Omega_i$  and  $\Omega_{i,j}$  are denoted by 30  $h_i$  and  $h_{i,j}$ , respectively. Also,  $H_b:=\max_i H_i$ ,  $H_s:=\max_i H_{i,j}$ ,  $\frac{H_b}{h}:=\max_i \frac{H_i}{h_i}$ ,  $\frac{H_s}{h}:=31$   $\max_{i,j} \frac{H_{i,j}}{h_{i,j}}$ . We introduce the following:

$$\Gamma_{gl}:=igcup_{i
eq j}\partial\Omega_i\cap\partial\Omega_j, ext{ potential contact surface between bodies,,}$$
 
$$\Gamma_{loc}^{(i)}:=igcup_{i
eq k}(\partial\Omega_{i,j}\cap\partial\Omega_{i,k}), ext{interface between subdomains, } i=1,\cdots,N. \tag{2}$$

Here, the subscripts gl and loc stand for global and local, respectively, referring to 33

nature of the interfaces. For each body,  $\Omega_i$ ,  $i = 1, \dots, N$ , two kinds of finite element spaces are introduced:  $\widehat{W}^{(i)}$  is a standard finite element space of continuous, 35 piecewise linear functions and, as such, is continuous across  $\Gamma_{loc}^{(i)}$ ;  $\widetilde{W}^{(i)}$  is a more 36 general space, consisting of finite element functions required to be continuous only 37 at the *primal* nodes (i.e., the vertex nodes of  $\Gamma_{loc}^{(i)}$  in this two-dimensional case; more 38 sophisticated continuity couplings, i.e., primal constraints, are required in  $\widetilde{W}^{(i)}$  for 39 three-dimensional problems; see [9, 10]), as in the FETI-DP (dual-primal FETI) 40 method. The trace spaces of  $\widetilde{W}^{(i)}$  and  $\widehat{W}^{(i)}$  on  $\Gamma_{loc}^{(i)} \cup (\partial \Omega_i \cap \Gamma_{gl})$  are denoted by  $\widetilde{V}^{(i)}$ and  $\widehat{V}^{(i)}$ , respectively. The trace space of  $\widehat{W}^{(i)}$  on  $\partial \Omega_i \cap \Gamma_{gl}$  is denoted by  $V_{Ol}^{(i)}$ , where 42 OL stands for "one level." The Schur complements of the stiffness matrices for  $\widetilde{W}^{(i)}$  43 and  $\widehat{W}^{(i)}$ , obtained by eliminating unknowns corresponding to the *subdomain inte-* 44 riors, that is, those not associated with  $\Gamma_{loc}^{(i)} \cup (\partial \Omega_i \cap \Gamma_{gl})$ , are denoted by  $\widetilde{S}_{\Gamma}^{(i)}$  and 45  $\widehat{S}_{\Gamma}^{(i)}$ , respectively. The Schur complement  $S_{OL}^{(i)}$  of the stiffness matrix for  $\widehat{W}^{(i)}$ , on the 46 other hand, is obtained by eliminating unknowns corresponding to the body interior, 47 i.e., those *not* associated with  $\partial\Omega_i\cap\Gamma_{gl}$ . Therefore  $\widetilde{S}_{\Gamma}^{(i)},\widehat{S}_{\Gamma}^{(i)}$ , and  $S_{OL}^{(i)}$  can be viewed 48 as operators on  $\widetilde{V}^{(i)},\widehat{V}^{(i)},$  and  $V_{OL}^{(i)},$  respectively. We note that applying  $S_{OL}^{(i)}$  requires 49 solving a Dirichlet problem on  $\Omega_i$ . Let  $\widetilde{V} := \prod_{i=1}^{N} \widetilde{V}^{(i)}, \widehat{V} := \prod_{i=1}^{N} \widehat{V}^{(i)}, V_{OL} := \prod_{i=1}^{N} V_{OL}^{(i)}, \widetilde{S} = \operatorname{diag}_{i=1}^{N} \widetilde{S}_{\Gamma}^{(i)}, \widehat{S} = \operatorname{diag}_{i=1}^{N} \widehat{S}_{\Gamma}^{(i)},$  51 and  $S_{OL} := \operatorname{diag}_{i=1}^{N} S_{OL}^{(i)}$ . We also introduce matrices  $\widetilde{B}, \widehat{B}$ , and  $B_{OL}$ , with elements of  $\{0,-1,1\}$ :  $\widetilde{Bu} \Leftrightarrow u \in \widetilde{V}$  is continuous across  $\Gamma_{loc}^{(i)}, \forall i$ , as well as  $\Gamma_{gl}$ ;  $\widehat{Bv} \Leftrightarrow v \in \mathfrak{ss}$  $\widehat{V}$  is continuous across  $\Gamma_{gl}$ ;  $B_{OL}w \Leftrightarrow w \in V_{OL}$  is continuous across  $\Gamma_{gl}$ .

### 2 Algorithms

With the matrices defined in Sect. 1, we can consider the following algorithm for 56 solving (1): 57

Algorithm: Active set method + Krylov subspace method

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1. Initialize  $u^0$ . Set k = 0. Set  $\mathscr{A}_k$ , a subset of the index set  $\{1, \dots, \#(\text{rows}(\widetilde{B}))\}$ (resp.  $\#(rows(\widehat{B}))$ ), according to the active set method of choice. 60

2. Solve 61

$$\min_{u \in \widetilde{V}} \frac{1}{2} u^T \widetilde{S} u - \widetilde{g}^T u, \quad \text{with} \quad Z^k \widetilde{B} u = 0$$
 (3)

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$$\min_{u \in \widehat{V}} \frac{1}{2} u^T \widetilde{S} u - \widetilde{g}^T u, \quad \text{with} \quad Z^k \widetilde{B} u = 0$$

$$\left( \text{resp.} \quad \min_{u \in \widehat{V}} \frac{1}{2} u^T \widehat{S} u - \widehat{g}^T u, \quad \text{with} \quad \widehat{Z}^k \widehat{B} u = 0 \right)$$
(4)

approximately to a given precision, using a Krylov subspace method. Set  $u^{k+1}$ to the resulting approximate solution. Find  $\mathcal{A}_{k+1}$  accordingly. 63

3. Set k = k + 1. Stop if  $\mathcal{A}_{k-1} = \mathcal{A}_k$ ; return to Step 2 otherwise.

Note that the linear problem in the kth iteration of the active set method is formulated as a minimization problem in terms of the interface variables in  $\widetilde{V}$  or  $\widehat{V}$ . Here, 66  $\widetilde{g} \in \widetilde{V}$  and  $\widehat{g} \in \widehat{V}$  are appropriate load vectors. The square, diagonal matrix  $Z^k$ , with 67 all elements equal to 0 or 1, is chosen such that  $Z^k\widetilde{B} = \widetilde{B}_{\mathcal{A}_k}$ , where  $\widetilde{B}_{\mathcal{A}_k}$  is obtained 68 by replacing the *i*th row of  $\widetilde{B}$  with zeros for  $\forall i \notin \mathscr{A}_k$ . The matrix  $\widehat{Z}^k$  is defined analogously. The minimization problems (3) and (4) are equivalent to the following saddle 70 point problems,

$$\begin{bmatrix} \widetilde{S} & (Z^k \widetilde{B})^T \\ Z^k \widetilde{B} & 0 \end{bmatrix} \begin{bmatrix} u \\ \lambda \end{bmatrix} = \begin{bmatrix} \widetilde{g} \\ 0 \end{bmatrix}, \tag{5}$$

and

$$\begin{bmatrix} \widehat{S} & (\widehat{Z}^k \widehat{B})^T \\ \widehat{Z}^k \widehat{B} & 0 \end{bmatrix} \begin{bmatrix} u \\ \lambda \end{bmatrix} = \begin{bmatrix} \widehat{g} \\ 0 \end{bmatrix}, \tag{6}$$

respectively. We now consider the preconditioning of (5) and (6).

The FETI-FETI method is a combination of the one-level FETI method with a 74 Dirichlet preconditioner [4] and the FETI-DP method [5], and was used in [1, 2] 75 to solve frictionless contact problems. For (6), it is natural to follow the approach in 76 the one-level and FETI-DP methods and form a Schur complement equation 77

$$\underbrace{Z^{k}\widetilde{B}\widetilde{S}^{\dagger}\widetilde{B}^{T}Z^{k}}_{\cdot -F}\lambda = Z^{k}\widetilde{B}\widetilde{S}^{\dagger}\widetilde{g} + Z^{k}\widetilde{B}R\alpha, \tag{7}$$

where  $\widetilde{S}^{\dagger}$  is a pseudoinverse of  $\widetilde{S}$ , range $(R) = \text{null}(\widetilde{S})$ , and the vector  $\alpha$  is to be 78 determined. We solve (7) with the preconditioned conjugate gradient (PCG) method, 79 using the following preconditioner:

$$P_F^{-1} := Z^k \widetilde{B}_D \widetilde{S} \widetilde{B}_D^T Z^k. \tag{8}$$

If  $\widetilde{S}$  is singular, then the PCG method needs to be confined to the following subspace: 81

$$V^{k} := \{ \lambda : Z^{k} \widetilde{B} \lambda \in \operatorname{range}(\widetilde{S}) \}. \tag{9}$$

Most of the computational work in each iteration of the PCG method goes into the ap- 82 plications of  $\widetilde{S}^{\dagger}$  and  $\widetilde{S}$ , in the applications of F and  $P_F^{-1}$ , respectively. The application 83 of  $\widetilde{S}$  involves solving a Dirichlet problem on each subdomain,  $\Omega_{i,j}, i=1,\cdots,N, j=84$  $1, \dots, N_i$ . The application of  $\widetilde{S}^{\dagger}$  involves solving a Dirichlet problem in each subdomain, with the Dirichlet boundary condition imposed only at subdomain vertices, 86 plus solving a coarse problem on each body, associated with the set of vertices of 87  $\Gamma_{loc}^{(i)}, i = 1, \dots, N$ ; for details, see, e.g., [13], [14, Chap. 6].

The hybrid method is a combination of the one-level FETI method with a Dirich- 90 let preconditioner and the BDDC (balancing domain decomposition by constraints) 91 method [3]. For (6), forming a Schur complement equation similar to (7) is much 92 more expensive because of the dense structure of S. Hence we keep the saddle point 93 formulation (6) as is and solve it with the preconditioned conjugate residual (PCR) 94 method. As in the FETI-FETI method, the PCR method needs to be confined to the 95 following subspace:

$$\widehat{V}^k := \{\lambda : \widehat{Z}^k \widehat{B} \lambda \in \operatorname{range}(\widehat{S})\}.$$

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Letting  $P^k$  denote an orthogonal projection onto  $V^k$ , we rewrite (6) as

$$\underbrace{\begin{bmatrix} \widehat{S} & (P^k \widehat{Z}^k \widehat{B})^T \\ P^k \widehat{Z}^k \widehat{B} & 0 \end{bmatrix}}_{:=\mathscr{A}} \begin{bmatrix} u \\ \mu \end{bmatrix} = \begin{bmatrix} \widehat{g} - \widehat{B}^T \lambda_0 \\ 0 \end{bmatrix}, \tag{10}$$

with  $\lambda_0$  satisfying  $(\widehat{Z}^k\widehat{B}^T)\lambda_0 \in \operatorname{range}(\widehat{S})$ . For details on how to recover a solution 99 of (6) from a solution of (10), see [8]. Letting  $P_R$  denote an orthogonal projection 100 onto range  $(\hat{S})$ , we introduce the preconditioner  $\mathcal{B}$ , where 101

$$\mathscr{B}^{-1} = \begin{bmatrix} P_R M_{BDDC}^{-1} P_R & 0\\ 0 & P^k M_D^{-1} P^k \end{bmatrix}. \tag{11}$$

Here,  $M_{BDDC}$  is a block diagonal matrix consisting of the BDDC preconditioners [3] 102 for the bodies: 103

$$M_{BDDC}^{-1} = \operatorname{diag}_{i=1}^{N} M_{BDDC}^{(i)^{-1}} = \operatorname{diag}_{i=1}^{N} \widetilde{R}_{D,\Gamma}^{(i)^{T}} \widetilde{S}_{\Gamma}^{(i)^{\dagger}} \widetilde{R}_{D,\Gamma}^{(i)},$$

where  $\widetilde{R}_{D,\Gamma}^{(i)T}$ ,  $i=1,\cdots,N$ , is a scaled restriction from  $\widetilde{V}^{(i)}$  to  $\widehat{V}^{(i)}$ , with the scaling factors determined by the material coefficients; similarly, B<sub>OL,D</sub> is a scaled version 106 of  $B_{OL}$ . For details on the definition of these matrices, see, for instance, [11, 13]. 107 Then  $M_D$  can be viewed as a Dirichlet preconditioner of the one-level FETI method, 108 obtained by viewing each body,  $\Omega_i$ , as a subdomain:

$$M_D^{-1} = \widehat{Z}^k B_{OL,D} S_{OL} B_{OL,D}^T \widehat{Z}^{k^T}.$$

Most of the computational work in each iteration of the PCR method goes into the 111 application of  $\widehat{S}$ , in the application of  $\mathscr{A}$ , and the application of  $\widetilde{S}_{\Gamma}^{(i)^{\dagger}}$ ,  $i=1,\cdots,N$ and  $S_{OL}$ , in the application of  $\mathscr{B}^{-1}$ . The application of  $\widehat{S}$  requires solving a Dirichlet problem on each subdomain,  $\Omega_{i,j}$ ,  $i=1,\cdots,N, j=1,\cdots,N_i$ . The application of  $\widetilde{S}_{\Gamma}^{(i)^{\dagger}}$ ,  $i=1,\cdots,N$ , which is carried out in the FETI-FETI method as well, requires solving a Dirichlet problem on  $\Omega_{i,j}$ ,  $j=1,\cdots,N_i$  with the Dirichlet boundary condition imposed only at the vertices, plus solving a coarse problem on  $\Omega_i$  associated with the vertices of  $\Gamma_{loc}^{(i)}$ . The application of  $S_{OL}$ , however, requires solving a Dirichlet problem on each body, which is expensive; therefore in practice such a Dirichlet problem needs only to be solved inexactly, for instance with a Krylov subspace method. A preconditioner for solving such a Dirichlet problem is proposed and tested in [11].

3 Theory 123

We now present condition number estimates for the FETI-FETI and hybrid methods. 124 Because of space limitations, details and proofs are given elsewhere; see [11, 12]. 125

**Theorem 1.** Let  $F, P_F$ , and  $V^k$  be defined as in (7) and (9), respectively. For any 126  $\lambda \in V^k$ , we have

$$\langle P_F \lambda, \lambda \rangle \le \langle F \lambda, \lambda \rangle \le C(H_b/H_s)(1 + \log(H_s/h))^2 \langle P_F \lambda, \lambda \rangle,$$
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where C > 0 is a constant independent of the sizes of the bodies, subdomains, and 129 elements.

Convergence of the PCR method for the hybrid method is determined by

$$\mathcal{K}(\mathcal{B}^{-1}\mathcal{A}) := \frac{\mu_{max}}{\mu_{min}} = \frac{\max\{|\lambda| : \lambda \in \sigma(\mathcal{B}^{-1}\mathcal{A})\}}{\min\{|\lambda| : \lambda \in \sigma(\mathcal{B}^{-1}\mathcal{A})\}},\tag{12}$$

where  $\sigma(\mathscr{B}^{-1}\mathscr{A})$  is the spectrum of  $\mathscr{B}^{-1}\mathscr{A}$  on range $(P_R) \times \widehat{V}^k$ .

**Theorem 2.** Let  $\mathcal{B}^{-1}$ ,  $\mathcal{A}$ , and  $\mathcal{K}(\mathcal{B}^{-1}\mathcal{A})$  be defined as in (11)–(12), respectively. 133 We then have the following bound:

$$\mathscr{K}(\mathscr{B}^{-1}\mathscr{A}) \le C(1 + \log(H_b/h))^2 (1 + \log(H_s/h))^2,$$
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where C > 0 is a constant independent of the sizes of the bodies, subdomains, and 136 elements.

## 4 Numerical Results: Auxiliary Linear Problems

We solve the following equality-constrained minimization problem:

$$\min \sum_{i=1}^{N_b \times N_b} \left( \frac{1}{2} \int_{\Omega_i} |\nabla u^i|^2 dx - \int_{\Omega_i} f u^i dx \right),$$
with equality constraints to be specified, (13)

			F	ETI-	-FETI		Hyl	orid
			I		II		I	II
$1/H_b$	$H_b/H_s$	$H_s/h$	cond	iter	cond	iter	iter	iter
2	fixed	fixed	2.89	7	2.31	7	10	10
4	at 2	at 2	4.41	12	2.85	10	11	8
6			4.51	13	2.91	10	11	9
8			4.55	14	2.93	10	11	8
10			4.56	14	2.94	10	11	8
12			4.57	13	2.95	10	11	7
14			4.58	14	2.96	10	11	7
16			4.58	14	2.96	10	11	7
fixed	4	fixed	7.68	10	5.02	9	10	10
at 2	6	at 2	12.70	12	7.46	10	10	10
	8		17.80	13	8.12	10	10	10
	10		22.93	15	10.96	11	10	8
	12		28.08	16	13.43	12	10	8
	14		33.25	17	14.01	12	9	8
	16		38.41	17	16.90	12	8	7
fixed	fixed	4	4.71	9	4.73	9	12	11
at 2	at 2	6	5.90	10	6.37	10	13	13
		8	6.90	10	7.08	10	13	13
		10	7.79	11	8.27	11	14	14
		12	8.55	11	9.25	11	14	14
		14	9.23	12	9.71	12	14	14
	V-	16	9.83	12	10.52	12	14	14

**Table 1.** Results of FETI-FETI and hybrid.

where  $\Omega_i \subset \mathbb{R}^2, i=1,\cdots,N_b \times N_b$  are square bodies with side length  $H_b:=1/N_b$ , 140 which collectively form the domain  $\bar{\Omega}=\bigcup_{i=1}^{N_b\times N_b}\bar{\Omega}_i=[0,1]\times[0,1]$ . We require  $u^i\in 1$ 141

 $H^1(\Omega_i), u^i|_{\partial\Omega_i\cap\partial\Omega} = 0$ . Each  $\Omega_i$  is decomposed into  $N_s \times N_s$  square subdomains, 142 each of which is discretized by square bilinear elements of side length h. Also,  $\Gamma := 143$  $\bigcup_{i\neq j}\partial\Omega_i\cap\partial\Omega_i$  denotes the interface between the bodies.

We supplement (13) with two different equality constraints, associated with different contact areas between the bodies. In the first problem, the entire  $\Gamma$  is considered as the contact area, that is, we require the continuity of the displacement 147 vector across the entire  $\Gamma$ . This case has already been considered by Klawonn and 148 Rheinbach [6] and Klawonn and Rheinbach [7]. In the second problem, continuity 149 is imposed only on the middle third of the faces between the bodies. We solve these 150 problems with both the FETI-FETI and hybrid methods. The PCG and PCR iter- 151 ations are stopped when the norm of the residual has been reduced by a factor of 152  $10^{-6}$ .

The results are shown in Table 1. We have three parameters to vary: the number of bodies across  $\Omega$  ( $N_b = 1/H_b$ ), the number of subdomains across each body 155

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 $(N_s = H_b/H_s)$ , and the number of elements across each subdomain  $(H_s/h)$ . We vary 156 one parameter while keeping the other two fixed. The results for the first set of experiments, with the entire  $\Gamma$  as the contact surface, are shown in column I; those for 158 the second set of experiments with a reduced contact area are shown in column II.

Note the linear dependence of the condition number on the number of subdomains across each body,  $H_b/H_s$ , for the FETI-FETI method, which confirms our 161 theoretical finding. Note also that the iteration counts of the hybrid method do not 162 increase as the number of subdomains is increased. Similar numerical results for 163 the FETI-FETI method have been obtained independently by Klawonn and Rheinbach [6] and Klawonn and Rheinbach [7].

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# **New Theoretical Coefficient Robustness Results for FETI-DP**

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1 Introduction 12

In this short note, we present new weighted Poincaré inequalities (WPIs) with 13 weighted averages that allow a robustness analysis of dual-primal finite element tear- 14 ing and interconnecting (FETI-DP) methods in certain cases where jumps of coefficients are not aligned with the subdomain partition.

Let  $\Omega$  be a bounded Lipschitz domain in  $\mathbb{R}^2$  or  $\mathbb{R}^3$ . We consider the weak form 17 of the scalar elliptic PDE 18

$$-\operatorname{div}(\alpha \nabla u) = f \quad \text{in } \Omega, \tag{1}$$

with a uniformly positive diffusion coefficient  $\alpha \in L^{\infty}(\Omega)$  that is piecewise constant with respect to a (possibly rather fine) partitioning of  $\Omega$ . The discretization by continuous and piecewise linear finite elements (FEs) on a mesh  $\mathcal{F}(\Omega)$  leads to the 21 sparse (but in general large) linear system

$$\mathbf{K}\mathbf{u} = \mathbf{f}$$
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We consider FETI-DP solvers (see [2, 4, 5]) for the fast (and parallel) solution 24 of this system, and we follow the structure described in [12, Sect. 6.4]. To this end, 25 we partition the domain  $\Omega$  into non-overlapping subdomains  $\Omega_i$ ,  $i=1,\ldots,N$  such 26 that the global mesh  $\mathscr{T}(\Omega)$  resolves the interface  $\bigcup_{i\neq j}\partial\Omega_i\cap\partial\Omega_j$ . The interface 27 itself can be divided into subdomain vertices, edges, and faces (for d = 3), cf. [12, 28] Sect. 4.21.

Without loss of generality, we assume that  $\alpha$  is constant on each element of 30  $\mathcal{T}(\Omega)$ . Crucially, we do *not* assume that  $\alpha$  is constant on each subdomain. However, 31 we need assumptions on the kind of jumps. Let  $\alpha_i$  denote the restriction of  $\alpha$  to  $\Omega_i$  32 and note that it has a well-defined trace in  $L^2(\partial \Omega_i)$ . For each subdomain edge (face) 33

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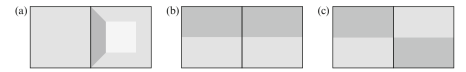


Fig. 1. Different types of coefficient jumps along an edge between two subdomains: (a) across (b) along (c) both across and along

 $\mathscr{E}$  on  $\Omega_i$ , let  $V^h(\mathscr{E})$  denote the restriction of the global FE space to  $\overline{\mathscr{E}}$  and let us define 34 the weighted average

$$\overline{v}^{\mathscr{E},\alpha_i} := \frac{\int_{\mathscr{E}} \alpha_i v}{\int_{\mathscr{E}} \alpha_i} \quad \text{for } v \in V^h(\mathscr{E}).$$
 (2)

**Assumption A1.** Whenever two  $\Omega_i$  and  $\Omega_j$  share an edge (face)  $\mathcal{E}$ , the weighted 36 averages of any function  $v \in V^h(\mathscr{E})$  coincide:  $\overline{v}^{\mathscr{E}}, \alpha_i = \overline{v}^{\mathscr{E}}, \alpha_j$ . 37

A sufficient condition for Assumption A1 is that the coefficient jumps either 38 across or along, but not both at the same time. For an illustration see Fig. 1. Our 39 assumptions rules out situations of type (c).

Following [12, Algorithm B], we define the *primal space*  $\widehat{W}_{\Pi}$  spanned by the 41 vertex nodal basis functions at subdomain vertices, the subdomain edge cut-off 42 functions and subdomain face cut-off functions (all of them extended discrete  $\alpha$ - 43 harmonically from the interface to the subdomain interiors). The dual space  $W_{\Lambda}$  44 contains FE functions that are discontinuous across the subdomain interfaces with 45 vanishing  $\alpha$ -weighted averages over the subdomain faces, edges, and vertices. We 46 formally perform a change of basis, such that we have a splitting of the degrees of 47 freedom (DOFs) into primal and dual ones, and work in the space  $\widehat{W} = \widehat{W}_{\Pi} \oplus W_{\Lambda}$ .

Let  $B: \widetilde{W} \to U$  be the usual jump operator. The FETI-DP system

$$F\lambda = B\widehat{K}^{-1}\widehat{f} \tag{3}$$

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is solved by preconditinioned conjugate gradients, where  $F := B\widehat{K}^{-1}B^{\top}$  and where 50  $\vec{K}$ ,  $\vec{f}$  denote the stiffness matrix and load vector partially assembled at the primal 51 DOFs, respectively. The overall solution is then given by

$$u = \widehat{K}^{-1}(\widehat{f} - B^{\top}\lambda).$$
 53

Next, we define a FETI-DP preconditioner that is slightly modified to allow for 54 certain coefficient jumps (cf. [3, 7]). Let i = 1, ..., N be fixed and let  $\mathcal{T}(\Omega_i)$  denote 55 the mesh restricted to subdomain  $\Omega_i$ . For each mesh node  $x^h$  on  $\overline{\Omega}_i$ , we set

$$\widehat{\alpha}_i(x^h) := \max_{T \in \mathscr{T}(\Omega_i): x^h \in \overline{T}} \alpha_{|T}. \tag{4}$$

Furthermore, if  $\mathcal{N}_{x^h}$  denotes the index set of subdomains sharing the mesh node  $x^h$ , 57 we define the weighted counting function 58

$$\delta_{i}^{\dagger}(x^{h}) := \begin{cases} \frac{\widehat{\alpha}_{i}(x^{h})}{\sum_{j \in \mathscr{N}_{x^{h}}} \widehat{\alpha}_{j}(x^{h})}, & \text{if } x^{h} \text{ lies on } \overline{\Omega}_{i}, \\ 0, & \text{otherwise.} \end{cases}$$

Using these counting functions we define the scaled jump operator  $B_D$  according 60 to [12, Sect. 6.4.1] (for details see also [9] where the same scaled jump operator 61 was used to define a one-level FETI preconditioner). The FETI-DP preconditioner is 62 finally given by

$$M^{-1} := B_D S B_D^{\top}, \tag{5}$$

where  $S = \operatorname{diag}(S_i)_{i=1}^N$  is the block-diagonal Schur complement of the block stiffness 64 matrix  $K = \operatorname{diag}(K_i)_{i=1}^N$ , eliminating the interior DOFs in each subdomain. Alternatively, one may replace B and  $B_D$  in (3), (5) by the respective operators which only 66 act on the dual DOFs, which reduces the number of redundancies in  $\lambda$ . 67

### 2 Weighted Poincaré Inequalities with Weighted Averages

Let D be a bounded Lipschitz polytope and let  $\{Y_\ell\}_{\ell=1}^n$  be a subdivision of D into 69 open Lipschitz polytopes such that 70

$$\alpha_{|\tilde{Y}_{\ell}} = \alpha_{\ell} = \text{const.}$$
 (6)

Furthermore, let  $\mathscr{X} \subset \partial D$  be a manifold of dimension  $0 \le d_{\mathscr{X}} \le d-1$  (usually a 71 vertex, an open subdomain edge or an open face, or a union of these). We define 72

$$\mathscr{X}_{\ell} := \overline{Y}_{\ell} \cap \mathscr{X}.$$
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Some of these sets may be empty or have lower dimension than  $\mathscr{X}$ . However, with 74 the index set  $I_{\mathscr{X}} := \{\ell : \operatorname{meas}_{d_{\mathscr{X}}}(\mathscr{X}_{\ell}) > 0\}$  we can write 75

$$\overline{\mathscr{X}} = \bigcup_{k \in I_{\mathscr{X}}} \overline{\mathscr{X}}_k.$$
 76

In general, for different indices k,  $\ell \in I_{\mathscr{X}}$ , the manifolds  $\mathscr{X}_k$  and  $\mathscr{X}_\ell$  may have a 77 non-trivial intersection or even coincide. For simplicity, we assume that 78

$$k \neq \ell \in I_{\mathscr{X}} \implies \operatorname{meas}_{d_{\mathscr{X}}}(\mathscr{X}_k \cap \mathscr{X}_\ell) = 0.$$

The general case needs more formalism and will be treated in an upcoming paper 80 [10]. Finally, we can define a meaningful trace  $\alpha_{tr} \in L^{\infty}(\mathcal{X})$  of  $\alpha$  by 81

$$\alpha_{\rm tr}(x) = \alpha_k \quad \text{for } x \in \mathscr{X}_k.$$

Let  $\{V^h(D)\}_h$  be a family of  $H^1$ -conforming FE spaces associated with a quasi- 83 uniform family of triangulations of D. For  $v \in V^h(D)$ , we define the weighted 84 (semi)norms and the weighted average on  $\mathscr{X}$  by

$$\|v\|_{L^2(D),\alpha}^2 := \int_D \alpha \, v^2 \,, \quad |v|_{H^1(D),\alpha}^2 := \int_D \alpha \, |\nabla v|^2 \quad \text{and} \quad \overline{v}^{\mathscr{X},\alpha_{\operatorname{tr}}} := \frac{\int_{\mathscr{X}} \alpha_{\operatorname{tr}} \, v}{\int_{\mathscr{X}} \alpha_{\operatorname{tr}}} \,. \qquad \text{86}$$

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We are interested in the following WPI with weighted average:

$$\|u - \overline{u}^{\mathscr{X}, \alpha_{\operatorname{tr}}}\|_{L^{2}(D), \alpha}^{2} \leq C_{P, \alpha}(D, \mathscr{X}; h) \operatorname{diam}(D)^{2} |u|_{H^{1}(D), \alpha}^{2} \quad \forall u \in V^{h}(D).$$
 (7)

In particular, we are interested under which assumptions the parameter  $C_{P,\alpha}(D,\mathcal{X};h)$ is independent of the values  $\{\alpha_{\ell}\}$ .

Sufficient conditions for robustness. We need two crucial assumptions for (7) to 90 be independent of the values  $\{\alpha_{\ell}\}$ . The first assumption is a quasi-monotonicity 91 assumption on  $\alpha$ . It has been introduced in [1] and generalized in [4, 8]. The second 92 assumption states that  $\mathscr X$  "sees" the largest coefficient. 93

**Definition 1.** Let  $0 \le m < d$  and let  $\ell^* := \operatorname{argmax} \alpha_{\ell}$  denote the index of the largest

coefficient.4 95

- (a) We call the region  $P_{\ell_1,\ell_s}:=(\overline{Y}_{\ell_1}\cup\ldots\cup\overline{Y}_{\ell_s})^\circ$ ,  $1\leq\ell_1,\ldots,\ell_s\leq n$  a type-m quasi-96 monotone path from  $Y_{\ell_1}$  to  $Y_{\ell_s}$  (with respect to  $\alpha$ ), if
  - (i) the regions  $Y_{\ell_i}$  and  $Y_{\ell_{i+1}}$  share a common m-dimensional manifold, and 98
  - (ii)  $\alpha_{\ell_1} \leq \alpha_{\ell_2} \leq \ldots \leq \alpha_{\ell_s}$ . 99
- (b) We say that  $\alpha$  is type-m quasi-monotone on D, if for all k = 1, ..., n there exists 100 a quasi-monotone type-m path from  $Y_k$  to  $Y_{\ell^*}$ . 101

**Assumption A2.**  $\alpha$  is type-*m* quasi-monotone on *D* for some  $0 \le m < d$ . 102

**Assumption A3.** 
$$\operatorname{meas}_{d_{\mathscr{X}}}(\mathscr{X} \cap \overline{Y}_{\ell^*}) > 0.$$

In order to formulate our main theorem, we first need some definitions of gener- 104 alized Poincaré constants/parameters. 105

**Definition 2.** (i) For any bounded Lipschitz domain  $Y \subset \mathbb{R}^d$  let  $C_P(Y)$  be the smallest constant such that 107

$$\|v - \overline{v}^Y\|_{L^2(Y)}^2 \le C_P(Y) \operatorname{diam}(Y)^2 |v|_{H^1(Y)}^2 \quad \forall v \in H^1(Y).$$

(ii) Let Z be the finite union of bounded Lipschitz polytopes such that  $\overline{Z}$  is con- 109 nected, and let  $\{\mathcal{T}^h(Z)\}_h$  be a quasi-uniform family of triangulations of Z with the associated continuous piecewise linear FE spaces  $\{V^h(Z)\}_h$ . Let X, 111  $W \subset \overline{Z}$  be manifolds/subdomains of (possibly different) dimension  $\in \{0,\ldots,d\}$ . 112 Let  $C_P(Z,X,W;h)$  be the best parameter such that 113

$$\|v - \overline{v}^X\|_{L^2(W)}^2 \le C_P(Z, X, W; h) \frac{|W|}{|Z|} \operatorname{diam}(Z)^2 |u|_{H^1(Z)}^2 \qquad \forall v \in V^h(Z).$$

|W| and |Z| denote the measures of W and Z (in the respective dimension).

<sup>&</sup>lt;sup>4</sup> We can assume without loss of generality that  $\ell^*$  is unique. By definition, type-m quasimonotonicity implies that otherwise all maximal subregions can be combined into a single subregion.

If Z is connected and if the dimensions of X and W are  $\geq d-1$ , we can define 116 a constant  $C_P(Z,X,W)$  independent of the discretization parameter h such that the 117 inequality in Definition 2(ii) holds for all functions in  $H^1(Z)$ .

**Theorem 1.** Let Assumptions A2 and A3 be satisfied. Then the parameter 119  $C_{P,\alpha}(D,\mathcal{X};h)$  in formula (7) is independent of the values  $\{\alpha_\ell\}_{\ell=1}^n$  and 120

$$C_{P,\alpha}(D,\mathcal{X};h) \leq 2\left[C^{*,1}(h) + C^{*,2}(h)\right]$$
 (8)

with 121

$$egin{aligned} C^{*,1}(h) &:= \sum_{\ell=1}^n rac{|Y_\ell| \operatorname{diam}(P_{\ell,\ell^*})^2}{|P_{\ell,\ell^*}| \operatorname{diam}(D)^2} C_P(P_{\ell,\ell^*},\mathscr{X}_{\ell^*},Y_\ell;h), \ C^{*,2}(h) &:= rac{|D|}{|\mathscr{X}_{\ell^*}|} \sum_{k \in I_{\mathscr{X}^*}} rac{|\mathscr{X}_k| \operatorname{diam}(P_{k,\ell^*})^2}{|P_{k,\ell^*}| \operatorname{diam}(D)^2} C_P(P_{k,\ell^*},\mathscr{X}_{\ell^*},\mathscr{X}_k;h). \end{aligned}$$

*Proof.* Without loss of generality, we may assume that  $\bar{u}^{\mathcal{X},\alpha_{\text{tr}}}=0$ . For each index 122  $\ell=1,\ldots,n$ ,

$$\frac{1}{2} \|u\|_{L^{2}(Y_{\ell})}^{2} \leq \|u - \overline{u}^{\mathscr{X}_{\ell^{*}}}\|_{L^{2}(Y_{\ell})}^{2} + |Y_{\ell}| (\overline{u}^{\mathscr{X}_{\ell^{*}}})^{2}.$$

Due to Assumption A2, there is a quasi-monotone path from  $Y_{\ell}$  to  $Y_{\ell^*}$ . With  $c_{\ell,\ell^*}$ := 124  $C_P(P_{\ell,\ell^*}, \mathscr{X}_{\ell^*}, Y_{\ell}; h)$ , summation over  $\ell = 1, \ldots, n$  yields

$$\frac{1}{2} \|u\|_{L^{2}(D),\alpha}^{2} \leq \sum_{\ell=1}^{n} c_{\ell,\ell^{*}} \frac{|Y_{\ell}|}{|P_{\ell,\ell^{*}}|} \operatorname{diam}(P_{\ell,\ell^{*}})^{2} \underbrace{\alpha_{\ell} |u|_{H^{1}(P_{\ell,\ell^{*}})}^{2}}_{\leq |u|_{H^{1}(D),\alpha}^{2}} + \underbrace{\sum_{\ell=1}^{n} \alpha_{\ell} |Y_{\ell}|}_{\leq \alpha_{\ell^{*}} |D|} (\overline{u}^{\mathscr{X}_{\ell^{*}}})^{2},$$

where we have used Definition 2(ii) and the quasi-monotonicity of  $P_{\ell,\ell^*}$ . The first 126 sum is bounded by  $C^{*,1}(h)\operatorname{diam}(D)^2|u|^2_{H^1(D),\alpha}$ . To bound the remaining term, we 127 use Cauchy's inequality and the definition of  $\alpha_{\mathrm{tr}}$ :

129

$$|lpha_{\ell^*}|D|\left(\overline{u}^{\mathscr{X}_{\ell^*}}
ight)^2 \le rac{|D|}{|\mathscr{X}_{\ell^*}|} |lpha_{\ell^*}||u||_{L^2(\mathscr{X}_{\ell^*})}^2 \le rac{|D|}{|\mathscr{X}_{\ell^*}|} ||u||_{L^2(\mathscr{X}),lpha_{\mathrm{tr}}}^2.$$

A variational argument yields

$$\begin{split} \|u\|_{L^{2}(\mathscr{X}),\alpha_{\mathrm{tr}}}^{2} & \leq \|u - \underline{\overline{u}}^{\mathscr{X},\alpha_{\mathrm{tr}}}\|_{L^{2}(\mathscr{X}),\alpha_{\mathrm{tr}}}^{2} = \inf_{c \in \mathbb{R}} \|u - c\|_{L^{2}(\mathscr{X}),\alpha_{\mathrm{tr}}}^{2} \\ & \leq \|u - \overline{u}^{\mathscr{X}_{\ell^{*}}}\|_{L^{2}(\mathscr{X}),\alpha_{\mathrm{tr}}}^{2} = \sum_{k \in I_{\mathscr{X}}} \alpha_{k} \|u - \overline{u}^{\mathscr{X}_{\ell^{*}}}\|_{L^{2}(\mathscr{X}_{k})}^{2}. \end{split}$$

Now, we have

$$\alpha_{k} \| u - \overline{u}^{\mathscr{X}_{\ell^{*}}} \|_{L^{2}(\mathscr{X}_{k})}^{2} \leq C_{P}(P_{k,\ell^{*}}, \mathscr{X}_{\ell^{*}}, \mathscr{X}_{k}; h) \frac{|\mathscr{X}_{k}|}{|P_{k,\ell^{*}}|} \operatorname{diam}(P_{k,\ell^{*}})^{2} \alpha_{k} |u|_{H^{1}(P_{k,\ell^{*}})}^{2}.$$
 131

Using the quasi-monotonicity of  $\alpha$  on  $P_{k,\ell^*}$  finally leads to (8).

Necessity of the conditions. As discussed in [8, Sect. 3.1], Assumption A2 is necessary to ensure that  $C_{P,\alpha}(D,\mathcal{X};h)$  is independent of the values  $\{\alpha_{\ell}\}$ .

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To see that A3 is necessary as well, assume that meas<sub>d or</sub>  $(\mathscr{X} \cap \overline{Y}_{\ell^*}) = 0$ . We 135 choose a function u which is one on  $Y_{\ell^*}$ . Since the average functional  $v \mapsto \overline{v}^{\mathscr{X},\alpha_{\text{tr}}}$  is 136 independent of  $\alpha_{\ell^*}$ , we can prescribe values of u on  $\mathscr X$  such that  $\overline{u}^{\mathscr X}, \alpha_{tr} = 0$  and 137 continuously extend u into  $D \subset \overline{Y}_{\ell^*}$ . The whole construction of u is independent 138 of  $\alpha_{\ell^*}$ , Since  $\nabla u=0$  on  $Y_{\ell^*}$ , the seminorm  $|u|_{H^1(D),\alpha}$  is independent of  $\alpha_{\ell^*}$  as well. 139 However,  $||u||_{L^2(D)}^2 \alpha \ge \alpha_{\ell^*} |Y_{\ell^*}|$ . Therefore, if  $\alpha \le \alpha_k$  on  $D \setminus Y_{\ell^*}$ , then  $C_{P,\alpha}(D, \mathcal{X}; h) =$  $\mathscr{O}(\frac{\alpha_{\ell^*}}{\alpha_k})$  for  $\alpha_{\ell^*}/\alpha_k \to \infty$ . This means that Assumptions A2 and A3 in some sense 141 characterize the robustness of the WPI with weighted average. 142

#### 3 Robustness Proof of FETI-DP

To analyze the robustness of FETI-DP, we need the following assumption.

**Assumption A4.** For each subdomain  $\Omega_i$  and for each subdomain edge (face)  $\mathscr{E}$  of 145  $\Omega_i$ , there is a Lipschitz domain  $D_{i,\mathscr{E}} \subset \Omega_i$ , such that  $\mathscr{E} \subset \partial D_{i,\mathscr{E}}$  and Assumptions A2 146 and A3 are satisfied for  $D = D_{i,\mathcal{E}}$  and  $\mathscr{X} = \mathcal{E}$ . The union of all the regions  $D_{i,\mathcal{E}}$ covers a boundary layer  $\Omega_{i,\eta_i}$  of width  $\eta_i \ge h$  of  $\Omega_i$  (see e.g. [6, Definition 2.6]). 148

**Theorem 2.** Let Assumptions A1 and A4 hold. Then the condition number  $\kappa(M^{-1}F)$  149 for the FETI-DP method is independent of the values of the coefficient α, in partic- 150 ular of any non-resolved jumps.

Due to space limitations we only give a sketch of the proof. A detailed proof will 152 be given in [10], together with a more detailed statement of Theorem 2 that makes 153 precise the dependence of  $\kappa(M^{-1}F)$  on geometric parameters, such as the ratios 154  $\operatorname{diam}(\Omega_i)/h$  and  $\operatorname{diam}(\Omega_i)/\eta_i$ .

Let  $\mathcal{H}_i$  denote the discrete  $\alpha$ -harmonic extension from  $\partial \Omega_i$  to  $\Omega_i$  and let

$$|w|_S^2 := \sum_{i=1}^N |\mathscr{H}_i w|_{H^1(\Omega_i),\alpha}^2.$$
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Then, following [12, Sect. 6.4.3], a bound of the kind

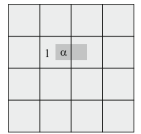
$$|P_D w|_S^2 \le \omega |w|_S^2 \qquad \forall w \in \widetilde{W}, \tag{9}$$

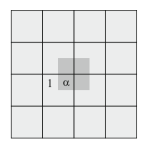
where  $P_D := B_D^\top B$ , implies that  $\kappa(M^{-1}F) \leq \omega$ .

As in the proof of [9, Lemma 5.6; formula (5.24)], we can introduce a set of 160 cut-off functions associated with each subdomain edge (face) & whose support is 161 contained in  $D_{i,\mathscr{E}}$ . It then follows that, for any  $w \in W_{\Pi} \oplus W_{\Delta}$ , 162

$$|P_D w|_S^2 \leq C \sum_{i=1}^N \left[ |\mathscr{H}_i w_i|_{H^1(\Omega_i),\alpha}^2 + \sum_{\mathscr{E}} \frac{1}{\operatorname{diam}(\Omega_i)^2} \|\mathscr{H}_i w_i - \overline{w_i}^{\mathscr{E}}\|_{L^2(D_{i,\mathscr{E}}),\alpha}^2 \right], \tag{163}$$

where C depends on diam $(\Omega_i)/h$  and diam $(\Omega_i)/\eta_i$ , but it is independent of the values 164  $\{\alpha_\ell\}$ . By Theorem 1, we can bound each of the weighted  $L^2$  norms by the weighted 165  $H^1$  seminorm of  $\mathcal{H}_i w_i$ , and thus obtain (9). 166





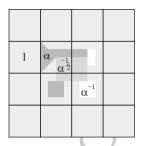


Fig. 2. Edge-island (left), cross-point island (middle), complicated coefficient (right)

$\alpha$	condition	#iterations	t1.1	$\alpha$	condition	#iterations	t2.1 $lpha$	condition	#iterations	t3.1
1	1.58	10	t1.2	1	1.58	10	t2.2 1	1.58	10	t3.2
$10^{1}$	1.57	10	t1.3	$10^{1}$	1.59	10	t2.3 $10^1$	1.61	11	t3.3
$10^{3}$	1.56	10	t1.4	$10^{3}$	1.59	10	$t2.4  ext{ } 10^2$	1.62	11	t3.4
$10^{5}$	1.56	10	t1.5	$10^{5}$	1.59	10	$t2.5  ext{ } 10^3$	1.62	11	t3.5
$10^{7}$	1.56	10	t1.6	$10^{7}$	1.59	10	t2.6 $10^4$	1.62	11	t3.6
$10^{-1}$	1.70	10		$10^{-1}$	1.57	10	t2.7 $10^{-1}$		11	t3.7
$10^{-3}$	1.74	10		$10^{-3}$		10	t2.8 $10^{-2}$		11	t3.8
$10^{-5}$	1.74	10	t1.9	$10^{-5}$	1.57	10	t2.9 $10^{-3}$	1.59	11	t3.9
$10^{-7}$	1.74	11	t1.10	$010^{-7}$	1.57	10	t2.10 $10^{-2}$	1.59	11	t3.10

**Table 1.** Edge-island (left), crosspoint-island (middle), complicated coefficient (right), H/h =32.

#### 4 Numerical Results

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We provide results for the three examples shown in Fig. 2. Note that in the last 168 example, the coefficient is not quasi-monotone on one of the subdomains, but satisfies Assumptions A1 and A4. In our implementation we used PARDISO [11]. 170 The estimated condition numbers and the number of PCG iterations are displayed 171 in Table 1. They clearly confirm Theorem 2.

5 Conclusion 173

We analyse a FETI-DP method for the scalar elliptic PDE (1) with possible jumps in 174 the diffusion coefficient alpha. We show that provided weighted edge/face averages 175 are used, the condition number of the preconditioned system is independent of coefficient jumps. The essential assumptions are A1 and A4, i.e., the coefficient does not 177 jump both across and along any interfaces between two subdomains and the coeffi- 178 cient is quasi-monotone in the vicinity of any edge/face within each subdomain. The 179 key theoretical tool that is of interest in itself is a novel weighted Poincaré inequality 180 for functions with suitably chosen vanishing weighted face/edge averages. We are 181 able to show that under Assumption A4, the Poincare constant of each neighborhood 182  $D_{i,\mathcal{E}}$  can be bounded independent of jumps.

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As in our previous work [8], the Poincaré constants (and thus also the condition 184 number) will also depend on the "geometry" of the coefficient variation. In partic- 185 ular, for piecewise constant coefficients it will in general depend on the geometry 186 of the subregions where the coefficient is constant. We did not give details of this 187 dependence here, but this will be done in an upcoming paper [10] (using [8]). Cases 188 where the coefficient jumps both along and across subdomain interfaces appear to be 189 substantially harder to be treated and are also the subject of our future investigations. 190

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## **Monotone Multigrid Methods Based on Parametric Finite Elements**

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Summary. In this paper, a particular technique for the application of elementary multilevel 8 ideas to problems with warped boundaries is studied in the context of the numerical simulation 9 of elastic contact problems. Combining a general multilevel setting with a different perspec- 10 tive, namely an advanced geometric modeling point of view, we present a (monotone) multigrid method based on a hierarchy of parametric finite element spaces. For the construction, a 12 full-dimensional parameterization of high order is employed which accurately represents the 13 computational domain.

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The purpose of the volume parametric finite element discretization put forward here is 15 two-fold. On the one hand, it allows for an elegant multilevel hierarchy to be used in preconditioners. On the other hand, it comes with particular advantages for the modeling of contact problems. After all, the long-term objective lies in an increased flexibility of hp-adaptive 18 methods for contact problems.

1 Introduction 20

In the numerical simulation of elastic contact problems, the treatment of the non- 21 penetration conditions at the potential contact boundary is of particular importance 22 for both the quality of a finite element approximation and the overall efficiency of the 23 algorithms. A vital challenge is to achieve an accurate description of geometric fea- 24 tures, e.g., of warped surfaces, often incorporated in three-dimensional models from 25 computer-aided design (CAD). Here, we investigate a new connection of different 26 numerical methods, namely modern discretization techniques for partial differential 27 equations on complex geometries on the one side and fast multilevel solvers for con- 28 strained minimization problems on the other side.

It is fair to say that the development of hp-adaptive methods for contact prob- 30 lems has not yet reached a mature state; see, e.g., [2] and the references therein. 31 Partly, this is due to the difficulties concerning the geometric representation of the 32 computational domain. A generally accepted paradigm is, though, that high order (fi- 33 nite element or boundary element) methods need high order meshes [11, 14]. This is 34 especially difficult for three-dimensional multi-body contact problems. In this case, 35 the application of non-conforming domain decomposition techniques [16] to realize 36

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an optimal information transfer across geometrically non-matching warped contact 37 interfaces is a highly demanding task. For low order finite elements, this has been 38 achieved, among others, by the authors; see [6].

The perspective we offer here is a parametric finite element method. For hp-40 adaptive methods, it is convenient to have a parameterization describing the geometry 41 accurately ready to hand. This is because a change of the computational domain 42 due to locally altered polynomial degree is not desirable. Therefore, it is reasonable 43 to uncouple the representation of the geometry on the one hand and of a scale of 44 approximation spaces for the discrete solution on the other hand. These two purposes 45 are usually not separated properly. But of course, one can find curved elements of 46 other than isoparametric structure in some form or another in the literature; see, e.g., 47 [8, 17] or the monograph [3] and the references therein. Note that, for similar reasons, 48 an "isogeometric" concept, which uses NURBS bases for both the description of the 49 geometry and the discrete solution of the differential equation, has been introduced 50 in [11].

For practical computations, the development of fast and robust solvers is equally 52 important. As this issue has not yet been in the main focus of, e.g., the isogeometric 53 analysis [11], we would like to contribute ideas from the field of multilevel meth- 54 ods for variational inequalities. More precisely, we show how to use a monotone 55 multigrid method to efficiently solve the non-linear contact problem discretized with 56 low order parametric finite elements. Note that the actual treatment of higher order 57 elements is beyond the scope of the present discussion.

To obtain multilevel parametric finite element spaces in case d=3, we use a 59 full-dimensional parameterization, constructed by tetrahedral transfinite interpola- 60 tion [15] of CAD data, to lift standard Lagrange elements to the computational do- 61 main. Note that, similarly, a surface parameterization has been used in a wavelet 62 Galerkin scheme for boundary integral equations; see [10]. Such a procedure may 63 serve as an essential prerequisite to tackle the problems mentioned above. In par- 64 ticular, many of the issues arising in the generation of p-version meshes for curved 65 boundaries [14] can be avoided in a quite elegant way. In this sense, although rather 66 expensive, the use of a high order parameterization permits maximal freedom in an 67 hp-adaptive discretization scheme. We presume that the present concept can also be 68 combined with the ideas in [6].

All in all, our results constitute real progress made in the development of an 70 efficient hp-adaptive simulation environment for elastic contact problems in case of 71 complex three-dimensional geometries.

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### 2 Parametric Finite Elements

In this section, we introduce a parametric finite element discretization. On the one 74 hand, this method uses much more geometric information from a CAD model than 75 standard finite elements; on the other hand, we do not use the same functions for the 76 discrete approximation of the displacement field as for the representation of the geometry, which is done in the so-called "isogeometric analysis" introduced in [11]. We 78

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use the associated space hierarchy in Sect. 3 to build a monotone multigrid method 79 for low order elements.

In the following, the symbols  $\varphi$  with some indices stand for certain full-dimensional parameterizations or finite element transformations. We denote the (closed) 82 *d*-simplex by  $\Delta^d$  and its faces by  $\Delta^d_j$ ,  $j \in \{1, ..., d+1\}$ . To describe the elastic 83 body (here, d=3) by a practicable parameterization, we consider a non-overlapping 84 simplicial decomposition of the computational domain  $\Omega\subset\mathbb{R}^d$  into a fixed number 85 of  $K \ge 1$  subdomains. Formally this reads as

$$\overline{\Omega} = \bigcup_{k=1}^K \overline{\Omega}_k = \bigcup_{k=1}^K \varphi_k(\Delta^d),$$

where the notation already indicates that the subdomains  $(\Omega_k)_{k=1,...,K}$  appear as particular images of the simplex  $\Delta^d$  under suitable parameterizations  $(\varphi_k)_{k=1,\dots,K}$ . This 88 is illustrated in Fig. 1 (right).

Let us assume that the faces of the simplicial cells  $\Omega_k$ , namely the surfaces 90  $\varphi_k(\Delta_i^d), k \in \{1, \dots, K\}, j \in \{1, \dots, d+1\},$  are given as B-patches. This way to rep- 91 resent polynomial surfaces is analyzed in [4]. In this case, the author of [15] pro- 92 poses to construct the full-dimensional mappings  $\varphi_k : \Delta^d \to \mathbb{R}^d$ ,  $k \in \{1, \dots, K\}$ , as 93 transfinite interpolations of the surface values from the CAD model using certain 94 blending functions. Particularly, the single parameterizations are smooth and they 95 match across these B-patch surfaces if the surfaces themselves match. This gives rise 96 to a consistent global parameterization which we do not write down explicitly. We 97 note that this global mapping is continuous but not necessarily differentiable across 98 the interior interfaces. In addition, one can guarantee that each parameterization  $\varphi_k$  99 satisfies the regularity assumption

$$\det(\nabla \varphi_k) > 0 \quad \text{in } \Delta^d. \tag{1}$$

In fact, this is one of the main results of [15].

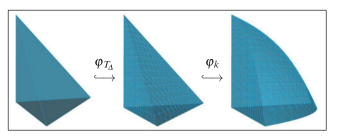
In the following, we define the parametric finite element spaces in a rather 102 straightforward way via a lift of standard Lagrange finite elements. For this purpose, 103 let  $(\mathscr{T}^k)_{\ell\in\mathbb{N}}$  be a family of nested simplicial meshes of  $\Delta^d$  for each  $k\in\{1,\ldots,K\}$ . 104 To keep the global finite element spaces conforming, we assume that, at each level 105  $\ell \in \mathbb{N}$ , the meshes meeting at the faces of the simplicial subdomains  $\Omega_k$  of  $\Omega$  match. 106 Let  $\widehat{T}$  be the reference element; here,  $\widehat{T} = \Delta^d$ . Then, for each  $T_\Delta \in \mathscr{T}^k_\ell$ , there is an 107 affine mapping  $\varphi_{T_{\Lambda}}: \widehat{T} \to \Delta^d$  such that  $\varphi_{T_{\Lambda}}(\widehat{T}) = T_{\Delta}$ .

Now, we give a concise description of the parametric elements in  $\Omega$  by employing the special finite element transformations

$$\varphi_T := \varphi_k \circ \varphi_{T_\Delta} : \widehat{T} \to \mathbb{R}^d, \tag{2}$$

which are diffeomorphisms between the reference element  $\widehat{T}$  and the actual elements. That way, the parametric elements at level  $\ell \in \mathbb{N}$  are identified as the images of the elements of the meshes  $(\mathscr{T}_{\ell}^k)_{k=1,\ldots,K}$ ; see Fig. 1. More precisely, a family of parametric meshes  $(\mathcal{T}_{\ell})_{\ell \in \mathbb{N}}$  of  $\Omega$  can be defined by

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Fig. 1. From left to right: the reference element  $\hat{T} = \Delta^3$ ; a mesh of the simplex  $\Delta^3$ ; a parametric mesh (here, K = 1) where each element is an image of an affine element; a tetrahedral decomposition of a cylinder with K = 8

$$\mathscr{T}_{\ell} := \left\{ T = \varphi_T(\widehat{T}) = \varphi_k(\varphi_{T_{\Delta}}(\widehat{T})) \mid 1 \leq k \leq K, \ T_{\Delta} \in \mathscr{T}_{\ell}^k \right\}, \quad \forall \ \ell \in \mathbb{N}.$$

Assume that this family of global meshes is shape regular and quasi-uniform. Note 115 that assumption (1), combined with the continuous differentiability of the mappings 116  $(\varphi_k)_{k=1,\ldots,K}$  in the compactum  $\Delta^d$ , implies that it is sufficient to ensure these regularity conditions for each sequence  $(\mathscr{T}_{\ell}^{k})_{\ell \in \mathbb{N}}$  separately as far as we keep K fixed.

Finally, let  $\mathbb{P} := \mathbb{P}_r(\widehat{T})$  be the space of polynomials of degree r in  $\widehat{T}$ . Then, for  $\ell \in \mathbb{N}$ , the parametric finite element space associated with the parametric mesh  $\mathscr{T}_{\ell}$  is 120

$$X_{\ell} := \left\{ v \in \mathcal{C}^{0}(\Omega) \mid \forall T \in \mathcal{T}_{\ell} \ \exists \ w \in \mathbb{P} : \ v(\mathbf{x}) = w(\varphi_{T}^{-1}(\mathbf{x})), \ \forall \ \mathbf{x} \in T \right\}$$

$$= \left\{ v \in \mathcal{C}^{0}(\Omega) \mid v \circ \varphi_{T} \in \mathbb{P}, \ \forall \ T \in \mathcal{T}_{\ell} \right\}.$$

$$(3)$$

Note that, in principle, the above definition makes sense for any reasonable set of 121 finite element transformations  $(\varphi_T)_{T\in\mathcal{T}_t}$ . In case the mappings are constructed as 122 in (2) via the high order parameterization from [15], this is a "superparametric" con- 123 cept if the degree r is small. This is in contrast to the subparametric or isoparametric 124 finite elements which are usually considered in the literature; see [3].

From a practical point of view, virtually every kind of parameterization can be 126 employed with the following qualification. For an efficient assembly of the stiffness 127 matrix and the right hand side via sufficiently accurate (at best exact) numerical 128 quadrature, the derivatives of the resulting finite element transformations (2) and the 129 mappings themselves must be easy to evaluate; see, e.g., [1].

### Discretization of Signorini's Problem

Let us now apply the above concept to a contact problem in elasticity to find the 132 deformation of a linear elastic body  $\Omega$  in contact with a rigid obstacle. For this 133 purpose, let the boundary be decomposed into pairwise disjoint parts:  $\partial\Omega=\overline{\Gamma}_D\cup$  134  $\overline{\Gamma}_N \cup \overline{\Gamma}_C$ . Assume that the Dirichlet boundary  $\Gamma_D$  is of positive Lebesgue measure in 135 dimension d-1. Moreover, the condition  $\overline{\Gamma}_C \cap \overline{\Gamma}_D = \emptyset$  may hold.

Let **n** be the outer normal vector field on  $\partial \Omega \in \mathscr{C}^1$ ; the initial gap to the rigid 137 obstacle in this direction is given as a function  $g:\Gamma_C\to\mathbb{R}_{>0}$ . Then, for sufficiently smooth prescribed volume and surface force densities  $\mathbf{f} = (f_i)$  and  $\mathbf{p} = (p_i)$ , the 139 displacement field  $\boldsymbol{u}:\Omega\to\mathbb{R}^d$  solves the boundary value problem 140

$$\begin{aligned}
-\sigma_{ij}(\boldsymbol{u})_{,j} &= f_i & \text{in } \Omega, \\
\boldsymbol{u} &= \boldsymbol{0} & \text{on } \Gamma_D, \\
\sigma_{ij}(\boldsymbol{u})n_j &= p_i & \text{on } \Gamma_N, \\
\boldsymbol{u} \cdot \boldsymbol{n} &\leq g & \text{on } \Gamma_C,
\end{aligned} \tag{4}$$

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where  $\sigma_{ij}(\mathbf{u}) = A_{ijlm}u_{l,m}$  are the stresses and  $\mathbf{A} = (A_{ijlm})$  is Hooke's tensor. The existence of a unique weak solution follows from Lions' and Stampacchia's lemma. 142

We use the vector-valued parametric finite element space  $\mathbf{X}_{\ell} := (X_{\ell})^d$  defined 143 by (3) with r=1 and denote the set of nodes by  $\mathcal{N}_{\ell}$ . As usual, the non-penetration 144 conditions on the possible contact boundary  $\Gamma_{\!C}$  are merely enforced at the potential 145 contact nodes  $\mathcal{N}_{\ell}^{C} = \mathcal{N}_{\ell} \cap \Gamma_{C}$ ; see below. Then, a discretization of Signorini's problem (4) with one-sided constraints is obtained by specifying a variational inequality

find 
$$\mathbf{u}_{\ell} \in \mathbf{K}_{\ell}$$
 such that  $a(\mathbf{u}_{\ell}, \mathbf{v} - \mathbf{u}_{\ell}) \ge f(\mathbf{v} - \mathbf{u}_{\ell}), \forall \mathbf{v} \in \mathbf{K}_{\ell},$  (5)

on a suitable set of admissible displacements

$$\mathbf{K}_{\ell} := \{ \mathbf{v} \in \mathbf{X}_{\ell} | \mathbf{v} = \mathbf{0} \text{ on } \Gamma_{D}, (\mathbf{v} \cdot \mathbf{n})(p) \leq g(p), \forall p \in \mathcal{N}_{\ell}^{C} \}.$$

In the discrete variational inequality (5), the (bi-)linear forms a and f representing the elastic energy and the applied forces, respectively, are given by  $a(\mathbf{u}, \mathbf{v}) := 150$  $\int_{\Omega} A_{ijlm} u_{l,m} v_{i,j} d\mathbf{x}$  and  $f(\mathbf{v}) := \int_{\Omega} f_i v_i d\mathbf{x} + \int_{\Gamma_{i}} p_i v_i d\mathbf{a}$ .

Although, from a modeling point of view, as much geometric information as 152 possible should be used for an accurate description of contact phenomena, we re- 153 mark that a strong pointwise non-penetration condition everywhere on  $\Gamma_C$  is usually not suitable for the variational formulation on which the (parametric) finite element 155 method relies. Besides, a decoupled set of constraints is preferable for a variety of 156 reasons. The common remedy is to prescribe the contact constraints with respect to 157 a suitable cone of Lagrange multipliers. This requires the introduction of appropri- 158 ate sets of functionals in  $(H^{\frac{1}{2}}(\Gamma_C))'$ . To retain inequality constraints which can be 159 enforced merely by looking at the nodes, one can employ discontinuous test spaces 160 described, e.g., in [7].

The quality of a priori error estimates for the above discretization certainly de- 162 pends on a number of aspects which have to be examined more closely. Beside regularity assumptions for the continuous solution, the balance of the primal degrees of 164 freedom and the constraints by means of an inf-sup condition and certain properties 165 of the parameterization, e.g., the regularity (1), influence the error analysis.

# 3 Monotone Multigrid Method for Parametric Elements

Similarly to some of the approaches reviewed in [5, Chap. 4], the scale of parametric 168 finite element spaces constitutes an adjusted discretization technique which allows 169 for an almost straightforward application of multilevel ideas. In this section, we examine the constructed space hierarchy, which we presume to possess the required 171 approximation properties, and the corresponding natural transfer operators in a little 172 more detail.

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For the solution of the discrete variational inequality, we propose a monotone 174 multigrid method [12]; see [13] for an overview of this and other solution strategies 175 for contact problems and more references. Here, the non-penetration conditions at 176 the potential contact nodes are treated by a non-linear block Gauß-Seidel smoother 177 at the finest level L. Let  $\tilde{u} \in K_L$  be a preliminary approximate solution (i.e., a current 178 admissible iterate). Then, in the next step, a linear multilevel preconditioner depending on  $\tilde{u}$  is employed, which acts only on the space  $\{v \in X_L | (v \cdot n)(p) = 0, \forall p \in 180\}$  $\mathcal{N}_L^C$  with  $(\tilde{\boldsymbol{u}} \cdot \boldsymbol{n})(p) = g(p)$ . The construction of the required coarse spaces from 181 the spaces  $(\boldsymbol{X}_\ell)_{\ell < L}$  involves local modifications of the coarse level matrices resulting from recursively truncated basis functions; see, e.g., [13].

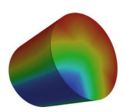
By construction, the spaces defined by (3) are nested. This is an immediate consequence of the fact that the parameterization is fixed and does not change with the 185 index  $\ell$ . Still, let us formulate this statement in the following lemma and give an 186 elementary proof of the assertion.

### **Lemma 1.** The parametric finite element spaces $(X_{\ell})_{\ell \in \mathbb{N}}$ are nested.

*Proof.* For  $\ell \geq 1$ , let  $\nu \in X_{\ell-1}$  be arbitrary. Then, for  $T \in \mathcal{T}_{\ell-1}$  there is a unique element  $T_{\Delta} \in \mathscr{T}^k_{\ell-1}$  for some  $k \in \{1, \dots, K\}$  such that  $\varphi_k(T_{\Delta}) = T$ . Let  $(T^i_{\Delta})_{i=1,\dots,N}$  be the children of  $T_{\Delta}$  in  $\mathcal{T}_{\ell}^{k}$ . In general,  $1 \le N \le 2^{d}$ ; in case of standard uniform refinement of the simplices, it is  $N=2^d$ . We have the corresponding set of elements  $(T^i)_{i=1,\dots,N}$  192 in  $\mathscr{T}_\ell$  with  $T^i=\varphi_k(T^i_\Delta)$  for  $i\in\{1,\ldots,N\}$ . By assumption,  $v\circ\varphi_T=v\circ\varphi_k\circ\varphi_{T_\Delta}\in\mathbb{P}$ . 193 Therefore, it is  $v \circ \varphi_{T^i} = v \circ \varphi_k \circ \varphi_{T^i} \in \mathbb{P}$  because  $T^i_{\Lambda} \subset T_{\Delta}$  and the finite element transformations are affine. As each element of  $\mathcal{T}_{\ell}$  appears as the child of an element in  $\mathscr{T}_{\ell-1}$  in the above fashion, we obtain  $v \in X_{\ell}$ . Consequently,  $X_{\ell-1} \subset X_{\ell}$  for all  $\ell \geq 1$ . 196

Therefore, no advanced transfer concepts need to be studied here as the canonical 199 inclusion  $\mathscr{I}_{\ell-1}^{\ell}: X_{\ell-1} \to X_{\ell}$  is the most natural operator to be used as prolongation. 200 Note that these operators only depend on the logical structure; as in the standard 201 nested case, the representing matrices contain the entries 0, 0.5 and 1 and may be 202 computed from the neighborhood relations in and between the simplicial meshes 203  $(\mathscr{T}_{\ell-1}^{\bar{k}})_{k=1,\ldots,K}$  and  $(\mathscr{T}_{\ell}^{\bar{k}})_{k=1,\ldots,K}$ . This is because the respective multilevel basis is 204 defined via a lift by proceeding as in (3). As a result, for a fixed finest level L, the 205 computation of the matrices  $\mathbf{I}_{\ell-1}^{\ell} \in \mathbb{R}^{|\mathcal{N}_{\ell}| \times |\mathcal{N}_{\ell-1}|}$  for  $\ell \in \{1, \dots, L\}$  between the nested 206 spaces  $(X_\ell)_{\ell=0,...,L}$  does not need the parameterization. However, the computation of 207 the outer normals  $(\mathbf{n}(p))_{p \in \mathcal{N}_L^C}$  and also of the values  $(g(p))_{p \in \mathcal{N}_L^C}$  for the prescription 208 of the contact constraints may require access to the mappings  $(\varphi_k)_{k=1,\ldots,K}$ .

We anticipate that the constructed coarse spaces have the desired multilevel ap- 210 proximation properties. More precisely, under mild assumptions on the employed 211 parameterization mappings  $(\varphi_k)_{k=1,...,K}$ , the relevant Jackson- and Bernstein-type inMonotone Multigrid Methods Based on Parametric Finite Elements



L	#elements	#dof	#steps	$ ilde{ ho}$	$ \mathscr{A}_L $
0	96	107	8 (2)	0.032	3
1	768	615	10(3)	0.031	15
2	6,144	3,915	11 (4)	0.065	58
3	49,152	27,795	13 (6)	0.091	199
4	393,216	209,187	14 (6)	0.102	753
5	3,145,728	1,622,595	15 (8)	0.114	2,984

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Fig. 2. Contact problem of a parameterized cylinder with a rigid obstacle shaped like a broad channel. The *colors* indicate the displacement in  $e_3$ -direction. Problem (5) is solved by a conjugate gradient method preconditioned by the monotone multigrid method ( $\mathcal{V}(3,3)$ -cycle)

equalities transfer from the standard finite element spaces to the parametric spaces; 213 see also [9].

Finally, we point out that no modifications are necessary in the code of the solver 215 provided that the local normal/tangential coordinate systems can be computed from 216 the parameterization. Consequently, a monotone multigrid method can be employed 217 for contact problems discretized with parametric finite elements in the quite straight- 218 forward way outlined above. Figure 2 shows a numerical example illustrating the 219 performance of the method for d=3. The number of active nodes where the constraints are binding is denoted by  $|\mathcal{A}_L|$ . We report on the asymptotic convergence rate 221  $\tilde{\rho}$  of a conjugate gradient method preconditioned by the monotone multigrid method 222  $(\mathcal{V}(3,3)$ -cycle). Starting with the initial iterate zero at each refinement level (i.e., 223 no nested iteration), we list the number of total steps needed to reduce the norm of 224 the residual to less than  $10^{-10}$ . The count of included non-linear steps is given in 225 brackets (e.g., for L = 5, the active set is found after 8 of the 15 cycles such that the remaining 7 steps are linear). Note that the pcg error reduction rate  $\tilde{\rho}$  corresponds to 227 this linear iteration phase where the active set has already been identified.

4 Conclusion 229

The results described in this paper certainly have preliminary character; the perfor- 230 mance of the presented algorithms needs to be studied in more detail. This is work in 231 progress. However, the experiments so far show that (monotone) multigrid methods 232 based on parametric finite elements work as expected; see Fig. 2. Still, the effort of 233 constructing a (high order) parameterization by the methodology developed in [15] 234 especially pays if there is also a considerable gain on the modeling side. Here, the 235 effect of this special resolution of the boundary on the discrete approximation of contact phenomena or general boundary effects needs to be investigated more closely.

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# TFETI Scalable Solvers for Transient Contact Problems

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Summary. We review our results obtained by application of the TFETI domain decomposition method to implement the time step of the Newmark scheme for the solution of transient contact problems without friction. If the ratio of the decomposition and discretization parameters is kept uniformly bounded as well as the ratio of the time and space discretization, then the cost of the time step is proved to be proportional to the number of nodal variables. The algorithm uses our MPRGP algorithm for the solution of strictly convex bound constrained quadratic programming problems with optional preconditioning by the conjugate projector to the subspace defined by the trace of the rigid body motions on the artificial subdomain interfaces. The optimality relies on our results on quadratic programming, the theory of the preconditioning by a conjugate projector for nonlinear problems, and the classical bounds on the spectrum of the mass and stiffness matrices. The results are confirmed by numerical solution of 3D transient contact problems.

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1 Introduction 20

The transient multibody contact problems are important in many applications arising in mechanical or civil engineering. However, it is not easy to provide a useful 22
solution to realistic problems. The reasons include the lack of smoothness, which 23
puts high demand on the construction of effective time discretization schemes, the 24
strong nonlinearity arising from the non-interpenetration boundary conditions, and 25
large dimension of the problems resulting from the space discretization. These complications stimulated extensive research activities both from the theoretical point of 27
view (see, e.g., [4]), or the numerical point of view (see, e.g., [10], or [11]).

Numerical solution of transient contact problems usually comprises several steps. 29 Starting from a week formulation of the conditions of equilibrium and boundary 30 conditions, the problem is first discretized in space by the finite element method in 31 a similar way as the related static problem. The resulting semidiscrete problem is then 32 discretized by a suitable time discretization scheme. The time integration requires a 33 special attention to guarantee stability of the algorithm and to avoid non-physical 34 oscillations that result from application of the standard time discretization methods 35 for unconstrained problems. Such schemes were proposed by many authors (see [6, 36])

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7, 9, 10]). In our approach, we use a combination of the standard finite element space 37 discretization with the contact stabilized Newmark scheme introduced by Krause and 38 Walloth [9] that reduces the solution of the transient contact problem to a sequence 39 of strictly convex quadratic programming (QP) problems with inequality constraints 40 that describe the non-interpenetration conditions.

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The final step amounts to the solution of QP problems of large dimension, pos- 42 sibly with millions of nodal variables and many inequality constraints. In this paper 43 we propose to resolve the auxiliary problems by our variant of the FETI domain de- 44 composition method called TFETI (total finite element tearing and interconnecting, 45 Dostál et al. [1]). Our research has been motivated by our recent results in develop- 46 ment of optimal algorithms for the frictionless static problems [1] that combine ef- 47 fective FETI preconditioning of both linear and nonlinear steps with our algorithms 48 for the solution of bound constrained QP problems [3]. An important feature of our 49 QP algorithms is the error estimate in terms of the bound on the condition number of 50 the Hessian matrix of the cost function.

### 2 Transient Contact Problem and Its Discretization Using TFETI 52

The starting point of our exposition is the discretized transient multibody contact 53 problem resulting from application of our TFETI domain decomposition. The reason 54 is that a little is known about the solvability of the weak formulation of the transient 55 contact problem (see, e.g., [4]), so we shall assume in what follows that its solution 56  ${\bf u}$  exists. Moreover, we shall assume that  ${\bf u}$  is sufficiently smooth so that  $\ddot{\bf u}$  exists in 57 some reasonable sense and can be approximated by finite differences. More specific 58 choice of the solution space can be found, e.g., in [4] or in [6].

To discretize the multibody contact problem using TFETI, we tear each body 60 from the part of the boundary with the Dirichlet boundary conditions, decompose 61 each body into subdomains, assign each subdomain a unique number, and introduce 62 new "gluing" conditions on the artificial subdomain interfaces and on the boundaries 63 with imposed Dirichlet conditions. We denote the subdomains and their number by 64  $\Omega^p$  and s, respectively. The gluing conditions require continuity of the displacements 65 and of their normal derivatives across the subdomain interfaces. The procedure is the 66 same as that for the static problem, [1].

Using finite element discretization in space we get the following semidiscrete 68 problem at time  $\tau$ 

$$\mathbf{M}\ddot{\mathbf{u}} + \mathbf{K}\mathbf{u} = \mathbf{f} - \mathbf{B}_{I}^{T} \boldsymbol{\lambda}_{I}^{T} - \mathbf{B}_{E}^{T} \boldsymbol{\lambda}_{E}, \tag{1}$$

$$\mathbf{B}_I \mathbf{u} \leq \mathbf{c}_I, \quad \mathbf{B}_E \mathbf{u} = \mathbf{c}_E, \quad \boldsymbol{\lambda}_I \geq \mathbf{o}, \quad \boldsymbol{\lambda}^T (\mathbf{B} \mathbf{u} - \mathbf{c}) = 0,$$
 (2)

with the discrete Newton equation of motion (1) and the equality and inequality constraints (2) resulting from the gluing, Dirichlet, and non-interpenetration conditions 71 enforced by Lagrange multipliers.

The TFETI based finite element semi-discretization in space of the subdomains 73  $\Omega^p$ ,  $p=1,\ldots,s$ , results in the block diagonal stiffness matrix  $\mathbf{K}=\mathrm{diag}(\mathbf{K}_1,\ldots,\mathbf{K}_s)$  74

of the order n with the sparse positive semidefinite diagonal blocks  $\mathbf{K}_p$  that correspond to the subdomains  $\Omega^p$ . The same structure has a positive definite mass matrix 76  $\mathbf{M} = \operatorname{diag}(\mathbf{M}_1, \dots, \mathbf{M}_s)$ . The decomposition induces also the block structure of the 77 vector of nodal forces  $\mathbf{f} = \mathbf{f}_{\tau} \in \mathbb{R}^n$  at time  $\tau$  and the vector of nodal displacements 78  $\mathbf{u} = \mathbf{u}_{\tau} \in \mathbb{R}^n$  at time  $\tau$ .

The matrix  $\mathbf{B}_I \in \mathbb{R}^{m_I \times n}$  and the vector  $\mathbf{c}_I \in \mathbb{R}^{m_I}$  describe the linearized non- 80 interpenetration conditions and the matrix  $\mathbf{B}_E \in \mathbb{R}^{m_E \times n}$  and the vector  $\mathbf{c}_E \in \mathbb{R}^{m_E}$  81 enforce the prescribed zero displacements on the part of the boundary with imposed 82 Dirichlet condition and the continuity of the displacements across the auxiliary interfaces.

Finally,  $\lambda_I \in \mathbb{R}^{m_I}$  and  $\lambda_E \in \mathbb{R}^{m_E}$  denote the components of the vector of Lagrange 85 multipliers  $\lambda = \lambda_{\tau} \in \mathbb{R}^m$ ,  $m = m_I + m_E$  at time  $\tau$ . We use the notation

$$\lambda = \begin{bmatrix} \lambda_I \\ \lambda_E \end{bmatrix}, \quad \mathbf{B} = \begin{bmatrix} \mathbf{B}_I \\ \mathbf{B}_E \end{bmatrix}, \quad \text{and} \quad \mathbf{c} = \begin{bmatrix} \mathbf{c}_I \\ \mathbf{c}_E \end{bmatrix}.$$
 (3)

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For the time discretization, we use the contact-stabilized Newmark scheme intro- 87 duced by Krause and Walloth [9] with the regular partition of the time interval [0, T], 88  $0 = \tau_0 < \tau_1 \dots < \tau_{n_T} = T$ ,  $\tau_k = k\Delta$ ,  $\Delta = T/n_T$ ,  $k = 0, \dots, n_T$ . The scheme 89 assumes that the acceleration vector is split at time  $\tau_k$  into two components 90

$$\ddot{\mathbf{u}}_k = \ddot{\mathbf{u}}_k^{int} + \ddot{\mathbf{u}}_k^{con}, \quad \ddot{\mathbf{u}}_k^{int} = \mathbf{M}^{-1} (\mathbf{f}_k - \mathbf{K} \mathbf{u}_k), \text{ and } \ddot{\mathbf{u}}_k^{con} = -\mathbf{M}^{-1} \mathbf{B}^T \boldsymbol{\lambda}_k. \tag{4}$$

We obtain the solution algorithm in the form

### Algorithm 2.1 Contact-stabilized Newmark algorithm.

Step 0. {Initialization} Set  $\mathbf{u}_0$ ,  $\hat{\mathbf{u}}_0$ ,  $\hat{\mathbf{K}} = \frac{4}{\Delta^2}\mathbf{M} + \mathbf{K}$ , T > 0,  $n_T \in \mathbb{N}$ , and  $\Delta = T/n_T$ . 94

Set 
$$\mathbf{u}_0$$
,  $\tilde{\mathbf{K}} = \frac{4}{\Delta^2}\mathbf{M} + \mathbf{K}$ ,  $T > 0$ ,  $n_T \in \mathbb{N}$ , and  $\Delta = T/n_T$ .

for 
$$k = 0, ..., n_T - 1$$
 do

Step 1. {Predictor displacement computation} 97

$$\min \left[ \frac{1}{2} \left( \mathbf{u}_{k+1}^{pred} \right)^T \mathbf{M} \mathbf{u}_{k+1}^{pred} - \left( \mathbf{M} \mathbf{u}_k + \Delta \mathbf{M} \dot{\mathbf{u}}_k - \mathbf{B}^T \boldsymbol{\lambda}_k^{pred} \right)^T \mathbf{u}_{k+1}^{pred} \right]$$
subject to  $\mathbf{B}_I \mathbf{u}_{k+1}^{pred} \leq \mathbf{c}_I$ , and  $\mathbf{B}_E \mathbf{u}_{k+1}^{pred} = \mathbf{c}_E$ 

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subject to 
$$\mathbf{B}_I \mathbf{u}_{k+1}^{pred} \leq \mathbf{c}_I$$
, and  $\mathbf{B}_E \mathbf{u}_{k+1}^{pred} = \mathbf{c}_E$ 

Step 2. {Contact-stabilized displacement computation} 100

$$\min \left[ \frac{1}{2} \mathbf{u}_{k+1}^T \widetilde{\mathbf{K}} \mathbf{u}_{k+1} - \left( \frac{4}{\Delta^2} \mathbf{M} \mathbf{u}_{k+1}^{pred} - \mathbf{K} \mathbf{u}_k + \mathbf{f}_k + \mathbf{f}_{k+1} - \mathbf{B}^T \boldsymbol{\lambda}_k \right)^T \mathbf{u}_{k+1} \right]$$
subject to  $\mathbf{B}_I \mathbf{u}_{k+1} < \mathbf{c}_I$  and  $\mathbf{B}_E \mathbf{u}_{k+1} = \mathbf{c}_E$  102

$$\dot{\mathbf{u}}_{k+1} = \dot{\mathbf{u}}_k + \frac{2}{\Delta} \left( \mathbf{u}_{k+1} - \mathbf{u}_{k+1}^{pred} \right)$$
 104

end 105

The matrix  $\mathbf{K}$  introduced in Step 0 is called an effective stiffness matrix. Let us 106 note that we omit the factor '1/2' in the term  $\mathbf{B}^T \boldsymbol{\lambda}_k^{pred}$  in the predictor step. 107

# 3 Optimal Solver with Bound on the Condition Number of the **Hessian of the Dual Energy Function**

The favorable distribution of the spectrum of the mass matrix M is sufficient to 110 implement Step 1 by using the dual theory and the standard MPRGP algorithm described in [3] with asymptotically linear complexity. To develop an optimal algorithm 112 for Step 2, we shall distinguish two cases. If the time steps are sufficiently short, then 113 the effective stiffness matrix can be considered as a perturbation of the well conditioned mass matrix, so it is enough to use again our MPRGP algorithm to prove 115 the numerical scalability and demonstrate it by numerical experiments. On the other 116 hand, if we use longer time steps, the effective stiffness matrix has very small eigen- 117 values which obviously correspond to the eigenvectors that are near the kernel of **K**. 118 This observation was fully exploited for linear problems by Farhat et al. [5] who used 119 the conjugate projectors to the natural coarse grid to achieve scalability with respect 120 to the time step. Unfortunately, this idea can not be applied in full extent to the contact problems as we do not know a priori which boundary conditions are applied to 122 the subdomains associated with the contact interface. However, we can still define 123 the preconditioning by the trace of the rigid body motions on the artificial subdomain 124 interfaces. To implement this observation, we use our preconditioning by conjugate 125 projector for partially constrained strictly convex quadratic programming problems 126 of the form

$$\min_{\lambda} \frac{1}{2} \lambda^T \widetilde{\mathbf{F}} \lambda - \lambda^T \mathbf{d} \text{ subject to } \lambda_{\mathscr{I}} \ge \mathbf{o}$$
 (5)

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which arises directly from the application of the dual theory on the problem in Step 128 2 of Algorithm 2.1. Such a method complies with our MPRGP-P algorithm for the 129 solution of strictly convex bound constrained problems described in [3]. We keep the 130 iterations in the subspace with the solution which is defined by the trace of the rigid 131 body motions on the artificial interfaces between subdomains excluding the contact 132 interface. Even though the necessity to keep the coarse grid away from the contact 133 interface prevented us from proving the optimality with respect to the time step, we 134 give the proof of optimality of our algorithm provided the ratio of the time step and 135 the space discretization parameter is kept uniformly bounded and show that the optimality can be observed by numerical experiments (see [2] for details). Moreover, 137 MPRGP-P algorithm has the rate of convergence in terms of the norm of the pro- 138 jected gradient and the bound on the condition number of the Hessian matrix of the cost functional. Therefore all we need to guarantee optimality is a uniform bound on 140 the condition number of the Hessian.

In [2], we used the standard arguments to prove the following lemma which gives 142 the required bound.

**Lemma 1.** Let  $B_1 \|\mathbf{\lambda}\|^2 \leq \|\mathbf{B}^T \mathbf{\lambda}\|^2 \leq B_2 \|\mathbf{\lambda}\|^2$  and let the elements have a regular 144 shape and size. Then 145

$$C_1 \frac{h^2 \Delta^2}{h^d (h^2 + \Delta^2)} \|\boldsymbol{\lambda}\|^2 \le \boldsymbol{\lambda}^T \widetilde{\mathbf{F}} \boldsymbol{\lambda} \le C_2 \frac{\Delta^2}{h^d} \|\boldsymbol{\lambda}\|^2, \tag{6}$$

with constants  $B_1$ ,  $B_2$ ,  $C_1$ , and  $C_2$  independent of h, H, and  $\Delta$ . Moreover, if C > 0 is 146 any constant, then for any  $0 < \Delta \le Ch$  the condition number  $\kappa(\widetilde{\mathbf{F}})$  satisfies  $\kappa(\widetilde{\mathbf{F}}) \le 147$   $\frac{C_2}{C_1}(1+C^2)$ .

# **4 Numerical Experiments**

The described algorithms were implemented in MatSol library [8] developed in Matlab environment and tested on the solution of 3D frictionless transient contact problems. For all computations we used the HP Blade system, model BLc7000 and as parallel programming environment we used Matlab Distributed Computing Engine. Sall the computations were carried out with the relative stopping tolerance  $\varepsilon = 10^{-4}$ . 154

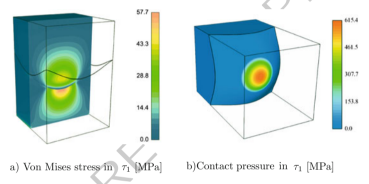
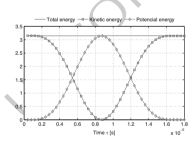
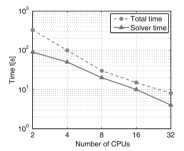


Fig. 1. Results of 3D benchmark



**Fig. 2.** Energy conservation  $(\text{ton} \cdot \text{mm}^2 \cdot \text{s}^{-2})$ 



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Fig. 3. Parallel scalability

#### 3D impact problem

Our first academic benchmark is a 3D impact between the curved 3D elastic boxes of size 10 (mm) depicted in Fig. 1. Material constants are defined by the Young modulus  $E = 2.1 \cdot 10^5$  (MPa), the Poisson ratio v = 0.3, and the density  $\rho = 7.85 \cdot 10^{-9}$  to 158 (ton/mm<sup>3</sup>). The initial gap between the curved boxes is set to 0.001 (mm). We prescribe the initial velocity -1,000 (mm/s) on the upper body in the  $x_3$  direction. The 160

upper body is floating in space and the lower body is fixed along the bottom side. The 161 linearized non-interpenetration condition was imposed on the contact interface. For 162 the time discretization, we use Algorithm 2.1 with the constant time step  $\Delta = 4 \cdot 10^{-7}$ and solve the impact of bodies in the time interval  $\tau = [0.45\Delta]$ .

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The von Mises stress distribution and the normal contact pressure along the contact interface in time  $\tau_1 = 22\Delta$  are depicted in Figs. 1a, b, respectively. The energy 166 development is shown in Fig. 2. We can see the constant total energy curve as expected.

In Table 1, we report the numerical scalability of our algorithm for the constant 169 time step  $\Delta_1 = 1 \cdot 10^{-3}$  and  $\Delta_2 = 1 \cdot 10^{-5}$  and with or without conjugate projectors. 170 We kept H/h = 10. Moreover, in last two lines of the table, we report the same 171 characteristics but with the time step dependent on the discretization step h, i.e., 172  $\Delta_{1,h} = 3h\Delta_1$ .

We can observe that the number of matrix-vector multiplications, the most expensive component of our algorithm, stays constant for the smaller time step  $\Delta_2$  as 175 expected and increases only mildly in agreement with the theory for the case of the 176 larger time step  $\Delta_1$  if we use conjugate projectors. If we simultaneously choose the 177 time step  $\Delta$  proportional to h, i.e.,  $\Delta = \Delta_h$ , then the number of matrix-vector multiplications stays the same as predicted by the theory.

Parallel scalability of our algorithm is depicted in Fig. 3, where we keep the number of elements fixed and increase the number of CPUs (subdomains).

Number of subdo	mains	16	54	128	250
Primal variables		196 608	663 552	1 572 864	3 072 000
Dual variables	21 706	81 652	214 699	443 920	
		I	Hessian n	nultiplication	ons
MPRGP	$\Delta_1$	67	86	113	191
MPRGP - P	$\Delta_1$	60	67	85	112
MPRGP	$\Delta_2$	39	40	40	42
MPRGP - P	$\Delta_2$	40	40	40	42
MPRGP	$\Delta_{1,h}$	67	72	76	78
MPRGP - P	$\Delta_{1,h}$	60	63	67	69

**Table 1.** Numerical scalability of 3D impact problem -  $\Delta$  constant or dependent on h

#### Impact of three bodies

We have also tested our algorithms on the impact of three bodies. We considered the 183 transient analysis of three elastic bodies in mutual contact (see Fig. 4). We prescribe 184 the initial velocity 5,000 (mm/s) on the sphere in the  $x_1$  direction. The L-shape body 185 is fixed along the bottom side. Material constants are defined by the Young mod- 186 ulus  $E = 2.1 \cdot 10^3$  (MPa), the Poisson ratio v = 0.3, and the density  $\rho = 6 \cdot 10^{-9}$  187 (ton/mm<sup>3</sup>). For the time discretization, we use the constant time step  $\Delta = 1 \cdot 10^{-3}$  (s) 188 and solve the impact of bodies in the time interval  $\tau = [0, 150\Delta]$  (s). The total displacement in times  $\tau_1 = 20\Delta$  and  $\tau_2 = 80\Delta$  (s) of the problem discretized by  $1.2 \cdot 10^5$  190 primal and  $8.5 \cdot 10^3$  dual variables and decomposed into 32 subdomains using METIS 191 is depicted in Fig. 4.

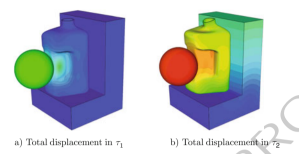


Fig. 4. Impact of bodies in time

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# **Model of Imperfect Interfaces in Composite Materials** and Its Numerical Solution by FETI Method

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Summary. Analysis of material interfaces in composite materials is in the center of attention of many material engineers. The material interface influences significantly the overall 8 behaviour of composite materials. While the perfect bond on material interface is modelled 9 without larger difficulties, the imperfect bond between different components of composite materials still causes some obstacles. This contribution concentrates on application of the FETI 11 method to description of the imperfect bond.

1 Introduction 13

The overall behavior of the engineering materials and structures is significantly affected or even dominated by the presence of interfaces, i.e. internal boundaries arising from material discontinuities. Therefore, considerable research efforts within the engineering community have been focused to adequately describe and simulate the 17 interfacial behavior under general loading conditions. A successful approach to this 18 problem is offered by the cohesive zone concept published in reference [3], in which the bulk material is assumed to be damage-free, whereas the interface response is 20 described by means of inelastic damage law. The interface model itself is formulated 21 in terms of displacement jumps and cohesive tractions bridging the interface, with 22 the elastic stiffness as the basic constitutive parameter. Initially, the stiffness is set 23 to a large value (modeling almost perfect bonding) that gradually decreases with in- 24 creasing load. For the standard displacement-based finite element approximations, 25 this gives a rise to numerical difficulties manifested in oscillations of interfacial trac- 26 tions for stiff interfaces and non-physical penetration of adjacent bodies for imperfect 27 bonding. The purpose of this contribution is to demonstrate that these limitations can 28 be overcome by duality solvers based on FETI method.

### 2 Interface Model

The constitutive description adopted in this work is based on the Ortiz-Pandolfi 31 model proposed in [7]. Detailed description of the model of the imperfect material 32

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interface can be found in reference [2]. The model is based on three state variables, 33 namely the domain displacement field,  $\mathbf{u}^{(j)}(\mathbf{x})$ , the interfacial displacement jump, 34  $[\![\mathbf{u}^{(i,j)}]\!](\mathbf{x})$ , and the interfacial damage parameter,  $\omega^{(i,j)}(\mathbf{x})$ . The superscript (j) denotes the subdomain number while the two superscripts (i,j) denote the interface 36 between the i-th and j-th subdomains.

The kinematics of the interface is quantified by the normal and tangential component of the displacement jump, provided by

$$[\![u_n^{(i,j)}]\!](\mathbf{x}) = [\![\mathbf{u}^{(i,j)}]\!](\mathbf{x}) \cdot \mathbf{n}^{(j)}(\mathbf{x}), \tag{1}$$

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where  $\mathbf{n}^{(j)}(\mathbf{x})$  denotes the normal vector and the tangential component is in the form 40

$$[\![\mathbf{u}_t^{(i,j)}]\!](\mathbf{x}) = [\![\mathbf{u}^{(i,j)}]\!](\mathbf{x}) - [\![\mathbf{u}_n^{(i,j)}]\!](\mathbf{x})\mathbf{n}^{(j)}(\mathbf{x}). \tag{2}$$

Note that the non-penetration condition hold, i.e. the normal component must remain 41 non-negative. Following [3], these quantities are combined into an effective opening 42

$$\delta(\mathbf{x}, [\![\mathbf{u}^{(i,j)}]\!](\mathbf{x})) = \sqrt{[\![u_n^{(i,j)}]\!]^2(\mathbf{x}) + \beta^2 |\![\![\mathbf{u}_t^{(i,j)}]\!](\mathbf{x})|\!|^2}$$
(3)

in which  $\beta$  denotes a constitutive parameter, also called the mode mixity parameter, 43 to be determined. This gives rise to an equivalent effective traction,  $\sigma$ , see [7]. In 44 addition, the state of an interface is quantified by an internal damage variable,  $\omega$ , 45 with  $\omega(\mathbf{x})=0$  corresponding to a perfect bonding at  $\mathbf{x}$ , whereas  $\omega(\mathbf{x})=1$  indicates 46 a fully damaged interface point.

In order to assemble the functional of energy, several energy densities are needed. 48 The density of internal energy has the form 49

$$e_{vol}^{(j)}(\mathbf{x}, \mathbf{u}^{(j)}(\mathbf{x})) = \frac{1}{2} \left( \varepsilon(\mathbf{u}^{(j)}(\mathbf{x})) \right)^T \mathbf{D} \varepsilon(\mathbf{u}^{(j)}(\mathbf{x})), \tag{4}$$

where  $\varepsilon^{(j)}(\mathbf{u}^{(j)}(\mathbf{x}))$  denotes the strain, **D** denotes the stiffness matrix of the material. 50 The internal energy functional can be written as

$$E_{vol}^{(j)}(\mathbf{u}^{(j)}(\mathbf{x})) = \int_{\Omega^{(j)}} e_{vol}^{(j)}(\mathbf{x}, \mathbf{u}^{(j)}(\mathbf{x})) d\Omega. \tag{5}$$

The potential energy of external forces has the form

$$E_{ext}^{(j)}(\mathbf{u}^{(j)}(\mathbf{x}),t) = -\int_{\Omega^{(j)}} \mathbf{u}^{(j)}(\mathbf{x}) \cdot \mathbf{b}(\mathbf{x},t) d\Omega - \int_{\Gamma^{(j)}} \mathbf{u}^{(j)}(\mathbf{x}) \cdot \mathbf{t}(\mathbf{x},t) d\Gamma, \tag{6}$$

where  $\mathbf{b}(\mathbf{x},t)$  denotes the vector of volume forces,  $\mathbf{t}(\mathbf{x},t)$  denotes the vector of surface 53 traction and  $\Gamma_l^{(j)}$  is the part of the boundary of the j-th subdomain where the surface 54 tractions are prescribed. The energy-based description involves the stored energy 55 function defined as

$$e_{int}(\mathbf{x}, [\![\mathbf{u}]\!](\mathbf{x}), \omega(\mathbf{x})) = \frac{1}{2} \frac{G}{\Delta^2} \frac{1 - \omega(\mathbf{x})}{\omega(\mathbf{x})} \delta^2, \tag{7}$$

where  $\Delta$  is the critical interface opening and G is the fracture toughness of an interface. This form is consistent with the linear softening law drawn in Fig. 1. Note that the stiffness associated with a partially damaged interface with the damage parameter,  $\omega$ , is obtained as a slope of the line 0A. The energy dissipated by changing the internal variable from  $\omega_1$  to  $\omega_2$  is given by

$$d = \begin{cases} G(\mathbf{x})(\omega_2(\mathbf{x}) - \omega_1(\mathbf{x})) & \forall \mathbf{x} \in \Gamma_{int} : \omega_1(\mathbf{x}) \le \omega_2(\mathbf{x}), \\ \infty & otherwise, \end{cases}$$
(8)

where the term  $\infty$  refers to the fact that the damage variable cannot decrease during 62 the loading process. The interfacial dissipation distance is defined 63

$$D(\omega_1(\mathbf{x}), \omega_2(\mathbf{x})) = \int_{\Gamma_{int}} d(\mathbf{x}, \omega_1(\mathbf{x}), \omega_2(\mathbf{x})) d\Gamma.$$
 (9)

The interfacial energy functional has the form

$$E_{int}(\llbracket \mathbf{u} \rrbracket(\mathbf{x}), \omega(\mathbf{x})) = \int_{\Gamma_{int}} e_{int}(\mathbf{x}, \llbracket \mathbf{u} \rrbracket(\mathbf{x}), \omega(\mathbf{x})) d\Gamma, \tag{10}$$

where  $\Gamma_{int}$  denotes the interface between subdomains.

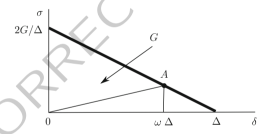


Fig. 1. Interfacial constitutive law

The description of the material interface is based on incremental solution where 66 the state variables at the k-th step  $\mathbf{u}_{k-1}(\mathbf{x})$ ,  $[\![\mathbf{u}]\!]_{k-1}(\mathbf{x})$ ,  $\omega_{k-1}(\mathbf{x})$  are known. Then, 67 the energy functional has the form

$$\Pi_k(\mathbf{u}(\mathbf{x}), [\![\mathbf{u}]\!](\mathbf{x}), \omega(\mathbf{x})) = \sum_{j=1}^n E_{vol}^{(j)}(\mathbf{u}^{(j)}(\mathbf{x})) +$$
(11)

$$\sum_{i=1}^{n} E_{ext}^{(j)}(\mathbf{u}^{(j)}(\mathbf{x})) + E_{int}(\llbracket \mathbf{u} \rrbracket(\mathbf{x}), \boldsymbol{\omega}(\mathbf{x})) + D(\boldsymbol{\omega}_{k-1}(\mathbf{x}), \boldsymbol{\omega}(\mathbf{x}))$$

and the following minimization problem is solved

$$(\mathbf{u}_k(\mathbf{x}), [\![\mathbf{u}]\!]_k(\mathbf{x}), \omega_k(\mathbf{x})) = \arg\min_{(\mathbf{u}(\mathbf{x}), [\![\mathbf{u}]\!](\mathbf{x}), \omega(\mathbf{x}))} \Pi_k(\mathbf{u}(\mathbf{x}), [\![\mathbf{u}]\!](\mathbf{x}), \omega(\mathbf{x})). \tag{12}$$

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The discretization of displacements and strains has the form

$$\mathbf{u}^{(j)}(\mathbf{x}) \approx \mathbf{u}_h^{(j)}(\mathbf{x}) = \mathbf{N}_{uh}^{(j)}(\mathbf{x})\mathbf{u}_h^{(j)},\tag{13}$$

$$\varepsilon^{(j)}(\mathbf{x}) \approx \varepsilon_h^{(j)}(\mathbf{x}) = \mathcal{B}_{u,h}^{(j)}(\mathbf{x})\mathbf{u}_h^{(j)},\tag{14}$$

where  $\mathbf{N}_{u,h}^{(j)}(\mathbf{x})$  denotes the matrix of basis functions and  $\mathcal{B}_{u,h}^{(j)}(\mathbf{x})$  denotes the strain-displacement matrix. The displacement jump is discretized in the form

$$[\![\mathbf{u}^{(i,j)}]\!](\mathbf{x}) \approx [\![\mathbf{u}_h^{(i,j)}]\!](\mathbf{x}) = \mathbf{N}_{[\![\mathbf{u}]\!],h}^{(i,j)}(\mathbf{x})[\![\mathbf{u}^{(i,j)}]\!]_h$$
(15)

and the damage parameter can be expressed

$$\boldsymbol{\omega}^{(i,j)}(\mathbf{x}) \approx \boldsymbol{\omega}_h^{(i,j)}(\mathbf{x}) = \mathbf{N}_{\boldsymbol{\omega},h}^{(i,j)}(\mathbf{x}) \boldsymbol{\omega}_h^{(i,j)}. \tag{16}$$

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After discretization, the functional of energy (11) has the form

$$\Pi_{k}(\mathbf{u}_{h}, [\![\mathbf{u}]\!]_{h}, \boldsymbol{\omega}_{h}) = \frac{1}{2} \sum_{j=1}^{n} \mathbf{u}_{h}^{(j)T} \mathbf{K}^{(j)} \mathbf{u}_{h}^{(j)} - \sum_{j=1}^{n} \mathbf{u}_{h}^{(j)T} \mathbf{f}_{h}^{(j)} + \frac{1}{2} [\![\mathbf{u}]\!]_{h}^{T} \mathbf{K}_{int}(\boldsymbol{\omega}_{h}) [\![\mathbf{u}]\!]_{h} + \boldsymbol{\omega}_{h}^{T} \mathbf{p}_{h}, \tag{17}$$

where the stiffness matrix has the classical form

$$\mathbf{K}^{(j)} = \int_{\Omega^{(j)}} \mathcal{B}_{u,h}^{(j)}^T \mathbf{D} \mathcal{B}_{u,h}^{(j)} \mathrm{d}\Omega$$
 (18)

and the vector of prescribed forces is defined as

$$\mathbf{f}_{h}^{(j)} = \int_{\Omega^{(j)}} \mathbf{N}_{u,h}^{(j)T}(\mathbf{x}) \mathbf{b}(\mathbf{x}) d\Omega + \int_{\Gamma_{t}^{(j)}} \mathbf{N}_{u,h}^{(j)T}(\mathbf{x}) \mathbf{t}(\mathbf{x},t) d\Gamma.$$
(19)

The stiffness matrix of the interface has the form

$$\mathbf{K}_{int}(\boldsymbol{\omega}_h) = \int_{\Gamma_{int}} \frac{G}{\Delta^2} \left( \frac{1}{\mathbf{N}_{\omega,h}(\mathbf{x})\boldsymbol{\omega}_h} - 1 \right) \mathbf{N}_{\llbracket u \rrbracket,h}^T(\mathbf{x}) \boldsymbol{\beta} \mathbf{N}_{\llbracket u \rrbracket,h}(\mathbf{x}) d\Gamma$$
(20)

and the vector  $\mathbf{p}_h$  is expressed as

$$\mathbf{p}_h = \int_{\Gamma_{int}} G(\mathbf{x}) \mathbf{N}_{\omega,h}(\mathbf{x}) d\Gamma. \tag{21}$$

The minimization (12) is done by the alternate minimization approach which can 79 be written as

$$(\mathbf{u}_{k}(\mathbf{x}), [\![\mathbf{u}]\!]_{k}(\mathbf{x}), \omega_{k}(\mathbf{x})) = \arg\min_{\omega(\mathbf{x})} \left( \min_{(\mathbf{u}(\mathbf{x}), [\![\mathbf{u}]\!](\mathbf{x}))} \Pi_{k}(\mathbf{u}(\mathbf{x}), [\![\mathbf{u}]\!](\mathbf{x}), \omega(\mathbf{x})) \right). \tag{22}$$

The minimization with respect to  $\mathbf{u}(\mathbf{x})$  and  $[\![\mathbf{u}(\mathbf{x})]\!]$  is associated with the Lagrangian 81 function in the form

**Page 358** 

$$L_{k,h}(\mathbf{u}_h, [\![\mathbf{u}]\!]_h, \lambda_h) = \frac{1}{2} \sum_{j=1}^n \mathbf{u}_h^{(j)T} \mathbf{K}^{(j)} \mathbf{u}_h^{(j)} - \sum_{j=1}^n \mathbf{u}_h^{(j)T} \mathbf{f}_h^{(j)} +$$

$$+ \frac{1}{2} [\![\mathbf{u}]\!]_h^T \mathbf{K}_{int}(\boldsymbol{\omega}_h) [\![\mathbf{u}]\!]_h + \lambda_h^T (\mathbf{B}_h \mathbf{u}_h - [\![\mathbf{u}]\!]_h).$$
(23)

Note that the displacement jumps  $[\![\mathbf{u}]\!]_h$  are subject to the non-penetration condition 83  $\mathbf{B}_h[\![\mathbf{u}]\!]_h \geq 0$ . In the current implementation, these constraints are converted to equalities by adopting a simple active set strategy based on the values of the Lagrange 85 multipliers  $\lambda_h$ . There are three stationary conditions 86

$$\frac{\partial L_{k,h}}{\partial \mathbf{u}_{h}^{(j)}} = \mathbf{K}^{(j)} \mathbf{u}_{h}^{(j)} - \mathbf{f}_{h}^{(j)} + \mathbf{B}_{u,h}^{(j)}^{T} \lambda_{h} = \mathbf{0}, \tag{24}$$

$$\frac{\partial L_{k,h}}{\partial \lambda_h} = \sum_{i=1}^n \mathbf{B}_{u,h}^{(j)} \mathbf{u}_h^{(j)} - [\![\mathbf{u}]\!]_h = \mathbf{0}, \tag{25}$$

$$\frac{\partial L_{k,h}}{\partial \|\mathbf{u}\|_h} = \mathbf{K}_{int}(\omega_h) [\![\mathbf{u}]\!]_h - \lambda_h = \mathbf{0}. \tag{26}$$

Equation (24) is the equilibrium equation for the *j*-th subdomain, (25) expresses the 87 interface conditions and (26) defines the relationship between the Lagrange multipliers and the displacement jumps on the interface.

3 FETI Method 90

This section summarizes the notation and the basic relationships of the FETI method 91 which is a non-overlapping domain decomposition method. More details can be 92 found in references [1, 4] or [5]. The vector of unknowns is denoted by **u**, the vector 93 of prescribed forces is denoted by **f** and the stiffness matrix is denoted by **K**. Interface 94 conditions for perfect and imperfect interaction have the form 95

$$\mathbf{B}\mathbf{u} = \begin{pmatrix} \mathbf{B}_c \\ \mathbf{B}_s \end{pmatrix} \mathbf{u} = \begin{pmatrix} \mathbf{0} \\ \mathbf{s} \end{pmatrix} = \mathbf{c}, \tag{27}$$

where s denotes the jump between subdomain displacements.

After space discretization, the functional of energy has the form

$$\Pi = \Pi(\mathbf{u}, \lambda) = \frac{1}{2} \mathbf{u}^T \mathbf{K} \mathbf{u} - \mathbf{u}^T \mathbf{f} + \lambda^T (\mathbf{B} \mathbf{u} - \mathbf{c}),$$
(28)

where  $\lambda$  denotes the vector of Lagrange multipliers.

The interface condition and the solvability condition define the coarse problem

$$\begin{pmatrix} \mathbf{F} & \mathbf{G} \\ \mathbf{G}^T & \mathbf{0} \end{pmatrix} \begin{pmatrix} \lambda \\ \alpha \end{pmatrix} = \begin{pmatrix} \mathbf{d} - \mathbf{c} \\ \mathbf{e} \end{pmatrix}, \tag{29}$$

where the well-known notation

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$$\mathbf{F} = \mathbf{B}\mathbf{K}^{+}\mathbf{B}^{T}, \quad \mathbf{G} = -\mathbf{B}\mathbf{R}, \quad \mathbf{d} = \mathbf{B}\mathbf{K}^{+}\mathbf{f}, \quad \mathbf{e} = -\mathbf{R}^{T}\mathbf{f}$$
 (30)

In reference [6], a constitutive law for the Lagrange multipliers and the discontinuity was introduced in the form

$$\mathbf{c} = \mathbf{H}\lambda$$
, (31)

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where the compliance matrix, **H**, was defined. The coarse problem can be rewritten to the form

$$\begin{pmatrix} \mathbf{F} + \mathbf{H} & \mathbf{G} \\ \mathbf{G}^T & \mathbf{0} \end{pmatrix} \begin{pmatrix} \lambda \\ \alpha \end{pmatrix} = \begin{pmatrix} \mathbf{d} \\ \mathbf{e} \end{pmatrix}. \tag{32}$$

The system of equations (32) is solved by the modified preconditioned conjugate 106 gradient method.

Comparison of (26) and (31) reveals the following equalities

$$\mathbf{c} = [\![\mathbf{u}]\!]_h = \mathbf{H}\lambda = \mathbf{K}_{int}^{-1}(\omega_h)\lambda_h. \tag{33}$$

# 4 Numerical Examples

is used.

The proposed strategy is applied to the end-notched flexure (ENF) test and the mixed-mode flexure (MMF) test used in reference [8]. The set up of the tests is depicted in Fig. 2. The material parameters are the following: Young's modulus of elasticity E =

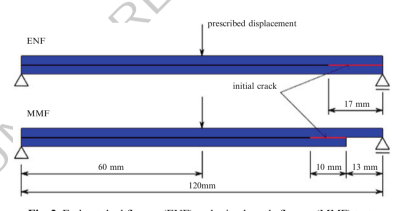


Fig. 2. End-notched flexure (ENF) and mixed-mode flexure (MMF) tests

75 GPa, Poisson's ratio v=0.3, critical stress  $\sigma_{max}=3.602$  MPa, critical opening 113  $\Delta=0.011$  mm, fracture toughness G=0.02 N/mm, mode mixity parameter  $\beta=114$  0.472. The structures are discretized by quadrilateral finite elements with bi-linear 115 basis functions. They are loaded by prescribed displacements in the center. 116

The load-deflection curves for both tests are depicted in Figs. 3 and 4 Very good agreement with results published in [8] and [7] is obtained.

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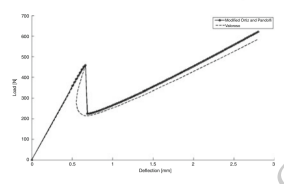


Fig. 3. Load-deflection curves for ENF test

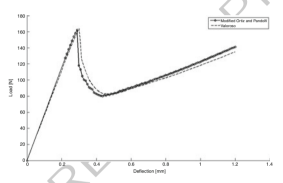


Fig. 4. Load-deflection curves for MMF test

5 Conclusions

Description of the imperfect material interface based on the compliance matrix H 120 introduced in [6] was generalized with help of the energy-based delamination model 121 described in [2]. This formulation uses piecewise constant approximation of damage 122 variables and as such it allows to express the interfacial stiffness matrix easily.

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# A Comparison of TFETI and TBETI for Numerical **Solution of Engineering Problems of Contact Mechanics**

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Summary. Since the introduction of Finite Element Tearing and Interconnecting (FETI) by 10 Farhat and Roux in 1991, the method has been recognized to be an efficient parallel technique 11 for the solution of partial differential equations. In 2003 Langer and Steinbach formulated its 12 boundary element counterpart (BETI), which reduces the problem dimension to subdomain 13 boundaries. Recently, we have applied both FETI and BETI to contact problems of mechanics. 14 In this paper we numerically compare their variants bearing the prefix Total (TFETI/TBETI) 15 on a frictionless Hertz contact problem and on a realistic problem with a given friction.

1 Introduction 17

One of the leading representatives of domain decomposition methods is the Finite 18 Element Tearing and Interconnecting (FETI) proposed by Farhat and Roux [8]. It re- 19 lies on a finite element discretization of a linear elliptic boundary value problem and 20 a nonoverlapping decomposition of the related geometric computational domain into 21 subdomains. Resulting local subproblems are glued by means of Lagrange multipli- 22 ers. The dual coarse problem is solved for the Lagrange multipliers by the method of 23 conjugate gradients. Farhat et al. [9] proved that the condition number of the Schur 24 complement, which arises from the elimination of the interior degrees of freedom, 25 preconditioned by a projector orthogonal to the kernel is proportional to H/h, where 26 H denotes the maximal subdomain diameter and h is the finite element discretization 27 parameter. Moreover, [15] proved a polylogarithmic bound on the condition num- 28 ber of the Schur complement preconditioned by the Dirichlet preconditioner. This 29 result was extended by Klawonn and Widlund [10] to the case of a redundant set of 30 Lagrange multipliers and the correct (multiplicity or stiffness) scaling.

As the Lagrange multipliers live on the skeleton of the decomposition, it is 32 very natural to employ a boundary integral representation of solutions to the local 33 subproblems. This is the Boundary Element Tearing and Interconnecting (BETI) 34 method, which was formulated and analyzed by Langer and Steinbach [13]. The 35

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resulting discretized Steklov-Poincaré operators, which relate the local Cauchy data, 36 are proved to be spectrally equivalent to the finite element Schur complements which eliminate interior degrees of freedom. An application of fully populated boundary 38 element (BE) matrices can be sparsified to a linear complexity (up to a logarithmic 39 factor), cf. [18]. Steinbach and Wendland [21] proposed a preconditioning of the BE 40 matrices by related opposite order BE operators. The latter two accelaration tech- 41 niques were exploited by Langer et al. [14] within the BETI method formulated in 42 a twofold saddle-point system. It turned to be natural to impose additional Lagrange 43 multipliers along the Dirichlet boundary, which was independently introduced as 44 Total FETI (TFETI) by Dostál et al. [6] and as All-Floating BETI by Of [16], see 45 also [17].

An extension of FETI and BETI methods to contact problems is a challenging 47 task due to the strong nonlinearity of the variational inequality under consideration. 48 To name a few of many research groups attacking this problem, see [1, 11, 20, 22]. 49 The base for our development is a theoretically supported scalable algorithm for 50 both coercive and semicoercive contact problems presented by Dostál et al. [7] and 51 in the monograph by Dostál [5]. The first scalability results using TBETI for the 52 scalar variational inequalities and the coercive contact problems were presented only 53 recently by Bouchala et al. [2, 3], respectively. We also refer to [19].

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The aim of this paper is to numerically compare TFETI and TBETI for two realistic problems. In Sect. 2 we recall the algebraic formulation of the TFETI and TBETI 56 methods for contact problems. In Sect. 3 we describe different representations of the 57 Schur complement. In Sect. 4 we compare the methods for the 3-dimensional (3d) 58 Hertz contact problem without a friction and for a 3d contact problem of a ball bearing with a given friction. In Sect. 5 we conclude.

### 2 TFETI/TBETI Formulations

Both TFETI and TBETI methods for contact problems of mechanics lead, after a 62 discretization, to the following problem: 63

$$\min_{u} \frac{1}{2} \langle Su, u \rangle - \langle f, u \rangle \text{ subject to } B_{\mathscr{I}} u \leq c_{\mathscr{I}} \text{ and } B_{\mathscr{E}} u = c_{\mathscr{E}},$$

where we search for the local boundary displacement fields  $u := (u_1, \dots, u_p)$  with 65 p being the number of subdomains. The Hessian  $S := \operatorname{diag}(S_1, \ldots, S_p)$  consists of 66 the Schur complements which are local Neumann finite element stiffness matrices 67 eliminated to subdomain boundaries in the case of TFETI, and which are symmetric boundary element discretizations of local Steklov-Poincaré operators in the case 69 of TBETI. Note that  $Ker S_i$  is the space spanned by six linearized local rigid body 70 modes. In  $f := (f_1, \dots, f_p)$  we cumulate local boundary tractions. Further,  $B_{\mathscr{E}}$  is 71 a full rank sign matrix, the first part of which interconnects teared degrees of freedom with corresponding first part of  $c_{\mathscr E}$  to be zero, while the second parts of  $B_{\mathscr E}$  73 and  $c_{\mathscr{E}}$  realize the Dirichlet boundary condition. Finally, the inequality with  $B_{\mathscr{I}}$ ,  $c_{\mathscr{I}}$  74 prescribes linearized non-penetration conditions.

Due to expensive projections onto the linear inequality constraints, we switch to 76 the dual formulation with simple bound and equality constraints 77

$$\min_{\lambda, f \geq 0} \frac{1}{2} \langle BS^{+}B^{T}\lambda, \lambda \rangle - \langle BS^{+}f - c, \lambda \rangle \text{ s.t. } (B^{T}\lambda - f) \bot \text{Ker } S,$$

where we introduce Lagrange multipliers  $\lambda := (\lambda_{\mathscr{I}}, \lambda_{\mathscr{E}})$  with  $\mathscr{I}$  and  $\mathscr{E}$  referring to 79 the inequality and equality constraints, respectively. Further, we cover  $B_{\mathscr{I}}, B_{\mathscr{E}}$  by B 80 and similarly  $c := (c_{\mathscr{I}}, c_{\mathscr{E}})$ . Let  $S^+$  be a pseudoinverse of S, i.e.,  $SS^+g = g$  for any 81  $g \perp \text{Ker } S$ . Let us denote by  $R := \text{diag}(R_1, \ldots, R_p)$  the column basis of Ker S consisting of local rigid body modes  $R_i$  and by P the orthogonal projector from Im B onto 83  $\text{Ker } R^TB^T = (\text{Ker } S)^\perp$ . To homogenize the linear (orthogonality) constraint, assume 84 we are given a feasible  $\lambda_0$  and search for  $\lambda := \tilde{\lambda} + \lambda_0$ . Returning to the old notation, 85 we arrive at the following constrained quadratic programming problem preconditioned by the projector P and regularized by the complementary projector P := I - P: 87

$$\min_{\lambda_{\mathscr{I}} \geq -(\lambda_0)_{\mathscr{I}}} \frac{1}{2} \left\langle \left( \frac{1}{\rho} PFP + Q \right) \lambda, \lambda \right\rangle - \left\langle \frac{1}{\rho} P(BS^+ f_0 - c), \lambda \right\rangle \text{ s.t. } R^T B^T \lambda = 0, \ (1)$$

where  $F := BS^+B^T$  and  $f_0 := f - B^T\lambda_0$ . Finally, we scale the cost function by  $\rho \approx 88$   $\|PFP\|$ . Now from Theorem 3.2 of [9] and from the spectral equivalence of local 89 boundary element and finite element Schur complements  $S_i$ , see Lemma 3.2 of [13], 90 we have the following optimality result valid for both TFETI and TBETI.

**Theorem 1.** Denote  $\mathcal{H} := (1/\rho)PFP + Q$ . There exist c, C > 0 independent of h, H 92 so that

$$\lambda_{\min}(\mathcal{H}|\mathrm{Im}P) \ge c\frac{h}{H}$$
 and  $\lambda_{\max}(\mathcal{H}|\mathrm{Im}P) = \|\mathcal{H}\| \le C.$  94

98

We are now in the position to use the augmented Lagrangian algorithm developed by 95 Dostál [4], see also [5], for the solution of our constraint minimization problem (1). 96 We mention that this algorithm is in some sense optimal. 97

# 3 Schur Complements

The local Schur complements  $S_i$  represent symmetric discretizations of the Steklov-Poincaré operator  $\tilde{S}_i$  mapping the Dirichlet data to the Neumann data. In particular, 100  $\tilde{S}_i(u_i) := \sigma_i(\varepsilon(\tilde{u}_i)) \cdot n_i$  in the case of elastostatics, where  $n_i$  is the outward unit normal 101 to the subdomain  $\Omega_i$ ,  $\sigma_i(\varepsilon(\tilde{u}_i))$  denotes the elastostatic stress evaluated using the 102 local linearized Hooke's law between the stress  $\sigma_i$  and the strain  $\varepsilon(\tilde{u}_i)$ , and where  $\tilde{u}_i$  103 solves the following inhomogeneous Dirichlet boundary value problem: 104

$$\operatorname{div} \sigma_i(\varepsilon(\tilde{u}_i(x))) = 0 \text{ in } \Omega_i, \quad \tilde{u}_i(x) = u_i(x) \text{ on } \partial \Omega_i.$$
 (2)

AQ1 In the case of TEETI we solve (2) approximately by the finite element method 108

In the case of TFETI we solve (2) approximately by the finite element method. 106 The approximation of  $\tilde{S}_i$  is then as follows:

$$S_i := (A_i)_{BB} - (A_i)_{BI}(A_i)_{II}^{-1}(A_i)_{IB},$$
 108

where  $(A_i)_{jk} := \int_{\Omega_i} \sigma_i(\varepsilon(\varphi_j^{(i)}(x))) : \varepsilon(\varphi_k^{(i)}(x)) dx$  is the Neumann finite element matrix assembled in the vector lowest order nodal basis functions  $oldsymbol{arphi}_i^{(i)},$  and where B and 110 I are the sets of indices of boundary and interior degrees of freedom, respectively.

In the case of TBETI the interior degrees of freedom are already eliminated in the 112 continuous formulation via a boundary integral representation of  $\tilde{u}_i(x)$  while making 113 use of the known elastostatic fundamental solution. After the lowest order Galerkin 114 boundary element discretization, we arrive at the following relation between the approximated nodal based Dirichlet data, still denoted by  $u_i$ , and the element-based 116 Neumann data, denoted by  $t_i \approx \sigma_i(\varepsilon(\tilde{u}_i)) \cdot n_i$ :

$$\begin{pmatrix} u_i \\ t_i \end{pmatrix} = \begin{pmatrix} (1/2)M_i - K_i & V_i \\ D_i & ((1/2)M_i + K_i)^T \end{pmatrix} \begin{pmatrix} u_i \\ t_i \end{pmatrix}$$
 118

with fully populated boundary element matrices  $V_i$ ,  $K_i$ , and  $D_i$ , which are referred 119 to as single-layer, double-layer, and hypersingular matrix, respectively, and with the 120 boundary mass matrix  $M_i$ . We then employ the following symmetric approximation 121 of the Schur complement  $\tilde{S}_i$ : 122

$$S_i := D_i + ((1/2)M_i + K_i)^T V_i^{-1} ((1/2)M_i + K_i).$$
123

### 4 Numerical Comparison

All the presented simulations are performed using a parallel Matlab within our Mat- 125 Sol library, see [12]. The implementations of TFETI and TBETI are consistent. The 126 only point where they differ is assembling of FEM and BEM matrices and subsequent 127 Cholesky factorizations. In the preprocessing phase times for the BEM matrices assembling dominate. Our simulations were run on a cluster of 48 cores with 2.5 GHz 129 and the infiband interface, which are equipped with licences of Matlab parallel computing engine.

First we consider a frictionless 3-dimensional Hertz problem, as depicted in 132 Fig. 1, with the Young modulus  $2.1 \cdot 10^5$  MPa and the Poisson ratio 0.3, where the 133 ball is loaded from top by the force 5,000 N. ANSYS discretization of the two bodies is decomposed by METIS into 1,024 subdomains. The comparison of TFETI 135 and TBETI in terms of computational times and number of Hessian multiplications 136 is given in Table 1. In Fig. 2 we can see a fine correspondence of contact pressures 137 computed by TFETI and TBETI to the analytical solution. The convergence criterion 138 was the decay of the dual error to  $10^{-6}$  relatively to the initial dual residuum.

In the second example we solve the contact problem of ball bearing, which 140 consists of 10 bodies. We impose Dirichlet boundary condition along the outer 141 perimeter and load the opposite part of the inner diameter with the force 4,500 N as 142 depicted in Fig. 3. The Young modulus and the Poisson ratio of the balls and rings are 143

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### Comparison of TFETI and TBETI on Contact Problems



Fig. 1. Geometry of the Hertz problem

method	number of primal DOFs		preprocessing time		number of Hessian applications	t1.1 t1.2
	4,088,832 1,849,344	926,435 926,435	21 min 1h 33 min	1 h 49 min 1 h 30 min		t1.3 t1.4

Table 1. Numerical performance of TFETI and TBETI applied to the Hertz problem

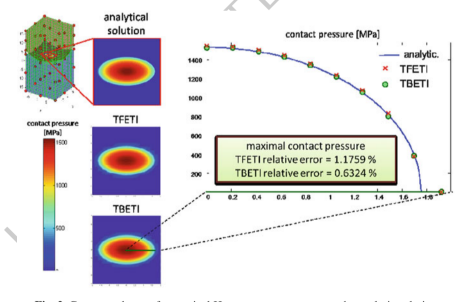


Fig. 2. Correspondence of numerical Hertz contact pressures to the analytic solution

 $2.1\cdot 10^5$  MPa and 0.3, respectively. Those of the cage are  $2\cdot 10^4$  MPa and 0.4, respectively. To get rid of the rigid body modes in the solution we introduce a small boundary gravitation term for each of the bodies. The discretized geometry was decomposed into 960 subdomains. Numerical comparison of TFETI and TBETI is shown 148 in Table 2 and the resulting vertical displacement field is depicted in Fig. 4.

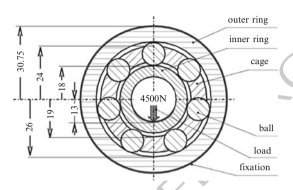


Fig. 3. Ball bearing: geometry, applied force and the Dirichlet boundary



Fig. 4. Ball bearing: vertical component of the computed displacement field

method	number of primal DOFs		preprocessing time	solution time	number of Hessian applications	t2 t2
	1,759,782	493,018	129 s	2 h 5 min	3203	t2
	1,071,759	493,018	715 s	1 h 52 min	2757	t2

Table 2. Numerical performance of TFETI and TBETI applied to the ball bearing problem

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5 Conclusion 150

In the paper we compared TFETI and TBETI and numerically documented their 151 performance for two engineering problems. Concerning timings and numbers of it- 152 erations it was shown that the methods are rather equal up to the assembling phase, 153 which is more expensive in TBETI case. On the other hand, the accuracy of the 154 boundary element discretization is usually much higher than the corresponding finite 155 element discretization. This statement is supported by the theory provided that the 156 solution is sufficiently regular. It can be also seen from Fig. 2, where one can guess 157 that the TFETI relative error of 1.1759% can be obtained with much less TBETI 158 degrees of freedom.

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### **AUTHOR QUERY**

AQ1. Please provide opening parenthesis for "div  $\sigma_i(\varepsilon(\tilde{u}_i(x)))$ " in Eq. 2.



# **FETI-DP** for Elasticity with Almost Incompressible **Material Components**

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1 Introduction 8

The purpose of this article is to present convergence bounds and some preliminary 9 numerical results for a special category of problems of compressible and almost incompressible linear elasticity when using FETI-DP or BDDC domain decomposition 11 methods.

We consider compressible and almost incompressible elasticity on the computational domain  $\Omega \subset \mathbb{R}^3$  which is partitioned into a number of subdomains. We 14 introduce nodes in the interior of the subdomains and on the interface. We distribute 15 the material parameters such that in a neighborhood of the interface we have compressible and in the interior of a subdomain we have almost incompressible linear 17 elasticity. Thus, each subdomain may contain an almost incompressible component 18 in its interior surrounded by a hull of compressible material. We will also refer to 19 this component as the incompressible inclusion.

By performing our analysis on the compressible hull, we can prove new condition 21 number bounds. Such bounds will depend on the variation of the Poisson ratio v in 22 a neighborhood of the interface of the subdomains. More precisely, for compressible 23 linear elasticity in a neighborhood of the interface and almost incompressible linear 24 elasticity in the interior of the subdomains, we can prove a polylogarithmic condition 25 number bound for the preconditioned FETI-DP system, which also depends on the 26 thickness  $\eta$  of the compressible hull.

The condition number estimate presented in this contribution is based on the theory developed in [8] for compressible linear elasticity. It can be seen as an extension 29 to certain configurations of incompressible components. For an algorithmic descrip- 30 tion of the FETI-DP method and the primal constraints applied in this paper, we refer 31 to [5, 6]. The current work can also be seen as an extension of the work of [13–15]. 32 There, the one-level FETI method for scalar elliptic problems is analyzed for special 33 cases of coefficient jumps inside subdomains.

Coarse spaces for iterative substructuring methods that are robust either with 35 respect to exact incompressibility constraints or with respect to almost incompressibility have been known for some time. For earlier work on Neumann-Neumann, 37

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FETI-DP, and BDDC methods for (almost) incompressible elasticity, see, e.g., 38 [4, 9, 10, 12].

### 2 Almost Incompressible Linear Elasticity

Let  $\Omega \subset \mathbb{R}^3$  be a polytope, which can be decomposed into smaller cubic subdomains. 41 We can allow also for subdomains that are images of cubes under a reasonable mapping. 43

The domain is fixed on  $\partial\Omega_D\subset\partial\Omega$ , i.e., we impose Dirichlet boundary conditions, and the remaining part  $\partial\Omega_N=\partial\Omega\setminus\partial\Omega_D$  is subject to a surface force g. 45 Let  $H^1_0(\Omega,\partial\Omega_D):=\left\{v\in (H^1(\Omega))^3: v\mid_{\partial\Omega_D}=0\right\}$  be the Sobolev space which is appropriate for the variational formulation. Furthermore, the linearized strain tensor 47  $\varepsilon=(\varepsilon_{ij})_{ij}$  is defined as  $\varepsilon(u)=\frac{1}{2}(\nabla u+(\nabla u)^T)$  with  $u\in (H^1(\Omega))^3$ .

Then, the linear elasticity problem is defined as follows. 49 Find the displacement  $u \in H_0^1(\Omega, \partial \Omega_D)$ , such that for all  $v \in H_0^1(\Omega, \partial \Omega_D)$ 

$$\int_{\Omega} G \, \varepsilon(u) : \varepsilon(v) \, dx + \int_{\Omega} G \, \beta \, \operatorname{div}(u) \, \operatorname{div}(v) \, dx = \langle F, v \rangle$$

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with the material parameters G,  $\beta$ , and the right hand side

$$\langle F, v \rangle = \int_{\Omega} f^T v \, dx + \int_{\partial \Omega_N} g^T v \, d\sigma.$$

The material parameters G and  $\beta$  can also be expressed using Young's modulus 53 E and the Poisson ratio  $\nu$  by  $G = \frac{E}{1+\nu}$  and  $\beta = \frac{\nu}{1-2\nu}$ . We analyze linear elasticity 54 problems with different material components. For the compressible part we use the 55 standard displacement formulation, i.e., we discretize the displacement by piecewise 56 quadratic tetrahedral finite elements.

For almost incompressible linear elasticity, i.e., when  $v \to \frac{1}{2}$ , the value of  $\beta$  tends 58 to infinity, and the discretization of the standard displacement formulation of linear 59 elasticity by low order finite elements leads to locking effects and slow convergence. 60 As a remedy the displacement problem is replaced by a mixed formulation. Therefore, we introduce the pressure  $p := G \beta \operatorname{div}(u) \in L_2(\Omega)$  as an auxiliary variable. 62

We consider the problem: Find  $(u, p) \in H_0^1(\Omega, \partial \Omega_D) \times L_2(\Omega)$ , such that

$$\int_{\Omega} G \, \varepsilon(u) : \varepsilon(v) \, dx + \int_{\Omega} \operatorname{div}(v) \, p \, dx = \langle F, v \rangle \quad \forall v \in H_0^1(\Omega, \partial \Omega_D)$$
$$\int_{\Omega} \operatorname{div}(u) \, q \, dx - \int_{\Omega} \frac{1}{G \, \beta} \, p \, q \, dx = 0 \quad \forall q \in L_2(\Omega).$$

It is well-known that in the case of almost incompressible linear elasticity, the solution of this mixed formulation exists and is unique.

For the discretization of this mixed problem we can in principle use any inf-sup 66 stable mixed finite element method. For simplicity we use  $Q_2 - P_0$  mixed finite el- 67 ements, i.e., we discretize the displacement with piecewise triquadratic hexahedral 68

finite elements and the pressure with piecewise constant elements. This discretization 69 is known to be inf-sup stable, which, in 3D, can be derived from the results in [11]. 70 To obtain again a symmetric positive definite problem, the pressure is statically con-71 densated element-by-element. We assume that a triangulation  $\tau_h$  of  $\Omega$  is given with 72 shape regular finite elements, having a typical diameter h. Additionally, we assume 73 that  $\Omega$  can be represented exactly as a union of finite elements.

The domain  $\Omega$  is now decomposed into N nonoverlapping subdomains  $\Omega_i$ , i = $1, \ldots, N$ , with diameter  $H_i$ . The resulting interface is given by  $\Gamma := \bigcup_{i \neq j} (\partial \Omega_i \cap \partial \Omega_j) \setminus \{0\}$  $\partial\Omega_D$ . We assume matching finite element nodes on the neighboring subdomains 77 across the interface  $\Gamma$ .

Then, for each subdomain we assemble the corresponding linear system

$$K^{(i)}u^{(i)} = f^{(i)}$$
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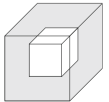
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From the local linear systems, we obtain the FETI-DP saddle point problem, 81 which is solved using a FETI-DP algorithm; see e.g., [1, 2, 5–8] for references on 82 this algorithm. In this article we consider in particular the algorithm given in [5, 6, 8]; 83 see the latter references for an algorithmic description of parallel FETI-DP methods 84 using primal edge constraints and a transformation of basis. Here, in particular, we 85 assume that all vertices are primal and all edge averages over all subdomain edges 86 are the same across the interface  $\Gamma$ .

In our analysis, each of the N subdomains may contain an almost incompressible 88 part, here also called an inclusion or a component, surrounded by a compressible 89 hull. We will specify the definitions of a hull as follows.

**Definition 1.** The hull of a subdomain  $\Omega_i$  with width  $\eta$  is defined as

$$\Omega_{i,\eta} := \{x \in \Omega_i : \operatorname{dist}(x,\partial\Omega_i) < \eta\}; \quad \text{see Fig. 1.}$$



**Fig. 1.**  $\Omega_{i,\eta}$ : hull of  $\Omega_i$ ; see Definition 1

# 3 Convergence Analysis

In this section we provide a condition number estimate for the preconditioned FETI- 94 DP matrix  $M^{-1}F$ , where F is the FETI-DP system matrix obtained from  $K^{(i)}$  and 95

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 $M^{-1}$  is the standard Dirichlet preconditioner; see [16]. We expand the convergence 96 analysis, given in [8] for compressible linear elasticity, to the case where each subdo- 97 main can contain an almost incompressible inclusion surrounded by a compressible 98 hull of thickness  $\eta$ . For the analysis, we make the following assumption; see [3] 99 where the full details are provided. 100

**Assumption 1** For each subdomain, we have an inclusion which can be either almost incompressible or compressible, surrounded by a hull  $\Omega_{i,n}$  of compressible material. The material coefficients G(x) and  $\beta(x)$  have a constant value in the interior inclusion and in the hull respectively, i.e.,

$$G(x) = \begin{cases} G_{1,i} \ x \in \overline{\Omega}_{i,\eta} \\ G_{2,i} \ x \in \Omega_i \setminus \Omega_{i,\eta} \end{cases} \qquad \beta(x) = \begin{cases} \beta_{1,i} \ x \in \overline{\Omega}_{i,\eta} \\ \beta_{2,i} \ x \in \Omega_i \setminus \Omega_{i,\eta}. \end{cases}$$
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Remark 1. Note that Assumption 1 allows that the Young modulus in the inclusion 106 can be different from the one in the hull and that their quotient can be arbitrarily 107 small or large.

The following assumption allows for the improved bound (2) in Theorem 1, 109 which contains a linear factor  $H/\eta$  compared to the factor  $(H/\eta)^4$  in (1). 110

**Assumption 2** For each subdomain  $\Omega_i$ , i = 1, ..., N, we assume that  $G_{1,i} \leq k_i \cdot G_{2,i}$ , 111 where  $k_i > 0$  is a constant independent of  $h, H, \eta, G_{1,i}$ , and  $G_{2,i}$ . 112

In the analysis provided in [3], for the edge term estimate, we need a further 113 assumption. 114

**Assumption 3** For any pair of subdomains  $(\Omega_i, \Omega_k)$  which have an edge in common, 115 we assume that there exists an acceptable path  $(\Omega_i, \Omega_{j_1}, \dots, \Omega_{j_n}, \Omega_k)$  from  $\Omega_i$  to  $\Omega_k$ , via a uniformly bounded number of other subdomains  $\Omega_{i_a}$ , q = 1, ...n, such that the coefficients  $G_{1,j_q}$  of the  $\Omega_{i_q}$  satisfy the condition

$$TOL \cdot G_{1,j_a} \geq \min(G_{1,i}, G_{1,k}), \ q = 1, \dots, n.$$

For a detailed description of the concept of acceptable paths, see [8, Sect. 5]. 120

The following theorem is proven in [3]. 121

**Theorem 1.** Under the Assumptions 1 and 3, the condition number of the preconditioned FETI-DP system satisfies 123

$$\kappa(M^{-1}F) \le C \max(1, TOL) \left(1 + \log\left(\frac{H}{h}\right)\right) \left(1 + \log\left(\frac{\eta}{h}\right)\right) \left(\frac{H}{\eta}\right)^4, \tag{1}$$

where C>0 is independent of  $h,H,\eta$ , and the values of  $G_i$  and  $\beta_i$ ,  $i=1,\ldots,N$  and hence also of  $E_i$  and  $v_i$ . 125

If additionally Assumption 2 is satisfied, we have

$$\kappa(M^{-1}F) \le C \max(1, TOL) \left(1 + \log\left(\frac{H}{h}\right)\right)^2 \left(\frac{H}{\eta}\right),$$
(2)

where C > 0 is independent of  $h, H, \eta$ , and the values of  $G_i$  and  $\beta_i$ , i = 1, ..., N and hence also of  $E_i$  and  $v_i$ . 128

#### 4 Numerical Results

In this section, we present our numerical results for a linear elasticity problem in 130 three dimensions. We consider almost incompressible inclusions in the interior of 131 the subdomains. The inclusions are always surrounded by a compressible hull with 132 v = 0.3. We use a FETI-DP algorithm with vertices and edge averages as primal 133 constraints to control the rigid body modes. For the algorithmic concept, see for 134 example [8]. The numerical results confirm our theoretical estimates.

Our tests are divided into different categories.

#### 4.1 Variable Thickness of the Compressible Hull

Here, we present results for  $3 \times 3 \times 3$  subdomains, a fixed H/h = 11, and a fixed 138 Poisson ratio v = 0.499999 in each inclusion and v = 0.3 in each hull. For these computations we vary the thickness of the hull, i.e.,  $\eta = 0, h, \dots, 5h$ ; see Table 1. 140 For the case  $\eta = 0$ , we obtain a large condition number of  $\kappa = 1,597.8$ . This is not surprising since we use a coarse space designed for compressible linear elasticity. In 142 this case using a different, larger coarse space in 3D is the remedy; see, e.g., [10] 143 or [12].

It is striking that already a hull with a thickness of one element, i.e.,  $\eta = h$ , is 145 sufficient to obtain a good condition number which is then not improved significantly 146 by further increasing  $\eta$ . As a result, the number of iteration steps does not change for 147  $\eta = h, \dots, 5h$ . In our theory, see Theorem 1, for this configuration of coefficients, our 148 bound is linear in  $H/\eta$ . From the numerical results in Table 1 we cannot conclude 149 that the bound is sharp. This might be due to the fact, that in 3D we cannot choose 150 our mesh fine enough. However, for 2D problems using very fine meshes the linear 151 dependence on  $H/\eta$  can be observed numerically; see Table 2.

iterations condition number 0 1597.8 50

**Table 1.** Growing  $\eta$ ; H/h = 11; 1/H = 3.

1h32 12.366 2h32 12.250 3h32 12.230 4h32 12.231

Growing  $\eta$  for  $3 \times 3 \times 3$  subdomains, E = 210 on the whole domain, v = 0.499999in each inclusion, and v = 0.3 in each hull. The results show only a weak dependence on  $\eta$ .

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**Table 2.** Growing  $\eta$ ; 2D; H/h = 200; 1/H = 3

η	iterations	condition number
1/100	47	199.906
2/100		102.081
3/100		70.719
4/100	36	54.674

#### 4.2 Variable Incompressibility in the Inclusions

In Table 3, we vary the Poisson ratio in the inclusions from v = 0.4 up to v = 154 0.499999 while choosing a fixed number of elements in each subdomain, i.e., H/h = 155 7, and a thickness of the hull of  $\eta = h$ . We see that the condition number is indeed 156 bounded independently of the almost incompressibility in the inclusions as expected 157 from Theorem 1.

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**Table 3.** Growing v; H/h = 7; 1/H = 3;  $\eta = h$ .

	ν	iterations	condition number
X	0.4	27	9.4841
	0.49	28	9.5038
	0.499	28	9.5063
	0.4999	28	9.5049
	0.49999	28	9.5066
	0.499999	29	9.5066

Growing v for  $3 \times 3 \times 3$  subdomains,  $\eta = h$ , v = 0.3 in the hulls, and E = 210 on the whole domain. A hull with a thickness of one element is clearly sufficient to obtain a good condition number.

# 4.3 Variable Young's Modulus in the Inclusions Combined with Variable Incompressibility in the Inclusions

In a last set of experiments, see Table 4, we consider subdomains with inclusions of 161 a high and low Young modulus, i.e., E = 1e + 4 and E = 1e - 4, either combined 162 with a Poisson ratio of v = 0.4 or v = 0.499999; see Fig. 2. The Young modulus of 163

**Page 378** 

the hull is always E = 1 and its Poisson ratio is always v = 0.3. The four different parameter settings are determined by the number of the subdomain modulo four; see 165 Fig. 2. In our theory, the condition number bound for such a configuration contains a 166 factor  $(H/\eta)^4$ . However, the results in Table 4 are not worse than in the configurations where bound (1) of Theorem 1 applies, which contains only a linear  $H/\eta$ . The 168 condition number is surprisingly low even if the thickness of the hull is only  $\eta \neq h$ . 169 While this is a favorable result it also means that it is difficult to confirm numerically 170 whether our theoretical bounds are sharp with respect to  $\eta$ .

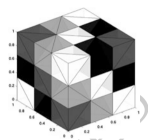


Fig. 2. Types of subdomains, see Table 4, identified by color

Table 4.	Growing	$\eta; H/$	h = 7;	1/H	= 3.
		li .			

distance $\eta$	iterations	condition number
0	> 250	13426
1 <i>h</i>	36	11.956
-2h	29	9.2575
3 <i>h</i>	29	9.4767
4 <i>h</i>	27	9.4812

Growing  $\eta$  for  $3 \times 3 \times 3$  subdomains. Four different kind of material parameter settings in the inclusions: E = 1e + 4 and v = 0.4; E = 1e - 4 and v = 0.4; E = 1e + 4and v = 0.499999; E = 1e - 4 and v = 0.499999; for all hulls: E = 1, v = 0.3.

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## An Alternative Coarse Space Method for Overlapping Schwarz Preconditioners for Raviart-Thomas Vector **Fields**

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Summary. The purpose of this paper is to introduce an overlapping Schwarz method for 7 vector field problems discretized with the lowest order Raviart-Thomas finite elements. The 8 coarse component of the preconditioner is based on energy-minimizing discrete harmonic 9 extensions and the local components consist of traditional solvers on overlapping subdomains. 10 The approach has a couple of benefits compared to the previous methods. The algorithm can 11 be implemented in an algebraic manner. Moreover, the method leads to a condition number 12 independent of the values and jumps of the coefficients across the interface between the substructures. Supporting numerical examples to demonstrate the effectiveness are also presented. 14

1 Introduction 15

Domain decomposition methods can be categorized in two classes: overlapping 16 Schwarz methods with overlapping subdomains and iterative substructuring methods 17 with nonoverlapping subdomains. In this paper, we consider two level overlapping 18 Schwarz algorithms. Such methods were originally developed for scalar elliptic prob- 19 lems; see [11, 15] and references therein. Later these methods have also been consid- 20 ered for solving vector fields problems posed in H(div) and H(curl); see [1, 9, 13]. 21 Other types of algorithms, such as multigrid methods, classical iterative substruc- 22 turing methods, balancing Neumann-Neumann, and FETI methods, have also been 23 suggested in [3, 8, 12, 14, 16, 17]. Many nonoverlapping methods have been stud- 24 ied for discontinuous coefficients cases for vector fields problems. However, only 25 few methods were introduced for the overlapping Schwarz methods in case of coef- 26 ficients which have jumps.

In the domain decomposition theory, methods can often provide good scalability, 28 i.e., the condition number of the preconditioned system will depend only on the size 29 of the subdomain problems and not on any other parameters, e.g., the number of subdomains and jumps of the coefficients. For the purpose of handling the discontinuity, 31 we borrow the advanced coarse space techniques of [6, 7] based on discrete harmonic 32 extensions of coarse trace spaces developed for almost incompressible elasticity.

27

The rest part of this paper is organized as follows. We introduce a model prob- 34 lem and its finite element approximation in Sect. 2. In Sects. 3 and 4, we recall the 35

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overlapping Schwarz method and we suggest the alternative coarse algorithm, re- 36 spectively. We next present the numerical results in Sect. 5. Finally, the conclusion 37 of this paper is given in Sect. 6.

#### 2 Discretized Problem

We consider the following second order partial differential equation for vector field 40 problem posed in H(div) in a bounded polyhedral domain  $\Omega$  with a homogeneous 41 boundary condition:

$$L\mathbf{u} := -\mathbf{grad} (\alpha \operatorname{div} \mathbf{u}) + \beta \mathbf{u} = \mathbf{f} \operatorname{in} \Omega,$$

$$\mathbf{u} \cdot \mathbf{n} = 0 \operatorname{on} \partial \Omega.$$
(1)

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Here we have positive coefficients  $\alpha, \beta \in L^{\infty}(\Omega)$  and assume that **f** is in  $(L^{2}(\Omega))^{3}$ . The main focus of our work is on the coefficients  $\alpha$  and  $\beta$  which have jumps across 44 between the substructures.

The model problem (1) has many important applications, such as a mixed and 46 least-squares formulation of certain types of second order partial differential equations [5, 17]. There are other types of applications related to H(div), e.g., iterative 48 solvers for the Reissner-Mindlin plate and the sequential regularization method for 49 the Navier-Stokes equations. For more detail, see [2, 10].

We next consider a variational formulation of (1):

$$\mathbf{a}(\mathbf{u}, \mathbf{v}) := \int_{\Omega} \alpha \operatorname{div} \mathbf{u} \operatorname{div} \mathbf{v} dx + \beta \mathbf{u} \cdot \mathbf{v} dx = \int_{\Omega} \mathbf{f} \cdot \mathbf{v} dx, \, \mathbf{v} \in H_0(\operatorname{div}; \Omega).$$
 (2)

We consider the lowest order Raviart-Thomas elements, conforming in H(div), 52 to obtain a discretized problem; see [4, Chap. 3]. We note that the degrees of freedom 53 of the Raviart-Thomas elements are defined by the average values of the normal 54 components over the faces. 55

Let us consider the variational problem (2). Restricting to the finite element space 56 of the lowest order Raviart-Thomas elements with shape regular and quasi-uniform 57 meshes, we obtain the following linear system:

$$Au = f. (3)$$

where the matrix A is a stiffness matrix, u is a vector of degrees of freedom, and f is 59 a known vector obtained from **f**. We note that A is symmetric and positive definite.

## 3 Overlapping Schwarz Preconditioner

We consider a decomposition of the domain  $\Omega$  into N nonoverlapping subdomains 62  $\Omega_i$ ,  $i=1,\cdots,N$ . We next introduce extended subregions  $\Omega_i'$  obtained from  $\Omega_i$  by 63 adding layers of elements and the interface  $\Gamma$  which is given by 64

$$\Gamma = \left(\bigcup_{i=0}^{N} \partial \Omega_{i}\right) \backslash \partial \Omega.$$
 65

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We consider a two-level overlapping Schwarz algorithm to solve the linear system (3). An overlapping Schwarz preconditioner usually has the following form:

$$P^{-1} = R_0^T A_0^{-1} R_0 + \sum_{i=1}^N R_i^T A_i^{-1} R_i, \tag{4}$$

where  $A_0$  is the matrix of the global coarse problem, the  $A_i$ 's are obtained from local 68 subproblems related to the extended subdomains  $\Omega'_i$ , and  $R_0$  and  $R_i$ 's are restriction 69 operators to the coarse space and local spaces, respectively; see [11, 15] for more 70 details.

In [9, 13], model problems were designed for constant coefficients and convex 72 domains to analyze the methods. In our work, we use more general assumptions: 73 convex subdomains and coefficients which have jumps across the interface  $\Gamma$ .

In order to deal with this situation, we consider an alternative coarse space 75 approach instead of traditional coarse interpolations. The basis functions for the 76 new algorithm are based on energy-minimizing discrete harmonic extensions with 77 given interface values. We use the corresponding discrete harmonic extensions of 78 the boundary values of standard basis functions to construct new basis functions. We 79 remark that this process can be performed locally and in parallel due to the fact that 80 the basis functions are supported in just two subdomains. We also note that we do 81 not need any coarse triangulation and this work can be done algebraically. With new 82 alternative basis functions, we obtain the operator  $R_0$  which defines the new basis 83 and the matrix  $A_0 = R_0 A R_0^T$  associated with the global coarse problem.

For the local components, we follow the traditional way. Each  $R_i$  is a rectangular 85 matrix with elements equal to 0 and 1 and provides the indices relevant to an indi- 86 vidual extended subdomain  $\Omega'_i$ . Each  $A_i = R_i A R_i^T$  is just the principal minor of the 87 original stiffness matrix A defined by  $R_i$ . By using these matrices, we can build the 88 local component  $\sum_{i=1}^{N} R_i^T A_i^{-1} R_i$  of the Schwarz preconditioner.

## 4 The Coarse Component

In this section, we explain our approach in detail. We focus on the restriction operator 91  $R_0$  onto the coarse space. Before we consider the alternative method, we introduce 92 the conventional method in [9, 13]. The restriction operator is obtained by the in- 93 terpolation from the subspaces defining the coarse component to the global space. 94 More precisely,  $R_0$  are exactly the coefficients obtained by interpolating the traditional coarse basis functions onto the fine mesh. We note that we need geometric 96 information, e.g., coordinate information, to construct  $R_0$ .

Instead of the conventional coarse basis, we will use discrete harmonic extensions to define the new coarse basis functions. We first consider two adjacent subdo- 99 mains  $\Omega_i$  and  $\Omega_j$ . We then have a coarse face  $F_{ij} = \partial \Omega_i \cap \partial \Omega_j$ . We note that each

**Page 383** 

coarse degree of freedom of our coarse component is related to each coarse face. Let u denote the vector of degrees of freedom for the original problem. Similarly, we 102 consider the vectors of degrees of freedom  $u_I^{(i)}, u_I^{(j)}$ , and  $u_{F_{ij}}$  associated with  $\Omega_i \backslash \Gamma$ , 103  $\Omega_j \backslash \Gamma$ , and  $F_{ij}$ , respectively. We then have restriction matrices  $R_I^{(i)}, R_I^{(j)}$ , and  $R_{F_{ij}}$ , i.e., 104  $u_I^{(i)} = R_I^{(i)}u$ ,  $u_I^{(j)} = R_I^{(j)}u$ , and  $u_{F_{ij}} = R_{F_{ij}}u$ . We note that each restriction matrix has 105 only one nonzero entry of unity per each row. We next introduce a submatrix of the 106 stiffness matrix A. It corresponds to the two subdomains which have  $F_{ij}$  in common: 107

$$\begin{bmatrix} A_{II}^{(i)} & 0 & A_{IF_{ij}}^{(i)} \\ 0 & A_{II}^{(j)} & A_{IF_{ij}}^{(j)} \\ A_{F_{ij}I}^{(i)} & A_{F_{ij}I}^{(j)} & A_{F_{ij}F_{ij}} \end{bmatrix}.$$

We choose  $u_{F_{ij}}^T=[1,1,\cdots,1]$  and introduce the local subproblems  $A_{II}^{(i)}u_I^{(i)}+A_{IF_{ij}}^{(i)}$  109  $u_{F_{ij}}=0$  and  $A_{II}^{(j)}u_I^{(j)}+A_{IF_{ij}}^{(j)}u_{F_{ij}}=0$  to consider discrete harmonic extensions; see [15, 110 Chap. 4.4]. Then,  $u_I^{(i)}$  and  $u_I^{(j)}$  are completely determined by  $u_{F_{ij}}$ , i.e.,  $u_I^{(i)}=E_iu_{F_{ij}}$  111 and  $u_I^{(j)}=E_ju_{F_{ij}}$ , where  $E_i:=-A_{II}^{(i)}-A_{IF_{ij}}^{(i)}$  and  $E_j:=-A_{II}^{(j)}-A_{IF_{ij}}^{(j)}$ . We then obtain 112 a coarse basis  $u_{ij}=R_I^{(i)}u_I^{(i)}+R_I^{(j)}u_I^{(j)}+R_{F_{ij}}^Tu_{F_{ij}}$  corresponding to  $F_{ij}$ . We can then 113 construct the following form of our coarse interpolation matrix  $R_0$  after the similar 114 process:

$$R_0 := \begin{bmatrix} \vdots \\ -u_{ij}^T - \\ \vdots \end{bmatrix}.$$

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As we mentioned earlier, we can obtain the coarse matrix  $A_0$  by the Galerkin product 117  $R_0AR_0^T$ . We remark that our alternative approach can be implemented in an algebraic 118 manner and in parallel. However, we need to solve additional local Dirichlet-type 119 subproblems to construct the coarse component compared to the conventional methods. 121

## **5 Numerical Experiments**

We apply the overlapping Schwarz method with the energy-minimizing coarse space to our model problem. We use  $\Omega=(0,1)\times(0,1)\times(0,1)$  and the lowest order hexahedral Raviart-Thomas elements. We decompose the domain into  $N\times N\times N$  identical subdomains. In each subdomain, we assume that the coefficients  $\alpha$  and  $\beta$  are the constant. We consider cases where the coefficients have jumps across the interface the three subdomains, in particular, a checkerboard distribution pattern. Each subdomain  $\Omega_i$  has side length H=1/N and each mesh cube has h as a minimum side length. We also introduce extended subdomains whose boundaries do not cut any last

mesh elements with an overlap parameter  $\delta$  between subdomains. We use the preconditioned conjugate gradient method to solve the preconditioned linear system

$$P^{-1}Au = P^{-1}f. (5)$$

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We stop the iteration when the residual  $l_2$ -norm has been reduced by a factor of  $10^{-6}$ . 133 We perform two different kinds of experiments. We first fix the overlap parameter 134  $H/\delta$  and vary H/h. We next fix the size of H/h and use various size of  $H/\delta$ . We 135 report the condition numbers estimated by the conjugate gradient method and the 136 number of iterations. Tables 1 and 3 show the first results and Tables 2 and 4 show 137 the results of the second experiments.

In the first set of experiments, we see that the condition numbers and the iteration counts do not depend on the size of H/h. In the second set, we can conclude 140 that the condition numbers grow linearly with  $H/\delta$ . For both cases, the condition 141 numbers and iteration counts are quite independent of coefficients and the jumps of 142 coefficients between the subdomains.

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**Table 1.** Condition numbers and iteration counts.  $\alpha_i = 1$  or specified values as indicated in a checkerboard pattern,  $\beta_i \equiv 1$ ,  $\frac{H}{\delta} = 4$ ,  $H = \frac{1}{3}$ , and  $h = \frac{1}{12}, \frac{1}{24}, \frac{1}{48}$ 

	$\alpha_i =$	0.01	$\alpha_i =$	0.1	$\alpha_i$	= 1	$\alpha_i =$	= 10	$\alpha_i =$	100	
$\frac{H}{h}$	cond	iters	cond	iters	cond	iters	cond	iters	cond	iters	
4	8.23	15	8.90	16	9.16	17	8.92	16	8.25	15	
1	8.39			~							
16	8.23	16	8.99	17	9.22	19	8.98	17	8.28	16	

**Table 2.** Condition numbers and iteration counts.  $\alpha_i = 1$  or specified values as indicated in a checkerboard pattern,  $\beta_i \equiv 1$ ,  $\frac{H}{h} = 16$ ,  $H = \frac{1}{3}$ , and  $h = \frac{1}{48}$ 

				$\alpha_i =$							
	$\frac{H}{\delta}$	cond	iters	cond	iters	cond	iters	cond	iters	cond	iters
	4	8.23	16	8.99	17	9.22	19	8.98	17	8.28	16
Ī	8	10.86	16	13.27	18	14.06	22	14.16	18	14.10	16
	16	16.22	18	22.94	22	25.03	24	25.30	22	25.32	20

6 Conclusion 144

An alternative coarse space technique based on energy-minimizing discrete harmonic 145 extensions for overlapping Schwarz algorithm for vector field problems posed in 146

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**Table 3.** Condition numbers and iteration counts.  $\beta_i = 1$  or specified values as indicated in a checkerboard pattern,  $\alpha_i \equiv 1$ ,  $\frac{H}{\delta} = 4$ ,  $H = \frac{1}{3}$ , and  $h = \frac{1}{12}$ ,  $\frac{1}{24}$ ,  $\frac{1}{48}$ 

	$\beta_i =$	0.01	$\beta_i =$	0.1	$\beta_i$ =	= 1	$\beta_i =$	= 10	$\beta_i =$	100
$\frac{H}{h}$	cond	iters								
4	8.18	15	8.36	16	9.16	17	8.68	17	8.36	16
8	8.18	17	8.46	18	9.20	18	8.65	18	8.37	18
16	8.18	17	8.45	18	9.22	19	8.62	18	8.37	18

**Table 4.** Condition numbers and iteration counts.  $\beta_i = 1$  or specified values as indicated in a checkerboard pattern,  $\alpha_i \equiv 1$ ,  $\frac{H}{h} = 16$ ,  $H = \frac{1}{3}$ , and  $h = \frac{1}{48}$ 

	$\beta_i =$	0.01	$\beta_i =$	0.1	$\beta_i =$	= 1	$\beta_i =$	10	$\beta_i =$	100
$\frac{H}{\delta}$	cond	iters								
4	8.18	17	8.45	18	9.22	19	8.62	18	8.37	18
	1		9.98				//			19
16	9.34	17	13.13	21	25.03	24	24.79	22	12.56	19

H(div) has been introduced and implemented. The numerical results show the usefulness of our method even in the presence of jumps of the coefficients between the 148 substructures.

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### **AUTHOR QUERY**

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## A Simultaneous Augmented Lagrange Approach for the Simulation of Soft Biological Tissue

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**Summary.** In this paper, we consider the elastic deformation of arterial walls as occurring, 12 e.g., in the process of a balloon angioplasty, a common treatment in the case of atherosclerosis. Soft biological tissue is an almost incompressible material. To account for this property 14 in finite element simulations commonly used free energy functions contain terms penalizing 15 volumetric changes. The incorporation of such penalty terms can, unfortunately, spoil the convergence of the nonlinear iteration scheme, i.e., of Newton's method, as well as of iterative 17 solvers applied for the solution of the linearized systems of equations. We show that the augmented Lagrange method can improve the convergence of the linear and nonlinear iteration 19 schemes while, at the same time, implementing a guaranteed bound for the volumetric change. 20 Our finite element model of an atherosclerotic arterial segment, see Fig. 1, is constructed from 21 intravascular ultrasound images; for details see [4].

Fig. 1. Finite element model of an atherosclerotic arterial segment 1.3M unknowns



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#### 1 Nonlinear Model and Algorithm

Biological tissues, such as arteries, are fiber enforced materials composed of an 24 almost incompressible matrix substance with embedded collagen fibers. The arrange- 25 ment of the fibers in arterial walls is characterized by two preferred directions heli- 26 cally wound along the artery. The material behavior of the collagen fiber bundles 27 is represented by the superposition of two transversely isotropic models; see [12]. 28 Thus, the strain energies are given by

$$\psi = \psi^{\text{iso}}(\mathbf{C}) + \psi^{\text{ti},(1)}(\mathbf{C}, \mathbf{M}^{(1)}) + \psi^{\text{ti},(2)}(\mathbf{C}, \mathbf{M}^{(2)}).$$
 (1)

Here,  $\mathbf{F} := \nabla \varphi$  is the deformation gradient,  $\mathbf{C} := \mathbf{F}^T \mathbf{F}$  the right Cauchy-Green-tensor, 30 and  $\mathbf{M}^{(a)} := \mathbf{a}^{(a)} \otimes \mathbf{a}^{(a)}$ , a = 1, 2 are the structural tensors characterizing the fiber 31 directions. There exist different possibilities to model the mechanical response of soft 32 biological tissue; see, e.g., [2, 12]. We are interested in polyconvex energy functions. 33 For the construction of anisotropic, polyconvex functions, see, e.g., [18]. Here, we 34 use the model due to [12], which was denoted model  $\psi_B$  in [3],

$$\psi = c_1 \left( I_1 I_3^{-1/3} - 3 \right) + \sum_{a=1}^{2} \frac{k_1}{2k_2} \left\{ \exp\left( k_2 \left\langle J_4^{(a)} I_3^{-1/3} - 1 \right\rangle^2 \right) - 1 \right\} 
+ \varepsilon_1 \left( I_3^{\varepsilon_2} + I_3^{-\varepsilon_2} - 2 \right)^{\alpha},$$

with the invariants  $I_1 = \operatorname{tr} \mathbf{C}, I_2 = \operatorname{tr}[\operatorname{Cof}(\mathbf{C})], I_3 = \det \mathbf{C}, J_4^{(a)} = \operatorname{tr}[\mathbf{C}\mathbf{M}^{(a)}], J_5^{(a)} = 36$  $\operatorname{tr}[\mathbf{C}^2\mathbf{M}^{(a)}]$ . Here,  $\langle \bullet \rangle$  denote the Macauly brackets,  $\langle \bullet \rangle = (|\bullet| + \bullet)/2$ . The penalty 37 term  $\varepsilon_1 \left( I_3^{\varepsilon_2} + I_3^{-\varepsilon_2} - 2 \right)^{\alpha}$  models the incompressibility. 38

We adjust our parameters to experimental results in [11]; for details, see [5]. The adjustment results in the parameters  $c_1 = 7.17 \, [\text{kPa}], k_1 = 3.69e - 3 \, [\text{kPa}], k_2 = 51.2$  40 for the adventitia and  $c_1 = 9.23 \, [\text{kPa}], k_1 = 193 \, [\text{kPa}], k_2 = 2.627e3$  for the media.

In the augmented Lagrange approach [10, 20] a Lagrange multiplier is introduced 42 on each finite element and  $\mu^T(\det \mathbf{F} - \mathbf{1})$  is added to the energy  $\psi$ . Here, we mean by 43 det F the vector of element-wise determinants of F. The Lagrange multiplier will be 44 computed iteratively by an Uzawa-like iteration  $\mu_{k+1} = \mu_k + \xi_k(\det \mathbf{F} - \mathbf{1})$ , where in 45 our computations in Sect. 3 the series  $\xi_k$  will be chosen as a constant  $\xi_k = \xi = 499.0$ . 46 We have chosen  $\xi$  by hand from the set  $\{99,499,999,1999,9999\}$ .

Our parameter fit is performed assuming incompressibility of the material. When 48 using the penalty approach we have to choose sufficiently large penalty parameters. 49 Here, our penalty parameters are  $\varepsilon_1 = 70.0 \, [\text{kPa}], \varepsilon_2 = 8.5, \alpha = 1$  for the adventitia 50 and  $\varepsilon_1 = 360.0 \, [\text{kPa}], \varepsilon_2 = 9.0, \alpha = 1$  for the media. Also in the augmented Lagrange 51 approach we need to choose our penalty parameters but here the penalty may be 52 relaxed significantly, i.e., we choose  $\varepsilon_1 = 10.0 \, [\text{kPa}], \varepsilon_2 = 4.0, \alpha = 1$  for adventitia 53 and media. The relaxation becomes evident when the penalty function is plotted for 54 the different sets of parameters. A sufficiently accurate stopping criterion has to be 55 chosen for the augmented Lagrange loop; here we chose a tolerance of  $|\det(\mathbf{F}) - 1| \le 56$ 0.01 on each element.

In our discretization, we have to avoid locking effects. We therefore replace 58 point-wise penalization by the penalization of the average volumetric change on 59 every finite element. This is accomplished, as in [3, 16], by applying a three-field 60 formulation, known as the  $\bar{\mathbf{F}}$ -approach; see [19]. We use 10-noded tetrahedral elements for the displacement.

In our nonlinear scheme we solve a sequence of linear problems obtained from 63 Newton's method, see, e.g., Fig. 2. This is also referred to as (pseudo) time stepping 64 or load stepping. To obtain a fair comparison, we have chosen an automatic time stepping strategy. For the penalty approach we increase  $\Delta t$  when the number of Newton 66 iterations is smaller than 6 and decrease  $\Delta t$  when it is larger than 9. This choice produced the best results. The simultaneous Augmented Lagrange approach, where the 68 iteration for the Lagrange multiplier simultaneously to the Newton correction, can 69 be viewed as an inexact Newton method. Thus, a quadratic convergence cannot be 70 expected. We therefore have chosen the bounds for the auto time stepping as 18 and 71 36. For all approaches the maximal time step size was bounded by  $\Delta t_{\rm max} = 0.4$ .

Fig. 2. Penalty for the incompressibility

```
Nonlinear Iteration (Penalty)  \begin{array}{l} \text{Set } k=0 \text{ and } t_0=\Delta t_0; \\ \text{Apply partial load } t_k \cdot \mathbf{f}_{\text{load}} \text{ if the full load is not yet reached;} \\ \text{Use Newton iteration to solve the nonlinear problem.} \\ \text{Use GMRES to solve linearized problem using the } \\ \text{FETI domain decomposition method as a preconditioner;} \\ \text{Apply Newton correction;} \\ \text{Adapt load step size } \Delta t_{k+1}, \text{ i.e.,} \\ \Delta t_{k+1} = 10^{1/5} \Delta t_k, \ \Delta t_{k+1} = 10^{-1/5} \Delta t_k, \text{ or } \Delta t_{k+1} = \Delta t_k;} \\ \text{Set } t_{k+1} = t_k + \Delta t_{k+1}; \\ \end{array}
```

#### 2 FETI-DP Method

We briefly introduce the well-known FETI-DP method. For a more detailed intro- 74 duction, see, e.g., [13, 16, 17, 21]. For algorithms of the Finite Element Tearing and 75 Interconnecting-type (FETI); see [6–9]. Using FETI-DP methods linear systems with 76 billions of unknowns have been solved, e.g., in [14, 16] on large parallel machines 77 (Fig. 3).

We decompose the domain  $\Omega$  into N nonoverlapping subdomains  $\Omega_i$ . For all subdomains  $\Omega_i$ , we assemble the local stiffness matrices  $\mathbf{K}^{(i)}$  and local load vectors  $\mathbf{f}^{(i)}$ , 80  $i=1,\ldots,N$ ,

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Fig. 3. Simultaneous augmented Lagrange for the incompressibility [10, 20]

#### Nonlinear Iteration (Simultaneous Augmented Lagrange)

Set k=0 and  $t_0=\Delta t_0$ ;

Apply partial load  $t_k \cdot \mathbf{f}_{load}$  if the full load is not yet reached; Set Lagrange parameter  $\mu_0 = 0$ ;

While Newton iteration has not converged and while elements with

TOL exist: Solve nonlinear problem with  $|\det(\mathbf{F}) - 1|$ simultaneous

Newton iteration and iteration for  $\mu$ 

Use GMRES to solve linearized problem using the FETI method

Apply Newton correction and update Lagrange parameter  $\mu_{k+1} = \mu_k + \xi_k(\det \mathbf{F} - 1)$ ;

Adapt load step size  $\Delta t_{k+1}$ , i.e.,  $\Delta t_{k+1} = 10^{1/5} \Delta t_k$ ,  $\Delta t_{k+1} = 10^{-1/5} \Delta t_k$ , or  $\Delta t_{k+1} = \Delta t_k$ . Set  $t_{k+1} = t_k + \Delta t_{k+1}$ ;

$$\mathbf{K} = \begin{bmatrix} \mathbf{K}^{(1)} & & \\ & \ddots & \\ & & \mathbf{K}^{(N)} \end{bmatrix}, \mathbf{u} = \begin{bmatrix} \mathbf{u}^{(1)} \\ \vdots \\ \mathbf{u}^{(N)} \end{bmatrix}, \mathbf{f} = \begin{bmatrix} \mathbf{f}^{(1)} \\ \vdots \\ \mathbf{f}^{(N)} \end{bmatrix}.$$
 83

The interface is  $\Gamma = \bigcup_{i=1}^N \partial \Omega_i \setminus \partial \Omega$ . The discrete problem can be formulated as 84 minimization problem with the interface continuity constraint  $\mathbf{B}\mathbf{u} = \mathbf{0}$ , where  $\mathbf{B} = 85$  $[\mathbf{B}^{(1)}, \dots, \mathbf{B}^{(N)}]$  with entries from 0, 1, -1. By introducing Lagrange multipliers  $\lambda$  to 86 enforce the continuity along the subdomain interface we obtain the problem: Find 87  $(\mathbf{u}, \lambda)$ , such that 88

$$\mathbf{K}\mathbf{u} + \mathbf{B}^T \lambda = \mathbf{f}$$

$$\mathbf{B}\mathbf{u} = \mathbf{0}.$$

This problem can be solved by eliminating the displacement variables **u** and solving 90 the resulting Schur complement system by conjugate gradients.

In FETI-DP methods some continuity constraints are enforced on primal dis- 92 placement variables  $\tilde{\mathbf{u}}_{\Pi}$  throughout iterations to enforce invertibility of the local 93 problems. This yields a saddle point problem of the form

$$\widetilde{\mathbf{K}}\widetilde{\mathbf{u}} + \mathbf{B}^T \lambda = \widetilde{\mathbf{f}}$$
 $\mathbf{B}\widetilde{\mathbf{u}} = \mathbf{0},$ 

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where the matrix  $\hat{\mathbf{K}}$  and right hand side  $\tilde{\mathbf{f}}$  are partially assembled in the primal variables, i.e., 97

$$\widetilde{\mathbf{K}} = \begin{bmatrix} \mathbf{K}_{BB}^{(1)} & \widetilde{\mathbf{K}}_{\Pi B}^{(1)T} \\ & \ddots & \vdots \\ & \mathbf{K}_{BB}^{(N)} \ \widetilde{\mathbf{K}}_{\Pi B}^{(N)T} \\ \widetilde{\mathbf{K}}_{\Pi B}^{(1)} \cdots \widetilde{\mathbf{K}}_{\Pi B}^{(N)} \ \widetilde{\mathbf{K}}_{\Pi \Pi} \end{bmatrix}, \qquad \widetilde{\mathbf{f}} = \begin{bmatrix} \mathbf{f}_{B}^{(1)} \\ \vdots \\ \mathbf{f}_{B}^{(N)} \\ \widetilde{\mathbf{f}}_{\Pi} \end{bmatrix}.$$

The coupling also provides the coarse problem for the method. Reducing the system  $_{99}$  of equations to an equation in  $\lambda$ , it remains to solve iteratively  $_{100}$ 

$$\mathbf{M}_{\mathrm{D}}^{-1}\mathbf{F}_{feti}\lambda = \mathbf{M}_{\mathrm{D}}^{-1}\mathbf{d},$$

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where  $\mathbf{F}_{feti} = \mathbf{B}\widetilde{\mathbf{K}}^{-1}\mathbf{B}^T$ , and  $\mathbf{M}_D^{-1} = \mathbf{B}_D\mathbf{R}_{\Gamma}^T\mathbf{S}\mathbf{R}_{\Gamma}\mathbf{B}_D^T$  is the Dirichlet preconditioner. 102 Here,  $\mathbf{S}$  is the Schur complement obtained by eliminating the interior variables in 103

every subdomain, i.e., 
$$\mathbf{S} = \begin{bmatrix} \mathbf{S}^{(1)} \\ \ddots \\ \mathbf{S}^{(N)} \end{bmatrix}$$
. The operator  $\mathbf{R}_{\Gamma}$  is a restriction matrix, 104

consisting of zeros and ones, that, when applied to a vector  $\tilde{\mathbf{u}}$ , removes the interior variables from  $\tilde{\mathbf{u}}$ . The matrices  $\mathbf{B}_D$  are scaled variants of the jump operator  $\mathbf{B}$  where, in the simplest case, the contribution from and to each interface node is scaled by the inverse of the multiplicity of the node. We define the multiplicity of a node as the number of subdomains it belongs to. For heterogeneous problems a more elaborate scaling, using an appropriate scaling factor, defined by the coefficients  $\rho_i$ , is necessary; see, e.g., [17, p. 1532, Formula (4.3)] and [15, p. 1403, Formula (6)].

#### 3 Numerical Results

A pressure of 200 mmHg is applied to the inside of the artery, see Fig. 1. The 113 FETI-DP iteration is stopped when the absolute residual is reduced to  $5 \times 10^{-9}$ ; 114 we have 224 subdomains. The total cost can be estimated by multiplying the number 115 of Newton steps by the corresponding average number of (inner) FETI-DP Krylov 116 iterations, see Tables 1 and 2.

Our results show that the use of the augmented Lagrange method can significantly improve the properties of the linearized systems occurring in the nonlinear solution scheme. The convergence of the nonlinear scheme is also improved, i.e., in our nonlinear scheme larger pseudo time steps  $\Delta t$  can be chosen. Of course, an additional iteration process for the Lagrange multiplier is introduced. Here, this iteration process is carried out simultaneously with the Newton iteration.

The results in Tables 1 and 2 show that the additional cost for the augmented 124 Lagrange iteration is more than amortized by the faster convergence of the nonlin- 125 ear scheme and the linear iterative solver. Moreover, in the augmented Lagrange 126 approach the volumetric change is exactly controlled during the iteration process, 127 i.e., we have satisfied element-wise the condition  $|\det(\mathbf{F}) - 1| \le 0.01$ . In the penalty 128 approach the volumetric change produced by the chosen penalty parameters is only 129

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**Table 1.** Newton iteration for the penalty formulation. Pseudo-time *t*, number of Newton steps, average number of Krylov iterations per Newton step.

t	Newton steps	Ø Krylov its
0.010	9	172.2
0.020	5	173.0
0.036	5	175.8
0.061	5	179.4
0.101	6	189.3
0.141	5	187.0
0.204	6	201.8
0.267	5	195.6
0.367	7	208.0
0.467	7	204.1
0.567	5	207.4
0.725	6	217.8
0.884	5	225.4
1.135	6	242.0
1.386	6	253.8
1.637	7	266.3
1.889	5	279.4
2.000	4	285.8
	Σ 104	Total ∅ 213.3

**Table 2.** Simultaneous Newton and augmented Lagrange (AL) iteration. Pseudo-time t, number of Newton-AL steps, average number of Krylov iterations per Newton-AL step.

t	Newton-AL steps	Ø Krylov its
0.010	9	99.3
0.026	4	100.5
0.051	5	101.4
0.091	6	101.3
0.154	6	102.8
0.254	7	104.3
0.412	11	105.4
0.664	14	109.4
1.062	14	119.0
1.462	16	139.7
1.862	17	167.0
2.000	15	180.8
	Σ 124	Total Ø 138.6

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known ex-post. In our example the solution using the penalty approach only satisfies 130  $|\det(\mathbf{F}) - 1| < 0.021$ .

In the results in Table 2, we see that the number of Newton-AL-iterations 132 increases during the simulation. This is due to the fact that in the beginning of the 133 simulation only a very small number of finite elements violate the element-wise condition  $|\det(\mathbf{F}) - 1| < 0.01$ .

The results in, both, Tables 1 and 2 also show an increase of the FETI-DP iterations during the simulation. We believe that this may in part be due to an increasing 137 influence of the incompressibility constraint during the simulation but also result 138 from the exponential stiffening behavior of the fibers. In [1], we have observed that 139 the anisotropies introduced to the material wall models by the terms modeling the 140 fibers can have a visible impact on the convergence of the nonlinear iteration scheme 141 as well as the convergence of the iterative linear solver. Ideas described in [16] may 142 improve the convergence of domain decomposition solvers for such anisotropic prob- 143 lems.

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#### **AUTHOR QUERIES**

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## **Techniques for Locally Adaptive Time Stepping Developed over the Last Two Decades**

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Adaptive mesh refinement techniques are well established and widely used for space 7 discretizations. In contrast, local time stepping is much less used, and the corre- 8 sponding techniques are less mature, needing delicate synchronization steps, which involve interpolation, extrapolation or projection. These operations can have adverse 10 effects on the stability, and can also destroy important geometric properties of the 11 scheme, like for example the conservation of invariants. We give here a survey on 12 the intensive research performed in this direction over the last two decades.

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## 1 Methods from the ODE Community

Local time stepping started in the ODE community with the development of split 15 Runge-Kutta methods with Rice [34]. Nowadays called multirate Runge-Kutta meth- 16 ods, these methods were first developed for naturally split systems of ordinary differential equations y' = b(y, z, t) and z' = c(y, z, t), in which the z components need 18 to be integrated on a finer time mesh than the y components. One then uses a Runge- 19 Kutta method for the fast, so called active components with a small time step, and 20 another one for the slow, so called latent components, with a large time step, and uses 21 either interpolation or extrapolation for the missing values, depending on which of 22 the components are computed first, see [27].

Multirate time integration methods were also proposed for linear multistep meth- 24 ods in [22], with two main approaches: fastest-first and slowest-first. Suppose an 25 implicit linear multistep method is used. In the fastest-first approach, one advances 26 the z components with small time steps h, and whenever one needs a component of  $\frac{1}{2}$ the slow part y, one uses a predictor step for it. Once the fine stepping scheme arrives 28 at a coarse step H, the slow solution component y is also computed. The major disadvantage of this approach is that it is very difficult to do adaptive time stepping. This 30 is easier in the slowest-first approach, where first the slow component is doing an 31 adaptive integration step, until one is accepted with step size H. Then the adaptive 32 fine integration is tried with small steps h, until one reaches with several accepted 33 small steps the coarse level H. For the slow adaptive step H however, one needs also 34

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an approximation of the fast component for coupled components, and the authors in 35 [22] say: "There are several possible ways to control the fast extrapolation error, none 36 of which is entirely satisfactory". The stability properties of such multirate schemes 37 were analyzed in [35] for Backward Euler multirate schemes; see also [23]. 38

In contrast to the multirate methods, multirate extrapolation methods aim at 39 integrating systems of ODEs without a priori knowledge of which components need 40 finer time integration steps than others. A method based on Richardson extrapolation 41 was proposed in [13]: one computes approximations for all components for a time 42 step sequence  $\{h_1, h_2, h_3, \ldots\}$ , e.g.  $h_2 = \frac{h_1}{2}$ ,  $h_3 = \frac{h_1}{3}$ ,..., and then builds the Richardson extrapolation table. As soon as a component has reached the desired accuracy 44 at step  $h_k$  (an error estimate is available automatically in the Richardson table), 45 extrapolation for this component is marked inactive, and only components needing 46 further accuracy continue the extrapolation. Inactive components must then however 47 be approximated in order for the extrapolation to continue. Using interpolation from 48 the continuous approximation obtained from the Richardson extrapolation can completely destroy the extrapolation process, which is based on the same error expansion 50 for all the components. The authors in [13] propose instead an elegant approximation 51 from the asymptotic expansion assumption itself, and also introduce a defect control 52 to avoid that inactivation fails in certain situations.

### 2 Methods from the PDE Community

Local time stepping schemes in the PDE community started with experimental work, 55 see for example [28]. Such ad hoc solutions were quite different for parabolic and 56 hyperbolic PDEs. 57

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Hyperbolic Problems: a first complete mathematical analysis of two space-time 58 adaptive schemes for the wave equation  $u_t = u_x$ , an interpolation based variant, and 59 the so called coarse mesh approximation method were given by Berger [2] (see also 60 [3], and an early analysis for a different technique based on finite volumes in [31]). 61 Using for example a three point explicit scheme, the interpolation based approach 62 starts with a coarse step at the interface, shown in red in Fig. 1 on the left, followed 63 by an interpolation for the fine grid values, shown in blue. In the coarse mesh approximation, one uses the coarse spatial mesh to compute small time steps  $\Delta t$ ,  $\Delta t$ , 65  $\Delta t$ , ... at the interface, instead of interpolating these values, as indicated in Fig. 1 on 66 the right for the second step  $\Delta t$  in red, where the blue value at  $\Delta t$  has already been 67

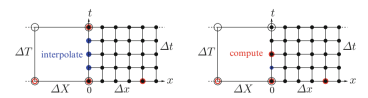
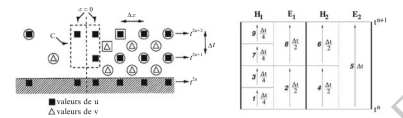


Fig. 1. Interpolation based approach on the *left*, and the coarse mesh method on the *right* 



**Fig. 2.** First energy-preserving local time stepping for the wave equation on the *left*, and symplectic scheme for Maxwell's equation on the *right* 

computed. The author proves for the hyperbolic model problem  $u_t = u_x$  that both 68 approaches are stable for the Lax-Wendroff scheme, but stability for the Leapfrog 69 scheme can only be achieved with overlap. Elegant recursive versions of such algorithms are in [33].

A key new ingredient to obtain stability for a Leapfrog type scheme for the 72 locally adaptive solution of the wave equation can be found in the seminal papers 73 by Collino et al. in [7, 8]: the introduction of a discrete energy conservation. In presentations, this approach was always introduced with an impressive movie, where a 75 wave passes a locally refined patch, and everything looks fine for quite a long time 76 after the wave has passed, until suddenly an instability forms at the boundary of the 77 patch, and the numerical solution explodes, if a simple interpolation based scheme 78 is used. The method was first described for the 1d Maxwell system  $u_t + v_x = 0$ , 79  $v_t + u_x = 0$ , which is equivalent to the 1d second order wave equation  $u_{tt} = u_{xx}$ , and 80 can best be described with the original picture from [7] shown in Fig. 2 on the left. 81 Thinking just about the second order wave equation, discretized with a centered fi- 82 nite difference scheme both in space and time, we get the five point star, well visible 83 with the black squares in Fig. 2 (the triangles would be for the unknowns v we do not 84 consider here). Now all points can be computed with this star at time levels  $t^{2n+1}$  and 85  $t^{2n+2}$ , given the values at earlier time levels, except for the values in the dashed box. 86 The key idea of the energy preserving scheme is now to permit two different values 87 at x = 0 at even time levels  $t^{2n}$ , and to introduce as additional equation the discrete 88 energy, which needs to be preserved. This leads naturally to a stable scheme, but it 89 requires the solution of a small linear system at the interface. Energy conservation 90 turned out to be a key tool for stability analysis, and is used now for other spacetime adaptive methods, see for example [11], where the authors introduce an unusual 92 energy, in order to analyze the stability of their space-time locally adaptive scheme. 93

A very elegant way of generalizing a symplectic integrator (which naturally preserves a nearby energy) for variable step size integration was presented in [26], and 95 adapted to Maxwell's system in [32]. The Störmer-Verlet scheme is symplectic for 96 these equations, and is shown in Fig. 2 on the right. Without refinement, the scheme 97 is visible in the right part under  $H_2, E_2$ : we see that first a half step denoted by 4 98 is performed for the magnetic field H, followed by a full step denoted by 5 for the 99 electric field E, and concluded by a second half step for H denoted by 6. In each of 100 these steps, the Störmer-Verlet scheme uses for H the newest values available from 101

the other field E, and vice versa. It turns out that doing the same over the locally 102 refined region shown in Fig. 2 on the right, and performing the steps in the given 103 order, starting with 1 and ending with 9, and using each time the newest information 104 available, is still symplectic! Since symplectic schemes preserve a nearby energy, 105 this scheme has all the good stability properties needed.

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In a finite volume or discontinuous Galerkin in the time domain setting (DGTD), 107 on unstructured meshes in space, the scheme in each subdomain with given time step 108 can be advanced until the new time value reaches that of its neighbor, according to 109 the stability constraint, see [12] for elastodynamics computations in the context of 110 ADER methods (Arbitrary high order, using high order DERivatives of polynomials). 111

Parabolic Problems are often integrated using implicit methods, which require 112 the solution of large systems of equations. These systems are obtained using the 113 same time step over the entire domain, and it is thus a priori not possible to use a 114 local time step. The first ideas to change this are based on domain decomposition 115 methods, where then interface values have to be predicted in some way, before the 116 subdomain problems are advanced in time by an implicit method.

A first interesting way to explicitly predict the interface values appeared in [9], 118 where a third spatial discretization size H is introduced, in addition to  $h_l$  and  $h_r$ , see 119 Fig. 3 on the left. The method then first does an explicit prediction step over the big 120  $\Delta t$ , stable because the corresponding spatial step H is big, as indicated in red. This 121 is followed by interpolation (in blue) to obtain all needed values at the interface, and 122 then on each side one can do implicit solves to advance the method. It is proved in 123 [9] that this scheme is stable for the heat equation with a centered finite difference 124 discretization in space, and forward/backward Euler in time, if  $\Delta t \leq \frac{1}{2}H^2$ , and the error satisfies the estimate max  $|\text{err}| \leq C(h_l^2 + h_r^2 + H^3 + \Delta t_l + \Delta t_r + H\Delta t)$ , which 126 shows impressively that the big prediction step  $\Delta t$ , H only affects the accuracy in 127 higher order terms!

A different approach was proposed by Blum et al. [4], as shown in Fig. 3 on the 129 right. The authors do not consider local refinement in time and space, their main 130 interest is to break up a large linear system from the implicit time integration into 131 smaller ones, but their idea can also be used for local adaptation in time and space. 132 The key idea is to use overlap, predict all values needed at the interfaces using a 133 higher order extrapolation method, and then solve implicitly on the corresponding 134 subdomains to advance the method. The authors prove for the heat equation without 135

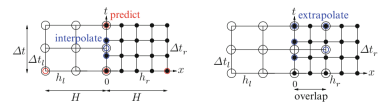
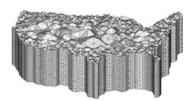
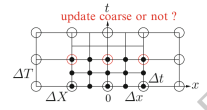


Fig. 3. Explicit prediction of the interface values on an intermediate spatial grid on the left, and by extrapolation with overlap on the right





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Fig. 4. A completely general space time mesh on the left, and the one-way and two way approaches on the right

local refinement,  $h_l = h_r = h$  and  $\Delta t_l = \Delta t_r = \Delta t$ , that the Crank-Nicolson scheme 136 is stable, provided that  $\Delta t \leq C \left(\frac{L}{\log L}\right)^2 h^2$ , where Lh is the overlap, and an error estimate 137 mate of the form  $O(\Delta t^2 + h^2)$ . So here increasing the overlap can lessen the stability 138 constraint on the time step. 139

If one wants to avoid any time step constraints, one can perform the coupling 140 fully implicitly, as proposed in [16]. Here, one simply writes the implicit scheme on 141 the fine and coarse subdomain, and the interpolation conditions into one big system 142 of linear equations, which is then solved. The authors show for a linear advection 143 reaction diffusion equation that a standard centered scheme with backward Euler in 144 time is unconditionally stable, and satisfies for  $\Delta t = O(h)$  the error estimate  $O(\Delta t +$  $h^2$ ) in 1d, but in 2d there is a loss of  $|\log h|^{\frac{1}{2}}$ , and in 3d a loss of  $\frac{1}{\sqrt{h}}$  in accuracy.

A more general approach based on domain decomposition can be found in [17]. 147 For the heat equation  $u_t = u_{xx}$ , and the decomposition of the domain  $\Omega = (-1,1)$  148 into two subdomains  $\Omega_1=(-1,0)$  and  $\Omega_2=(0,1)$ , the authors propose to discretize 149 the coupling conditions  $u_1(0) = u_2(0)$ ,  $\partial_x u_1(0) = \partial_x u_2(0)$  using a conservative finite 150 volume discretization over non-matching time grids. They also obtain, for each vari- 151 ant of the method, a very large system of equations to solve, but propose to solve it 152 using one or several steps of an iterative Dirichlet-Neumann algorithm. They show 153 that these schemes are conservative, provided one stops the iteration after a Neumann 154 step, and satisfy an error estimate  $O(\Delta t + h)$  under certain conditions. One can show 155 that one of their methods corresponds to the approach in [16].

**Space-Time Finite Element Methods** consider the time direction like one of the 157 spatial directions, and discretize the problem directly in space-time by a finite element method, which leads to a large discrete problem in space-time. These methods 159 have their roots in the work of C. Johnson and co-authors, see for instance [15] for 160 a review. Discontinuous Galerkin methods were used, and the adaptation was done 161 through a posteriori estimates. In the first versions of the method, the space-time 162 finite elements were still special, since they always had boundaries in time aligned 163 with the time direction, for example prisms. Completely general triangular meshes 164 in space time require special meshing techniques, since they need to satisfy certain 165 angle constraints, in order to avoid total global coupling in space-time, see [36] for 166 applications to Burger's equation and elastodynamics. An impressive example of 167 such a mesh from [14] is reproduced in Fig. 4 on the left. A very recent contribution 168 using discontinuous Galerkin methods can be found in these proceedings, see [30].

One-Way and Two-Way Methods are in principle very different from all the 170 methods we considered earlier, since they have both a coarse and a fine mesh in parts 171 of the domain. They have their roots in weather and climate simulations, which often use a global model over a large region, for example the entire planet, and then 173 refined models over a small region, for example a country. The question is then how 174 to compute a refined solution based on the solution of the global coarse problem. In 175 [10] and [6], the so called one-way (or "offline") and two-way (or "online") methods are proposed. In the one-way method, the coarse model is first solved once and 177 for all, and stored. Then boundary data is extracted to be imposed on the boundary 178 of the smaller refined region. The simplest approach is to use Dirichlet conditions, 179 which can however lead to large errors. A more refined approach is to use so called 180 open boundary conditions, which are related to absorbing boundary conditions, but 181 different, see [6, 29]. Open boundary conditions lead in general to substantially more 182 accurate fine models. In the two way approach, one only performs one or a few time 183 steps of the coarse model, then solves the fine model in the refined region as before, 184 but updates the coarse result whenever a more accurate fine result is available, before 185 continuing the next coarse time step, see Fig. 4 on the right. If one simulates only one 186 time step of the coarse model before solving the fine model and uses Dirichlet conditions, this approach is very much related to the first approach for hyperbolic problems 188 described earlier.

Schwarz waveform relaxation methods are the most flexible methods for solving evolution problems locally adaptively in space time, since they permit not only 191 refined time steps, but even different numerical methods, or different models in different regions. They were first described in [20] and are based on a decomposition 193 in space of the domain over which the evolution problem is posed and a subdomain 194 iteration in space-time: starting with an initial guess on each space-time interface 195 between subdomains, on each subdomain the evolution problem is solved over an 196 entire so called time window. Then information is exchanged between subdomains 197 using transmission conditions, and the subdomain problems are solved again and 198 again until a suitable matching is reached. So the price to pay for this flexibility and 199 generality is the iteration. The method from [17] we have seen earlier is in this class 200 of methods, but much faster convergence can be obtained when optimized transmis- 201 sion conditions are used, see [1, 18, 21, 24, 25], and references therein. Very general 202 non-matching space-time grids can be coupled like this using a projection algorithm 203 with optimal linear complexity from [19]. For recent realistic applications in a com- 204 plex setting, see [5].

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## Newton-Schwarz Optimised Waveform Relaxation Krylov Accelerators for Nonlinear Reactive Transport

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1 Introduction 9

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Krylov-type methods are widely used in order to accelerate the convergence of 10 Schwarz-type methods in the linear case. Authors in [2] have shown that they accelerate without overhead cost the convergence speed of Schwarz methods for different types of transmission conditions. In the nonlinear context, the well-known 13 class of Newton-Krylov-Schwarz methods (cf. [5]) for steady-state problems or time-14 dependent problems uses the following strategy: time-dependent problems are discretised uniformly in time first and then one proceeds as for steady-state problems, 16 i.e. the nonlinear problem is solved by a Newton method where the linear system 17 at each iteration is solved by a Krylov-type method preconditioned by an algebraic 18 Schwarz method. The major limitation is that NKS methods do not allow different 19 time discretisations in the subdomains since the problem is discretised in time uniformly up from the beginning.

In this work, we are interested in applying the well-established technique from 22 the linear case in the context of Schwarz Waveform Relaxation methods (SWR, cf. 23 [8]) to nonlinear time-dependent problems in order to benefit from its accelerating 24 properties. We emphasise the use of SWR methods since within this approach, it is 25 possible to use different discretisations in time and space in the subdomains, even the 26 coupling of different models is possible. In many applications, time step restrictions 27 in implicit approaches are highly localised in space due to heterogeneity and SWR 28 methods are perfectly suited to localise and isolate them in subdomains which are 29 treated with different time discretisations.

Our motivation of balancing time step restrictions in the time-dependent nonlinear case on subdomains is close to the approach in [6, 11] where the balancing of 32 nonlinearities on subdomains in the steady-state case is achieved using the permutation of domain decomposition methods and Newton's method in combination with 34 Krylov accelerators.

The paper is organised as follows: In Sect. 2 we set up the problem to solve. 36 In Sect. 3 we describe the Schwarz waveform relaxation (SWR) algorithm and the 37

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reduction to the interface variables. The new approach is described in Sect. 4. Nu- 38 merical issues and results are given in Sect. 5. 39

#### 2 Problem Description

In this paper we consider the following model in  $\Omega \times (0,T)$ ,  $\Omega \subset \mathbb{R}^d$ :

$$\partial_t(\phi w) + \mathcal{L}w + \mathcal{F}(w) = q \text{ in } \Omega \times (0, T), \tag{1}$$

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$$w(\cdot,0) = w_0 \text{ in } \Omega, \quad \mathscr{G}w = g \text{ on } \partial\Omega \times (0,T).$$
 (2)

where  $\phi(x) > 0$  is the porosity,  $w \in \mathbb{R}^s$  the vector containing the concentrations of the s chemical species.  $\mathcal{L}[\cdot] = \nabla \cdot (-a\nabla + \mathbf{b})$  is a linear operator which models diffusion 43 described by a positive scalar diffusion coefficient a > 0 and advection described by a Darcy field  $\mathbf{b} \in \mathbb{R}^d$ . The transport operator can be zero for non-mobile species.  $\mathscr{F}$  45 is a nonlinear chemical coupling operator. We impose initial conditions on  $\Omega$  given 46 by  $w_0$  and linear boundary conditions represented by  $\mathcal{G}$ , for instance Neumann or 47 Dirichlet conditions. The data g and q are source terms depending on space and time. 48

## 3 The Schwarz Waveform Relaxation Algorithm and the Classical Approach

We decompose the domain  $\Omega$  into two non-overlapping domains  $\Omega_1$  and  $\Omega_2$  and 51 call the common boundary  $\Gamma = \partial \Omega_1 \cap \partial \Omega_2$  the interface between the subdomains. 52 We introduce the following SWR algorithm with Robin transmission conditions to 53 approximate the solution of (1): given the iterate  $w_i^{k-1}$  which is equal to an initial 54 guess for the first iteration, then one step of the algorithm consists in computing 55 in parallel  $w_i^k$  for subdomains  $\Omega_i = 1, 2$ , with data coming from the neighbouring 56 subdomain  $\Omega_{\times}$ , with  $\tilde{1} = 2$  and  $\tilde{2} = 1$ .

$$\partial_t(\phi w_i^k) + \mathcal{L}w_i^k + \mathcal{F}(w_i^k) = q \quad \text{in } \Omega_i \times (0, T),$$
 (3)

$$(\partial_{n_i} + p)w_i^k = (\partial_{n_i} + p)w_{\times}^{k-1} \quad \text{on } \Gamma \times (0, T), \tag{4}$$

$$w_i^k(\cdot,0) = w_0 \text{ in } \Omega_i, \quad \mathscr{G}w_i^k = g \text{ on } \partial\Omega_i \setminus \Gamma \times (0,T),$$
 (5)

with  $n_i$  the unit outward normal of  $\Omega_i$  on  $\Gamma$  and  $p \in \mathbb{R}$ , p > 0 a constant.

It is possible to reduce algorithm (3)–(5) to the so-called interface variables. De- 59 fine the operators  $\mathcal{M}_i$ :  $(\lambda_i, f) \mapsto w_i$  solution of 60

$$\partial_t(\phi w_i) + \mathcal{L}w_i + \mathcal{F}(w_i) = q \quad \text{in } \Omega_i \times (0, T),$$
 (6)

$$(\partial_{n_i} + p)w_i = \lambda_i \quad \text{on } \Gamma \times (0, T),$$
 (7)

$$(\partial_{n_i} + p)w_i = \lambda_i \quad \text{on } \Gamma \times (0, T),$$

$$w_i^k(\cdot, 0) = w_0 \text{ in } \Omega_i, \quad \mathscr{G}w_i^k = g \text{ on } \partial\Omega_i \setminus \Gamma \times (0, T).$$
(8)

Here  $f = (q, w_0, g)$  represents all source terms except the ones on the interface  $\Gamma$  that 61 are represented separately by  $\lambda_i$ . With these definitions, the transmission conditions 62 (4) can be written as  $\lambda_i^{k+1} = -\lambda_{\times}^k + 2p\mathcal{M}_{\times}(\lambda_{\times}^k, f)$ , and as a system 63

Domain Decomposition for Nonlinear Reactive Transport

$$\begin{pmatrix} \lambda_1^k \\ \lambda_2^k \end{pmatrix} = \begin{pmatrix} -\lambda_2^{k-1} + 2p\mathcal{M}_2(\lambda_2^{k-1}, f) \\ -\lambda_1^{k-1} + 2p\mathcal{M}_1(\lambda_1^{k-1}, f) \end{pmatrix}.$$
 (9)

The SWR algorithm (3) is therefore a fixed point algorithm for the nonlinear 64 interface problem 65

 $\begin{pmatrix} \lambda_1 \\ \lambda_2 \end{pmatrix} = \begin{pmatrix} -\lambda_2 + 2p\mathcal{M}_2(\lambda_2, f) \\ -\lambda_1 + 2p\mathcal{M}_1(\lambda_1, f) \end{pmatrix}. \tag{10}$ 

AQ1 Each iterate requires solving the nonlinear problem (6)–(8). This can be achieved by a Newton method, or a semi-implicit discretisation in time. The latter method 67 has been implemented in [4] for the advection diffusion reaction equation, where the 68 convergence of the fixed point algorithm has been proved. The extension of the proof 69 to the system (1) should be easy.

### 4 Newton-Schwarz Optimised Waveform Relaxation

The new approach consists first in solving the system (10) by a Newton algorithm. If 72 the interface problem is well-posed, and if the initial data for Newton is sufficiently 73 closed to the solution, the algorithm converges to that solution. According to the 74 interface problem (10), we seek the zeros of the nonlinear function 75

$$\Theta(\lambda) := -(\lambda_1 + \lambda_2) \begin{pmatrix} 1 \\ 1 \end{pmatrix} + 2p \Upsilon(\lambda), \quad \Upsilon(\lambda) := \begin{pmatrix} \mathcal{M}_2(\lambda_2, f) \\ \mathcal{M}_1(\lambda_1, f) \end{pmatrix}.$$

One step  $k-1 \to k$  of Newton's algorithm consists in solving the linear system 76  $\Theta'(\lambda^{k-1}) \cdot (\lambda^k - \lambda^{k-1}) = -\Theta(\lambda^{k-1})$ . To evaluate the derivative of  $\Theta$ , we must 77 calculate the derivative of the functions  $\lambda_i \mapsto \mathscr{M}_i(\lambda_i, f)$ . If  $w_i = \mathscr{M}_i(\lambda_i, f)$  and 78  $W_i = \mathscr{M}_i(\lambda_i + \tilde{\lambda}_i, f)$ , we see by subtracting equations (6) for  $w_i$  and  $W_i$ , that  $W_i - w_i$  79 is solution of

$$\partial_t(\phi(W_i-w_i))+\mathscr{L}(W_i-w_i)+\mathscr{F}(W_i)-\mathscr{F}(w_i)=0.$$
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Introducing the derivative of  $\mathscr{F}$ ,  $\mathscr{F}(W_i) - \mathscr{F}(w_i) = \mathscr{F}'(w_i)(W_i - w_i) + \mathscr{O}((W_i - w_i)^2)$ , and therefore  $W_i - w_i = \tilde{w}_i + o(\tilde{w}_i^2)$ , where  $\tilde{w}_i$  is solution of the linear equation 83

$$\partial_t(\phi \tilde{w_i}) + \mathcal{L}\tilde{w}_i + \mathcal{F}'(w_i)\tilde{w}_i = 0. \tag{11}$$

$$(\partial_{n_i} + p)\tilde{w}_i = \tilde{\lambda}_i \tag{12}$$

$$\tilde{w}_i(x,0) = 0 \text{ in } \Omega_i, \quad \mathscr{G}\tilde{w}_i = 0 \text{ on } \partial\Omega_i \setminus \Gamma \times (0,T).$$
 (13)

Therefore  $\partial_{\lambda_i} \mathcal{M}_i(\lambda_i, f) \cdot \tilde{\lambda}_i = \tilde{w}_i := \mathcal{M}^{lin}(\mathcal{F}'(w_i); \tilde{\lambda}_i)$ , and

$$\Theta'(\lambda) \cdot \tilde{\lambda} = -(\tilde{\lambda}_1 + \tilde{\lambda}_2) \begin{pmatrix} 1 \\ 1 \end{pmatrix} + 2p \begin{pmatrix} \mathscr{M}^{lin}(\mathscr{F}'(w_2); \tilde{\lambda}_2) \\ \mathscr{M}^{lin}(\mathscr{F}'(w_1); \tilde{\lambda}_1) \end{pmatrix}.$$

After these computations, the algorithm takes the form

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$$w_{i}^{k-1} = \mathcal{M}_{i}(\lambda_{i}^{k-1}, f),$$

$$-\sum_{i=1}^{2} (\lambda_{i}^{k} - \lambda_{i}^{k-1}) \begin{pmatrix} 1\\1 \end{pmatrix} + 2p \begin{pmatrix} \mathcal{M}_{2}^{lin}(\mathcal{F}'(w_{2}^{k-1}); \lambda_{2}^{k} - \lambda_{2}^{k-1}) \\ \mathcal{M}_{1}^{lin}(\mathcal{F}'(w_{1}^{k-1}); \lambda_{1}^{k} - \lambda_{1}^{k-1}) \end{pmatrix} =$$

$$-\sum_{i=1}^{2} \lambda_{i}^{k} \begin{pmatrix} 1\\1 \end{pmatrix} + 2p \begin{pmatrix} \mathcal{M}_{2}(\lambda_{2}^{k-1}, f) \\ \mathcal{M}_{1}(\lambda_{1}^{k-1}, f) \end{pmatrix}$$

$$(14)$$

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The approach requires in every iteration to solve two nonlinear problems in the 86 subdomains. Therefore, a nested iterative procedure is necessary (Newton, or semiimplicit time stepping). Once this is done,  $\lambda^{n+1} - \lambda^n$  is a solution of a linear problem 88 solved in parallel in the subdomains.

## 5 Implementation Using Newton-Krylov Methods and Numerical Results

We have implemented both the classical and the new approach for a special case 92 of problem (1). We assume that s=2 and w=(u,v) where u denotes a mobile 93 species and v denotes a fixed species. The nonlinear function  $\mathscr{F}$  is given by  $\mathscr{F}(w) = 94$ (R(u,v), -R(u,v)) where R(u,v) is the overall reaction rate of the reversible reaction  $u \stackrel{\longleftarrow}{\longrightarrow} v$ .

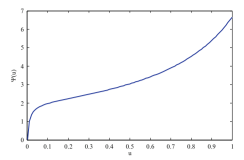
For the computation of  $\mathcal{M}_i(\lambda_i^{k-1}, f)$ , we use an implicit Euler scheme in time 97 and a hybrid finite volume scheme (based on [7]) in space. The nonlinear systems 98 are then treated with a global implicit approach by means of Newton's method with 99 exact LU-decomposition. The linear interface problems (14) for  $\lambda_i^k$  are solved using GMRES as Krylov-type method with a precision strategy in the spirit of inexact 101 Newton methods: we adapt the precision of the linear solver with respect to the 102 residuals of the Newton iterates and save therefore costly subdomain evaluations.

Concerning the stopping criterion for the Newton-Schwarz optimised algorithm, 104 it is classically controlled by both the residual and the correction  $(\Delta\lambda)$  norm. The 105 Schwarz optimised algorithm is only controlled by the correction norm.

For all tests, we set the simulation domain to  $\Omega = [0,1] imes [0,1] \subset \mathbb{R}^2$  with 107 the subdomains  $\Omega_1 = [0, 0.5] \times [0, 1]$  and  $\Omega_2 = [0.5, 1] \times [0, 1]$ . The time window considered is  $t \in [0, 1]$ . Physical parameters are  $\phi = 1$ , a = 1.5,  $(b_x, b_y) =$  $(5 \cdot 10^{-2}, 1 \cdot 10^{-3})$ . The nonlinear coupling term is defined by  $R(u, v) = k(v - \Psi(u))$  110 where the function  $\Psi$  is a BET isotherm law defined by

$$\Psi(u) = \frac{Q_s K_L u}{(1 + K_L u - K_S u)(1 - K_S u)}.$$

BET theory is a rule for the physical adsorption of gas molecules on a solid surface 112 and serves as the basis for an important analysis technique for the measurement of 113 the specific surface area of a material (cf. [3]). This law is insofar mathematically interesting as it is neither convex nor concave (cf. Fig. 1) and is therefore a challeng- 115 ing problem for standard nonlinear solvers like Newton's method. We set k = 100, 116



**Fig. 1.** BET Isotherm law function  $\Psi$  with  $Q_S = 2$ ,  $K_S = 0.7$ ,  $K_L = 100$ 

 $Q_S=2$ ,  $K_S=0.7$  and  $K_L=100$ . Initial values are set to  $(u_0,v_0)=(\frac{1}{2},\frac{1}{3})$ . By defining the function  $g(x, y, t) = (\sin(\pi x)\cos(\pi y)\cos(2\pi t) + \cos(\pi x)\sin(\pi y)\cos(2\pi t) +$  118  $\cos(\pi x)\cos(\pi y)\sin(2\pi t)+1)/2$  we impose Dirichlet boundary conditions with values set to u(x, y, t) = g(x, y, t) for  $(x, y) \in \partial \Omega$ .

As a first experiment, we are interested in the sensitivity of the new approach with 121 respect to the parameter p of the Robin transmission condition. Indeed the theory of 122 optimised Schwarz waveform relaxation for linear problem relies on the fact that the 123 convergence properties of the algorithm heavily depend on this parameter. A best 124 parameter for the advection diffusion reaction equation can be found analytically by 125 solving a best approximation problem, see [1, 8]. No such analysis is available for 126 the nonlinear problem, it is therefore interesting to study the issue numerically.

We discretise the numerical domain with  $\Delta x = \Delta y = 1/40$  and  $\Delta t = 1/10$  and impose a random initial guess on the interface for the first iteration. As both subdomains 129 are the same size, the number of overall matrix inversions is a meaningful criterion 130 for measuring the numerical performance. We run the two approaches for different 131 parameters p of the Robin transmission condition and plot in Fig. 2 (left) the num- 132 ber of matrix inversions as a function of the parameter p in the Robin transmission 133 condition. One observes first that the performance of the classical approach depends 134 highly on the parameter p of the Robin transmission condition, as in the linear case. 135 The best parameter is  $p^* \approx 40$ . We observe that the new approach also shows the best performance at  $p^*$  but is much less sensitive to the choice of the parameter. The loss 137 of sensitivity with respect to the parameter is still an open question.

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It turns out that the new method has a cost overhead, that becomes non negli- 139 gible if space discretisations are chosen too coarse. For this reason, we study the 140 asymptotic behaviour of the two approaches using always the optimal parameter of 141 the classical approach. We refine the problem in space using always  $\Delta x = \Delta y$ . Note 142 that we keep the time step constant at  $\Delta t = 0.1$ . Refining the discretisation also in 143 time would lead to a problem that is quasi stationary at every time step since we use a 144 global implicit approach. We measure again the overall number of matrix inversions 145 in the two approaches and plot them in Fig. 2 (right) versus the discretisation size. 146 One observes that the overhead cost of the new approach compared to the classical 147 approach becomes negligible starting at a discretisation with about 150 grid points 148

per dimension for the new method. For problems finer than the respective thresholds, 149 the new approach is always faster than the classical approach with the best parameter 150 for the transmission condition. Moreover, the finer the discretisation, the larger the 151 problem, the more important the accelerating property of the new approach. Note that 152 the new approaches has a slope of  $O(N^{1/7})$  in the asymptotic behaviour which is considerably less than the slope of the classical approach which behaves like  $O(N^{1/2.75})$ . 154 The slopes have been determined graphically, no theoretical justification is available. 155 However, this plot shows that the method is much less dependent of the size of the 156 problems than the classical one.

In order to exemplify the accelerating property of the new approach, we perform a 158 simulation with  $N_x = N_y = 200$  points in each dimension keeping the number of time 159 steps constant and compare the convergence behaviour of the stopping criteria of the 160 two methods. In Fig. 3 we plot the convergence criterion versus the number of matrix 161 inversions. Note that, for a better comparison, we set the residual norm of the nonlinear interface problem evaluated at the initial guess for both methods at zero matrix 163 inversions. The classical approach exhibit a linear convergence followed by a superlinear convergence, similar to the behaviour of the linear algorithm. We observe the 165 quadratic convergence of the new approach, the characteristic feature of the Newton 166 algorithm.

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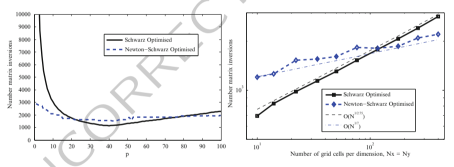
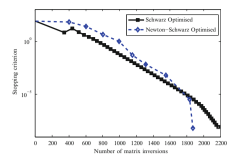
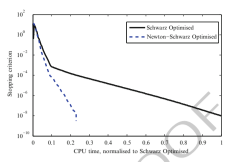


Fig. 2. Number of matrix inversions for the classical approach and new approach, synthetic test case. Left: Varying parameter p of the Robin transmission conditions with fixed discretisation in space and time. Right: Varying the number of discrete points per dimension  $(N_x = N_y)$  with fixed discretisation in time and optimal parameter for the Robin transmission condition

Finally, we want to apply the new approach to a benchmark test case in the context of CO<sub>2</sub> geological storage. The 3D test case is based on the benchmark for 169 the SHPCO2 project (Simulation haute performance pour le stockage géologique 170 du CO<sub>2</sub>) which is described in [10]. The global domain is set to  $\Omega = [0, 4,750] \times 171$  $[0, 3,000] \times [-1,100, -1,000]$  with (38, 24, 8) grid cells in (x, y, z)-direction. The domain is decomposed into the two nonoverlapping subdomains  $\Omega_1 = [1,000,2,500] \times 173$  $[0, 3,000] \times [-1,050, -1,000]$  and  $\Omega_2 = \Omega \setminus \Omega_1$ . We call  $\Omega_1$  the reactive subdomain 174 since in this subdomain an injection of the mobile species u is modelled by a source 175 term. The initial state is zero for the mobile and immobile species. We consider 176





**Fig. 3.** Convergence history with 200 points per space dimension for the classical approach and new approach, synthetic test case

Fig. 4. Convergence history for the classical approach and new approach, SHPCO2 benchmark case

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again the BET isotherm law as nonlinear coupling term. The injected mobile species 177 is partially adsorbed by the reaction and partially transported by mainly advection. 178 Simulation time is [0, 100]. The SWR approach allows us to use different discretisa- 179 tions in the subdomains. We choose to use ten time steps in the reactive subdomain 180  $\Omega_1$  and only five time steps in the subdomain  $\Omega_2$ . This choice is insofar justified 181 since the rapid injection in the reactive subdomain restricts the time step size by imposing a maximum number of Newton iterates of ten. As in the subdomain  $\Omega_2$ , the 183 mobile species appears only by transport processes on a slower time scale than the 184 injection, one can choose a larger time step in order to respect the maximum number 185 of Newton iterations. Concerning the parameter of the Robin transmission condition, 186 we use a low frequency approximation of the optimal parameter. The initial guess on 187 the interface is zero for both subdomain interfaces. In Fig. 4 we plot the convergence 188 histogram, i.e. the stopping criterion in a logarithmic scale versus the CPU time (normalised to the CPU time of the classical approach). Note that both subdomains have 190 a different size of unknowns and therefore the number of matrix inversions, as used 191 in the previous examples, is no longer a valid tool to measure the effort. One ob- 192 serves that the new approach needs only about 20 % of the CPU time of the classical 193 approach.

6 Conclusion 195

Based on a nonlinear coupled reactive transport system we have developed a new 196 approach for solving the interface problem in the nonlinear case using Krylov- 197 accelerators. In contrast to NKS methods the use of SWR methods allows us to use 198 different time discretisations in the subdomains and so to localise time stepping constraints. We have implemented and tested the method, comparative results with the 200 classical approach have been provided.

The numerical tests showed that, besides an overhead cost for coarse space dis- 202 cretisations, the method has an accelerating property and shows much less sensitiv- 203 ity with respect to the choice of the parameter of the Robin condition. The quadratic 204 convergence behaviour of the new approach outperforms the superlinear convergence 205 behaviour of the classical approach. Nevertheless, the new approach does have significant overhead costs that are not negligible in the case of coarse problems. Note 207 that a third approach is possible, namely to start with a Newton algorithm for the 208 nonlinear problem, and to solve the so obtained linear problem by a Schwarz-Krylov 209 algorithm (cf. [9]). 210

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# Alternating and Linearized Alternating Schwarz Methods for Equidistributing Grids

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1 Introduction 7

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The solution of partial differential equations (PDEs) with disparate space and time 8 scales often benefit from the use of nonuniform meshes and adaptivity to successfully 9 track local solution features.

In this paper we consider the problem of grid generation using the so-called 11 equidistribution principle (EP) [3] and domain decomposition (DD) strategies. In 12 the time dependent case, the EP is used to evolve an initial (often uniform) grid by 13 relocating a fixed number of mesh nodes. This leads to a class of adaptive meth-14 ods known as r-refinement or moving mesh methods. A thorough recent review of 15 moving mesh methods for PDEs can be found in the book [11].

In general, the appropriate grid for a particular problem depends on features of the (typically unknown) solution of the PDE. Here we will focus on the grid generation problem for the time independent, given function u(x) of a single spatial variable  $x \in [0,1]$ . Given some positive measure M(x) of the error or difficulty in the solution u(x), the EP requires that the mesh points are chosen so that the error contribution on each interval  $[x_{i-1},x_i]$  is the same. The function M is known as the monitor or mesh density function. Mathematically, we may write this as

$$\int_{x_{i-1}}^{x_i} M(\tilde{x}) d\tilde{x} \equiv \frac{1}{N} \int_0^1 M(\tilde{x}) d\tilde{x} \quad \text{or} \quad \int_0^{x(\xi_i)} M(\tilde{x}) d\tilde{x} = \frac{i}{N} \theta \equiv \xi_i \theta,$$
 (EP)

where  $x(\xi_i) = x_i$  and  $\theta \equiv \int_0^1 M(\tilde{x}) d\tilde{x}$  is the total error in the solution. The EP defines 24 a one–to–one co-ordinate transformation between the physical co–ordinate x and 25 underlying computational co–ordinate  $\xi$ . This will naturally concentrate mesh points 26 where the error in the solution is large.

Differentiating the continuous formulation of EP gives the required mesh transformation,  $x(\xi)$ , as the solution of the nonlinear boundary value problem

$$\frac{d}{d\xi}\left\{M(x(\xi))\frac{d}{d\xi}x(\xi)\right\} = 0, \quad x(0) = 0 \quad \text{and} \quad x(1) = 1.$$
 (1)

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If M is chosen properly, we expect the solution u(x) to be easy to represent on 30 a uniform grid in  $\xi$ . In general, the physical solution u is not known and instead 31 satisfies a PDE. In that case, the mesh transformation, satisfying (1), and the physical 32 PDE, are coupled and often solved in an iterative fashion.

We will assume (1) has a unique solution, see [8] for details. In [8], the authors 34 consider the solution of (1) and time dependent extensions using classical parallel, 35 optimized and optimal Schwarz methods. In this paper we continue the work of [8] 36 by providing details of the nonlinear and linearized alternating Schwarz approaches. 37 The reader is also referred to the experimental papers [7, 9, 10], which proposed 38 various strategies to couple DD and moving meshes. See [1, 2, 4-6, 12-15] for a 39 discussion of DD methods applied to other nonlinear PDEs.

In Sect. 2 we propose a new nonlinear alternating Schwarz method to solve (1) 41 and prove convergence in  $L^{\infty}$ . In Sect. 3 we avoid the nonlinear subdomain problems 42 and propose and analyze a linearized alternating Schwarz algorithm. Brief numerical 43 results are presented in the final section.

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# 2 A Nonlinear Alternating Schwarz Method

In [8] we consider the solution of (1) by a parallel, classical nonlinear Schwarz iteration. On each subdomain a nonlinear BVP is solved and Dirichlet transmission conditions are used at the subdomain interfaces. Convergence of the iteration can be accelerated if we are willing to compute sequentially. Consider the nonlinear alternating Schwarz iteration

$$(M(x_1^n)x_{1,\xi}^n)_{\xi} = 0, \ \xi \in \Omega_1, \qquad (M(x_2^n)x_{2,\xi}^n)_{\xi} = 0, \ \xi \in \Omega_2,$$

$$x_1^n(0) = 0, \qquad x_2^n(\alpha) = x_1^n(\alpha), \qquad (2)$$

$$x_1^n(\beta) = x_2^{n-1}(\beta), \qquad x_2^n(1) = 1,$$

where  $\Omega_1 = (0, \beta)$  and  $\Omega_2 = (\alpha, 1)$  with  $\alpha < \beta$ .

Direct integration and enforcing the boundary conditions gives the following implicit representation of the subdomain solutions. 48

**Lemma 1.** The subdomain solutions on  $\Omega_1$  and  $\Omega_2$  of (2) are given implicitly as

$$\int_0^{x_1^n(\xi)} M(\tilde{x}) d\tilde{x} = \frac{\xi}{\beta} \int_0^{x_2^{n-1}(\beta)} M(\tilde{x}) d\tilde{x},\tag{3}$$

and

$$\int_0^{x_2^n(\xi)} M(\tilde{x}) d\tilde{x} = \frac{1-\xi}{1-\alpha} \int_0^{x_1^n(\alpha)} M(\tilde{x}) d\tilde{x} + \frac{\xi-\alpha}{1-\alpha} \int_0^1 M(\tilde{x}) d\tilde{x}. \tag{4}$$

Let  $\|\cdot\|_{\infty}$  denote the usual  $L^{\infty}$  norm. We now relate  $x_{1,2}^n$  to  $x_{1,2}^{n-1}$  and obtain the following result. 52

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**Theorem 1.** Assume M is differentiable and there exist positive constants a and A 53 satisfying  $0 < a \le M(x) \le A < \infty$ . Then the alternating Schwarz iteration (2) con-54 verges for any initial guess  $x_2^0(\beta)$  and we have the error estimates

$$||x - x_1^{n+1}||_{\infty} \le \rho^n \frac{A}{a} |x(\beta) - x_2^0(\beta)|, \quad ||x - x_2^{n+1}||_{\infty} \le \rho^n \frac{A}{a} |x(\alpha) - x_1^0(\alpha)|, \quad (5)$$

with contraction factor  $\rho := \frac{\alpha}{\beta} \frac{1-\beta}{1-\alpha} < 1$ .

*Proof.* Evaluating (3) at  $\xi = \alpha$  and using the expression for  $x_2^{n-1}(\beta)$  from (4) we 57 have

$$\int_0^{x_1^n(\alpha)} M \, d\tilde{x} = \frac{\alpha}{\beta} \left\{ \frac{\beta - 1}{\alpha - 1} \int_0^{x_1^{n-1}(\alpha)} M \, d\tilde{x} + \frac{\beta - \alpha}{1 - \alpha} \int_0^1 M \, d\tilde{x} \right\}.$$
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Defining the two quantities

o quantities 60 
$$K_1^n = \int_0^{x_1^n(\alpha)} M(\tilde{x}) \, d\tilde{x} \qquad \text{and} \qquad C = \int_0^1 M(\tilde{x}) \, d\tilde{x}, \qquad 61$$
 near iteration 62

we obtain the linear iteration

$$K_1^n = \frac{\alpha}{\beta} \frac{\beta - 1}{\alpha - 1} K_1^{n-1} + \frac{\alpha}{\beta} \frac{\beta - \alpha}{1 - \alpha} C. \tag{6}$$

This iteration converges with rate  $\rho := \frac{\alpha}{\beta} \frac{1-\beta}{1-\alpha} < 1$ , and has the limit

$$K_1^* = \frac{\alpha}{\beta} \frac{1 - \beta}{1 - \alpha} K_1^* + \frac{\alpha}{\beta} \frac{\beta - \alpha}{1 - \alpha} C \implies K_1^* = \alpha C. \tag{7}$$

Since the monodomain solution also satisfies

$$\int_0^{x(\alpha)} M(\tilde{x}) d\tilde{x} = \alpha C,$$

and  $M(x) \ge a > 0$ , we have convergence at the interface to the correct limit.

Subtracting (6) from (7) we have

$$\int_{x_1^n(\alpha)}^{x(\alpha)} M(\tilde{x}) d\tilde{x} = \rho^n \int_{x_1^0(\alpha)}^{x(\alpha)} M(\tilde{x}) d\tilde{x}.$$
 (8)

Subtracting (4) from the equivalent expression for the exact solution and using (8) 67 we obtain

$$\int_{x_2^{n+1}(\xi)}^{x(\xi)} M(\tilde{x}) d\tilde{x} = \frac{1-\xi}{1-\alpha} \int_{x_1^n(\alpha)}^{x(\alpha)} M(\tilde{x}) d\tilde{x} = \frac{1-\xi}{1-\alpha} \rho^n \int_{x_1^0(\alpha)}^{x(\alpha)} M(\tilde{x}) d\tilde{x}.$$

Taking the modulus and using the boundedness of M we obtain, for all  $\xi \in [\alpha, 1]$ ,

$$|x(\xi) - x_2^{n+1}(\xi)| \le \frac{1-\xi}{1-\alpha} \rho^n \frac{A}{a} |x(\alpha) - x_1^0(\alpha)|.$$

Taking the supremum gives the second estimate in (5). The estimate on subdomain one is obtained similarly. 

## 3 A Linearized Alternating Schwarz Method

We may avoid nonlinear solves on each subdomain in (2) by considering a linearized alternating Schwarz iteration,

$$(M(x_1^{n-1})x_{1,\xi}^n)_{\xi} = 0, \ \xi \in \Omega_1 \qquad (M(x_2^{n-1})x_{2,\xi}^n)_{\xi} = 0, \ \xi \in \Omega_2$$

$$x_1^n(0) = 0, \qquad x_2^n(\alpha) = x_1^n(\alpha),$$

$$x_1^n(\beta) = x_2^{n-1}(\beta), \qquad x_2^n(1) = 1.$$
(9)

At iteration n we evaluate the nonlinear diffusion coefficient M using the solution 72 obtained from the previous iterate and obtain the updated solution by a single linear 73 BVP solve on each subdomain. A simple calculation yields the following represen- 74 tation of the subdomain solutions.

**Lemma 2.** The subdomain solutions of (9) are given by

$$x_1^n(\xi) = x_2^{n-1}(\beta) \frac{\int_0^{\xi} \frac{d\tilde{\xi}}{M(x_1^{n-1}(\tilde{\xi}))}}{\int_0^{\beta} \frac{d\tilde{\xi}}{M(x_1^{n-1}(\tilde{\xi}))}},$$
(10)

and

$$x_{1}^{n}(\xi) = x_{2}^{n-1}(\beta) \frac{\int_{0}^{\xi} \frac{d\tilde{\xi}}{M(x_{1}^{n-1}(\tilde{\xi}))}}{\int_{0}^{\beta} \frac{d\tilde{\xi}}{M(x_{1}^{n-1}(\tilde{\xi}))}},$$

$$(10)$$

$$x_{2}^{n}(\xi) = x_{1}^{n}(\alpha) + (1 - x_{1}^{n}(\alpha)) \frac{\int_{\alpha}^{\xi} \frac{d\tilde{\xi}}{M(x_{2}^{n-1}(\tilde{\xi}))}}{\int_{\alpha}^{1} \frac{d\tilde{\xi}}{M(x_{2}^{n-1}(\tilde{\xi}))}}.$$

Convergence of the linearized alternating Schwarz iteration (9) follows by proving 78 convergence at the interior interfaces and showing we have converged to the correct 79 limit. 80

**Theorem 2.** Under the assumptions of Theorem 1, the linearized alternating Schwarz 81 iteration (9) converges for any smooth initial guesses  $x_1^0(\xi)$  and  $x_2^0(\xi)$ . 82

*Proof.* Evaluating the subdomain solutions (10) and (11) at the interfaces, we obtain 83 for the interface values the iterations 84

$$x_1^n(\alpha) = \mathscr{C}_{\alpha}^n x_1^{n-1}(\alpha) + \mathscr{D}_{\alpha}^n \quad \text{and} \quad x_2^n(\beta) = \mathscr{C}_{\beta}^n x_2^{n-1}(\beta) + \mathscr{D}_{\beta}^n.$$

where 85

$$\mathscr{C}^n_{\alpha} = \frac{\int_{\beta}^{1} \frac{d\xi}{M(x_{2}^{n-2}(\xi))}}{\int_{\alpha}^{1} \frac{d\xi}{M(x_{2}^{n-2}(\xi))}} \frac{\int_{0}^{\alpha} \frac{d\xi}{M(x_{1}^{n-1}(\xi))}}{\int_{0}^{\beta} \frac{d\xi}{M(x_{1}^{n-1}(\xi))}}, \quad \mathscr{D}^n_{\alpha} = \frac{\int_{\alpha}^{\beta} \frac{d\xi}{M(x_{2}^{n-2}(\xi))}}{\int_{\alpha}^{1} \frac{d\xi}{M(x_{2}^{n-2}(\xi))}} \frac{\int_{0}^{\alpha} \frac{d\xi}{M(x_{1}^{n-1}(\xi))}}{\int_{0}^{\beta} \frac{d\xi}{M(x_{1}^{n-1}(\xi))}},$$

and

$$\mathscr{C}^{n}_{\beta} = \frac{\int_{\beta}^{1} \frac{d\tilde{\xi}}{M(\chi_{2}^{n-1}(\tilde{\xi}))}}{\int_{\alpha}^{1} \frac{d\tilde{\xi}}{M(\chi_{1}^{n-1}(\tilde{\xi}))}} \int_{0}^{\alpha} \frac{d\tilde{\xi}}{M(\chi_{1}^{n-1}(\tilde{\xi}))}, \qquad \mathscr{D}^{n}_{\beta} = \frac{\int_{\alpha}^{\beta} \frac{d\tilde{\xi}}{M(\chi_{2}^{n-1}(\tilde{\xi}))}}{\int_{\alpha}^{1} \frac{d\tilde{\xi}}{M(\chi_{1}^{n-1}(\tilde{\xi}))}}.$$

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It is possible to show the quantities  $\mathscr{C}^n_{\alpha}, \mathscr{D}^n_{\alpha}, \mathscr{C}^n_{\beta}$  and  $\mathscr{D}^n_{\beta}$  satisfy

$$0<\mathscr{C}^n_\alpha,\mathscr{C}^n_\beta\leq\rho<1,\quad 0<\mathscr{D}^n_\alpha\leq D_\alpha<1,\quad \text{and}\quad 0<\mathscr{D}^n_\beta\leq D_\beta<1,$$

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$$\rho := \frac{1}{1 + \frac{a}{A} \frac{\beta - \alpha}{1 - \beta}} \frac{1}{1 + \frac{a}{A} \frac{\beta - \alpha}{\alpha}}, \quad D_{\alpha} := \frac{1}{1 + \frac{a}{A} \frac{\beta - \alpha}{\alpha}} \frac{1}{1 + \frac{a}{A} \frac{1 - \beta}{\beta - \alpha}}, \quad \text{and} \quad D_{\beta} := \frac{1}{1 + \frac{a}{A} \frac{1 - \beta}{\beta - \alpha}}. \quad 90$$

To establish these bounds let F(x) := 1/M(x). The assumptions on M imply  $\frac{1}{4} \le$ 91  $F(x) \leq \frac{1}{a}$ . As an example, the upper and lower bounds on F then imply

$$\frac{\int_0^\alpha F(x(\xi)) d\xi}{\int_0^\beta F(x(\xi)) d\xi} \le \frac{1}{1 + \frac{a}{A} \frac{\beta - \alpha}{\alpha}} \quad \text{and} \quad \frac{\int_0^1 F(x(\xi)) d\xi}{\int_\alpha^1 F(x(\xi)) d\xi} \le \frac{1}{1 + \frac{a}{A} \frac{\beta - \alpha}{1 - \beta}}.$$

Consider now the iteration for  $x_1^n(\alpha)$  only. Using the recursion, we have

$$x_1^n(\alpha) = \prod_{k=1}^n \mathscr{C}_{\alpha}^k x_1^0(\alpha) + \sum_{k=1}^n \mathscr{D}_{\alpha}^k \left( \prod_{l=k+1}^n \mathscr{C}_{\alpha}^l \right),$$
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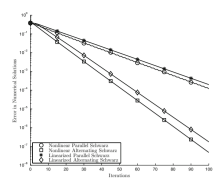
where the product in the k-th term of the sum is assumed to be one if the lower index 96 of the product exceeds the upper index. Since  $\rho < 1$ , the product multiplying  $x_1^0(\alpha)$ must go to zero as  $n \to \infty$ . The infinite series converges by direct comparison with 98  $\sum_{k=1}^{\infty} D_{\alpha} \rho^{k-1}$ . A corresponding argument applies to show convergence of  $x_2^n(\beta)$ .

Denote the limits of  $\{x_1^n(\alpha)\}\$  and  $\{x_2^n(\beta)\}\$  as  $\tilde{x}_{\alpha}$  and  $\tilde{x}_{\beta}$  respectively. Since the interface values converge, the subdomain solutions defined by (9) converge to functions  $\tilde{x}_1$  and  $\tilde{x}_2$  both satisfying the nonlinear PDE. Since  $\tilde{x}_1(\alpha) = \tilde{x}_2(\alpha)$  and  $\tilde{x}_1(\beta) = \tilde{x}_2(\beta)$ , both  $\tilde{x}_1$  and  $\tilde{x}_2$  satisfy the same PDE in the overlap with the same two boundary conditions, and by assumption of uniqueness,  $\tilde{x}_1$  and  $\tilde{x}_2$  must coincide in the overlap. One can therefore simply glue these two solutions together in order to obtain a function which satisfies the PDE everywhere, and also the two original boundary conditions at 0 and 1. Again by uniqueness, this must now be the desired solution.

### 4 Numerical Results

In this section we numerically demonstrate the results above using a simple finite 101 difference discretization of the BVP (1) and iterations (2) and (9). We also include 102 results using nonlinear and linearized parallel Schwarz algorithm from [8] for comparison. Details of the numerical approach and convergence of the discrete DD algorithm will be considered elsewhere.

We solve EP for  $u(x) = (1 - e^{\lambda x})/(1 - e^{\lambda})$  on the interval  $x \in [0, 1]$ . For large values of  $\lambda$  this function exhibits a boundary layer at x = 1. We use the arc-length 107 monitor function  $M(x,u(x)) = \sqrt{1+u_x^2}$  and choose  $\lambda = 20$ . The errors reported in Figs. 1 and 2 are the differences between the single domain numerical solution and 109 the domain decomposition solution over the first subdomain. 110



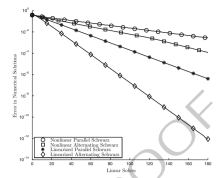


Fig. 1. Error versus # of DD iterations

Fig. 2. Error versus # of linear solves

t1

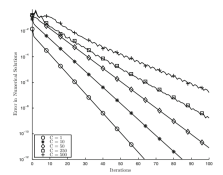
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In Fig. 1 we solve (1) on two subdomains with a 5 % overlap using linearized and 111 nonlinear, parallel and alternating Schwarz iterations. We see that the convergence 112 of the alternating iteration is faster than the parallel algorithms for both the nonlinear and linearized versions of the algorithms. In terms of number of iterations the 114 nonlinear algorithms outperform the linearized variants. It is important, however, to 115 keep in mind that each nonlinear DD iteration is more expensive than its linearized 116 counterpart. In Fig. 2 we repeat the convergence history as a function of a *work unit* 117 which we take to be the cost of a linear solve. Each iteration of a linearized Schwarz 118 algorithm requires one linear solve while each iteration of a nonlinear Schwarz algorithm requires many linear solves – one for each Newton step. Each linear solve 120 required by both algorithms has roughly the same cost due to the structure of the Jacobian matrix. As a function of the work effort the efficacy of the linearized Schwarz 122 algorithms is obvious for this example.

In Table 1 we demonstrate the quality of the computed grids by calculating the  $\|\cdot\|_{\infty}$  error between u(x) and the piecewise linear interpolant for u(x) on grids obtained by the nonlinear and linearized alternating Schwarz algorithms, as a function of the number of iterations. The last column shows the interpolation error obtained with the single domain grid: the solution of (1) computed on a uniform  $\xi$  grid consisting of 101 points. All interpolation errors are computed using a very fine grid. The results show that the nonlinear Schwarz method is quickly able to find an appropriate grid transformation after a few DD iterations. The linearized Schwarz algorithm, as expected, requires more DD iterations but is able to find a quality grid efficiently due to the smaller relative cost per iteration.

Iterations		3	5	7	9	11	$\infty$
Nonlinear	0.3625	0.0498	0.0462	0.0436	0.0449	0.0517	0.0366
Linearized	0.3625	0.1290	0.1019	0.0625	0.0453	0.0435	0.0366

**Table 1.** Interpolation errors for the grids obtained by Schwarz iterations.



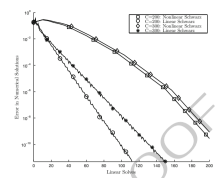


Fig. 3. Linearized Schwarz: error for varying C

4. Non-linear versus Fig. linearized Schwarz with varying C

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The quantities  $\rho$ ,  $D_{\alpha}$  and  $D_{\beta}$  corresponding to iteration (9) and the error estimates in Theorem 1 indicate a dependence on the shape of M for the linearized alternating 135 Schwarz iteration. To test this effect, we consider the performance of (9) for M(x) = 136 $C(x-0.5)^2+1$ . The parameter C controls the ratio a/A. As  $C\to\infty$ ,  $a/A\to0$ , and the contraction rate could diminish. This is demonstrated in Fig. 3. Figure 4 illustrates 138 the effect of changing the value of C on both the nonlinear and linearized Schwarz 139 algorithms. We see that the linearized Schwarz algorithm is affected more by an 140 increase in C.

In summary, we have proposed, analyzed and provided brief numerical comparisons for two alternating Schwarz algorithms to solve the steady grid generation 143 problem using the EP. Ongoing work includes the analysis of DD approaches to 144 moving mesh PDEs for the time dependent mesh generation problem, the discrete 145 analysis and extensions to higher dimensions.

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# Stability Analysis of the Matrix-Free Linearly Implicit 2 **Euler Method**

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Summary. Implicit time stepping methods are useful for the simulation of large scale PDE 9 systems because they avoid the time step limitations imposed by explicit stability conditions. 10 To alleviate the challenges posed by computational and memory constraints, many applica- 11 tions solve the resulting linear systems by iterative methods where the Jacobian-vector products are approximated by finite differences. This paper explains the relation between a linearly 13 implicit Euler method, solved using a Jacobian-free Krylov method, and explicit Runge-Kutta 14 methods. The case with preconditioning is equivalent to a Rosenbrock-W method where the 15 approximate Jacobian, inverted at each stage, corresponds directly to the preconditioner. The 16 accuracy of the resulting Runge-Kutta methods can be controlled by constraining the Krylov 17 solution. Numerical experiments confirm the theoretical findings.

1 Introduction 19

Large systems of time dependent partial differential equations (PDEs), arising in 20 multi-physics simulations, are often discretized using the method of lines approach. 21 The independent time and space numerical schemes allow the coupling of multiple 22 physics modules, and provide maximum flexibility in choosing appropriate algo- 23 rithms. After the semi-discretization in space the system of PDEs is reduced to a 24 system of ordinary differential equations (ODEs) 25

$$y' = f(y), \quad t_0 \le t \le t_f, \quad y(t_0) = y_0.$$
 (1)

Here  $y(t) \in \mathbb{R}^d$  is the solution vector and  $y_0$  the initial condition. We denote the 26 Jacobian of the ODE function by  $J(y) = f_v(y) \in \mathbb{R}^{d \times d}$ , and the identity matrix by 27  $\mathbb{I} \in \mathbb{R}^{d \times d}$ .

Stability requirements (e.g., the CFL condition for discretized hyperbolic PDEs) 29 limit the time steps allowable by explicit time discretizations of (1). When the fastest 30 time scales in the system (1) are short, e.g., in the presence of fast waves, the stability 31 condition imposes time steps much smaller then those required to achieve the target 32

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accuracy. The step size limitation by linear stability conditions is referred to as stiffness. In order to overcome this computational inefficiency, it is desirable to use implicit, unconditionally stable discretizations which allow arbitrarily large time steps 35 [2]. Implicit methods have a high cost per step due to the need to solve a (non)linear 36 system of equations.

To reduce the computational and memory costs of direct linear system solvers, 38 and to aid parallelization, iterative Krylov space methods are employed. Further- 39 more, matrix-free implementations approximate Jacobian vector products by finite 40 differences [4]. This approach avoids additional coding for the Jacobian, preserves 41 the parallel scalability of the explicit model, and has become popular in many appli- 42 cations, e.g., [1, 5, 6]. The hope is that the properties of the implicit time discretization remain unaltered, provided that the iterative solutions are carried out to sufficient 44 accuracy. We show here that the matrix-free approach does alter the properties of the 45 underlying implicit time stepping method.

This study treats a linearly implicit method, together with the Krylov subspace 47 iterations for solving the linear system, as a single numerical scheme. The analysis 48 reveals that matrix-free implementations of linearly implicit methods are equivalent 49 to explicit Runge Kutta methods. Consequently, the unconditional stability property 50 of the base method is lost. When preconditioning is used, the matrix-free implicit 51 methods are equivalent to Rosenbrock-W (ROS-W) methods where the approximate 52 Jacobians correspond directly to the preconditioners.

# 2 The Matrix-Free Linearly Implicit Euler Method

Consider the linearly implicit Euler (LIE) method applied to (1)

$$(\mathbb{I} - \Delta t J(y_n)) \cdot w = f(y_n), \quad y_{n+1} = y_n + \Delta t \cdot w.$$
 (2)

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When the linear system is solved exactly (modulo roundoff errors) by LU factoriza- 56 tion the method (2) is unconditionally stable, and thus suitable for the solution of 57 stiff systems. For many PDEs semi-discretized in the method of lines framework, 58 however, the dimension of the linear system (2) is very large, and the computational 59 and memory costs associated with a direct solution are prohibitive. Moreover, the 60 construction of the explicit Jacobian matrix J is difficult when the space discretization is based on a domain decomposition approach. To alleviate these problems, a 62 popular approach is to solve (2) by matrix-free iterative methods. We seek to analyze 63 the impact that this approximate solutions have on the stability and accuracy of the 64 implicit time stepping scheme. Our approach is to treat the original discretization 65 (2) together with the iterations as a single numerical method applied to solve the 66 ODE(1).

To be specific, we solve the linear system in (2) by a Krylov space method. The 68 initial guess is  $y_{n+1} = y_n$ , i.e., w = 0. After m iterations the following m-dimensional 69 Krylov space is built: 70

$$\mathcal{K}_{m} = \operatorname{span}\left\{f(y_{n}), \dots, \left(\mathbb{I} - \Delta t J(y_{n})\right)^{m-1} f(y_{n})\right\}.$$

In the matrix-free approach, the basis is constructed recursively and the Jacobian- 71 vector products are approximated by finite differences 72

$$\ell_i = \ell_{i-1} - \Delta t \, \varepsilon^{-1} \, f(y_n + \varepsilon \, \ell_{i-1}) + \Delta t \, \varepsilon^{-1} \, \ell_1, \quad i = 2, \dots, m. \tag{3}$$

We assume that the same scaling factor  $\varepsilon$  is used to compute the finite differences in 73 all iterations. (The analysis can be easily extended to the case where a different  $\varepsilon$  is 74 used in each iteration.) Denote 75

$$k_1 = f(y_n); \quad k_i = f(y_n + \varepsilon \ell_{i-1}), \quad i = 2, \dots, m.$$
 (4)

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The recurrence (3) can be expressed in terms of  $k_i$  as:

$$k_i = f\left(y_n + \Delta t \left(\Delta t^{-1} \varepsilon + (i-2)\right) k_1 - \Delta t \sum_{j=2}^{i-1} k_j\right), i = 2, \dots, m.$$
 (5)

The solution  $w = \sum_{i=1}^{m} \alpha_i \ell_i \in \mathcal{K}_m$  can be expressed in terms of  $k_i$ 's:

$$w = \left(\sum_{i=1}^{m} \alpha_i + \Delta t \,\varepsilon^{-1} \,\sum_{i=2}^{m} (i-1) \,\alpha_i\right) k_1 - \Delta t \,\varepsilon^{-1} \,\sum_{i=2}^{m} \left(\sum_{j=i}^{m} \alpha_j\right) k_i. \tag{6}$$

Equations (5) and (6), together with the relation  $y_{n+1} = y_n + \Delta t w$ , are compared with 78 the *m*-stage explicit Runge Kutta (ERK) method [3]

$$k_i = f\left(y_n + \sum_{j=1}^{i-1} a_{ij} k_j\right), i = 1, \dots, m; \quad y_{n+1} = y_n + \Delta t \sum_{i=1}^{m} b_i k_i.$$

The comparison reveals the following.

**Theorem 1.** The matrix-free LIE (2) method is equivalent to an explicit Runge Kutta 81 method. The number m of Krylov iterations defines the number of Runge Kutta stages. 82

Equations (5) and (6) define the coefficients of the ERK method:

$$a_{i,1} = \Delta t^{-1} \varepsilon + (i-2); \ a_{i,j} = -1, \ \text{for } i = 2, \dots, m, \ j = 2, \dots, i-1;$$
  
 $b_1 = \sum_{i=1}^m \alpha_j + \Delta t \ \varepsilon^{-1} \ \sum_{i=2}^m (j-1) \alpha_j; \ b_i = -\Delta t \ \varepsilon^{-1} \ \sum_{i=i}^m \alpha_j, \ i = 2, \dots, m.$ 

### 2.1 Stability Considerations

The solution of the linear system (under the initial guess w = 0) is part of the Krylov space  $\mathcal{K}_m$  and can be represented by a matrix polynomial 86

$$w = p_{m-1} \left( \mathbb{I} - \Delta t J(y_n) \right) \cdot f(y_n).$$

The matrix-free LIE method applied to the Dahlquist test problem  $y' = \lambda y$ , y(0) = 1, 88 gives the following solution:

$$y_{n+1} = y_n + \Delta t w = (1 + z p_{m-1} (1 - z)) y_n = R(z) y_n,$$
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with  $z = \Delta t \lambda$ . The stability function of the equivalent ERK method is the degree m 91 polynomial  $R(z) = 1 + z p_{m-1} (1 - z)$ .

**Theorem 2.** The stability region of the LIE method, with a Krylov matrix-free linear 93 solver, is necessarily finite. The unconditional stability of the original LIE method is 94 lost.

Similar considerations hold for Krylov space methods that use an orthogonal basis 96 of the Krylov space, built by Arnoldi iterations [7].

#### 2.2 Accuracy Considerations

The method accuracy is difficult to assess, as the coefficients depend on the time 99 step. The relation between the finite difference scaling factor  $\varepsilon$  and the time step  $\Delta t$ is important in determining accuracy.

Assume that the finite difference scaling factor is a constant fraction of the time 102 step,  $\varepsilon/\Delta t = \text{const.}$  This is a reasonable assumption: in order to increase accuracy 103 one decreases both  $\Delta t$ , to reduce the truncation error, and  $\varepsilon$ , to reduce the finite 104 difference error. (Of course, for very small  $\varepsilon$  the finite difference error becomes again 105 large due to roundoff.) Also assume that the coefficients  $\alpha_1, \dots, \alpha_m$  do not depend 106 on  $\varepsilon$  or  $\Delta t$ .

In this case the accuracy can be assessed using the classical approach. The order 108 conditions depend on the Krylov space coefficients  $\alpha$  as follows: 109

Order 1: 
$$\sum_{i=1}^{m} b_i = \sum_{j=1}^{m} \alpha_j = 1$$
, (7a)  
Order 2:  $\sum_{i=1}^{m} b_i c_i = -\sum_{i=2}^{m} (i-1) \alpha_i = \frac{1}{2}$ . (7b)

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Order 2: 
$$\sum_{i=1}^{m} b_i c_i = -\sum_{i=2}^{m} (i-1) \alpha_i = \frac{1}{2}.$$
 (7b)

Neither condition (7a) nor (7b) are automatically satisfied by the Krylov iterative 110 methods. In particular, 111

Lemma 1. The first order accuracy of the matrix-free LIE is not automatic when 112  $arepsilon/\Delta t=const.$  Additional constraints need to be imposed on the Krylov solution 113 coefficients. 114

Consider now the case where  $\varepsilon$  is constant (does not depend on  $\Delta t$ ). Assume 115 that the coefficients  $\alpha_1, \ldots, \alpha_m$  do not depend on  $\varepsilon$  or  $\Delta t$ . A necessary condition for the method to be accurate of order p is that its stability function approximates the 117 exponential,  $R(z) = e^z + \mathcal{O}(z^{p+1})$ . The stability function does not depend on either 118  $\varepsilon$  or  $\Delta t$ . The conditions (7a) and (7b) on the Krylov solution coefficients  $\alpha_1, \ldots, \alpha_m$ , 119 which are sufficient when  $\varepsilon = const \cdot \Delta t$ , seem to be necessary in the case  $\varepsilon = const$ . 120

In the general case the Krylov solution coefficients  $\alpha_1, \ldots, \alpha_m$  do depend on  $\Delta t$ . 121 For  $\Delta t \to 0$  we have that  $w \to f(y_n)$  and therefore  $\alpha_1 \to 1, \alpha_2, \alpha_3, \ldots \to 0$ . Asymp- 122 totically the condition (7a) holds. Moreover, the number of iterations m also depends 123 on  $\Delta t$  through the convergence speed. Consequently, it is difficult to extend the classical accuracy analysis to matrix-free linearly implicit methods. It seems reasonable, 125 however, to modify the Krylov method and impose at least condition (7a) on the 126 Krylov coefficients.

#### 3 Preconditioned Iterations

Consider the case where a preconditioner matrix M is used to speed up the iterations. 129 The linear system (2) becomes 130

$$M^{-1}\left(\mathbb{I} - \Delta t J(y_n)\right) \cdot k = M^{-1} f(y_n). \tag{131}$$

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The Krylov space constructed in this case is

$$\mathcal{K}_m = \operatorname{span}\left\{f(y_n) \ldots, \left(M^{-1}\left(\mathbb{I} - \Delta t J(y_n)\right)\right)^{m-1} M^{-1} f(y_n)\right\}$$

In the matrix-free approach the following basis is constructed recursively

$$\ell_1 = M^{-1} f(y_n), \ell_i = M^{-1} \ell_{i-1} - \Delta t \, \varepsilon^{-1} M^{-1} f(y_n + \varepsilon \ell_{i-1}) + \Delta t \, \varepsilon^{-1} \ell_1, \quad i = 2, \dots, m.$$

Denote  $k_1 = \Delta t \ \ell_1$  and  $k_i = \Delta t \ \ell_1 - \varepsilon \ell_i$  for  $i = 2, \dots, m$ . We have

$$Mk_1 = \Delta t f(y_n)$$

$$Mk_i = \Delta t f(y_n + k_1 - k_{i-1}) + k_{i-1} - k_1, i = 2, ..., m.$$
(8)

Consider, for comparison, a Rosenbrock-W (ROW) method in the implementationfriendly formulation [2, Sect. IV.7]

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$$\left[\mathbb{I} - \Delta t \, \gamma \widehat{J}_n\right] k_i = \Delta t \, \gamma f\left(y_n + \sum_{j=1}^{i-1} a_{ij} k_j\right) + \gamma \sum_{j=1}^{i-1} c_{ij} k_j,$$

$$y_{n+1} = y_n + \sum_{i=1}^{s} m_i k_i. \tag{9}$$

Here  $\widehat{J_n} \approx J(y_n)$  is an approximation of the exact Jacobian at the current step. We dentify the method coefficients  $\gamma = 1$  and

$$c_{i,1} = -1$$
;  $c_{i,i-1} = 1$ ;  $a_{i,1} = 1$ ;  $a_{i,i-1} = -1$ ,  $i = 2, \dots, m$ .

From the solution  $w = \sum_{i=1}^{m} \alpha_i \ell_i = \sum_{i=1}^{m} b_i k_i \in \mathcal{K}_m$  we identify the weights

$$b_1 = \alpha_1 \Delta t^{-1} + \varepsilon^{-1} \sum_{i=2}^{m} \alpha_j; \quad b_i = -\varepsilon^{-1} \alpha_i, \ i = 2, ..., m.$$

The preconditioner defines the Jacobian approximation in the ROW method,

$$M = \mathbb{I} - \Delta t \, \gamma \widehat{J}_n \quad \Rightarrow \quad \widehat{J}_n = \Delta t^{-1} \, (\mathbb{I} - M) \; .$$

**Theorem 3.** The preconditioned matrix-free LIE is equivalent to a linearly-implicit 141 ROW method. The choice of the preconditioner, besides accelerating convergence, 142 improves the stability of the matrix-free LIE method. The preconditioner defines the 143 Jacobian approximation in the ROW method.

Note that the general approach can be applied to ROW methods [2, Sect. IV.7] 145 by solving the linear system of each stage with an iterative matrix free algorithm. 146 The resulting scheme is an explicit Runge Kutta method (or a ROW method) with 147  $\sum_{i=1}^{s} m_i$  stages. 148

#### 4 Numerical Results

Consider the one dimensional scalar advection-diffusion equation

 $u_t + (au)_x = Du_{xx}, u(x,t=0) = u_0(x).$ (10)

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A spectral discontinuous Galerkin spatial discretization is used with 20 elements and 151 polynomials of order 8. The diffusive term discretization is stabilized using the internal penalty method [8]. The LIE time stepping is used with the matrix-free GMRES 153 solver [7].

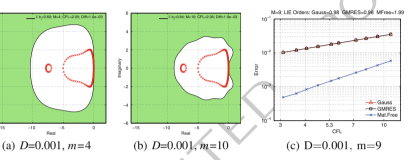


Fig. 1. (a) and (b) The ERK stability regions for different numbers of GMRES iterations. (c) The accuracy of the LIE scheme using various approaches to invert the Jacobian matrix. The GMRES weights are restricted by (7b) such as to obtain a second order method. Advectiondiffusion equation (10),  $\Delta t = CFL$ ,  $\varepsilon = 10^{-6} \Delta t$ 

In Fig. 1a, b, the stability regions generated by the GMRES iterations are plotted 155 for a varying number of Krylov vectors. The regions grow quickly and encompass 156 the eigenvalues of the discrete advection-diffusion operator. Subsequent iterations 157 improve solution accuracy but do not improve linear stability. Additional experi- 158 ments (not reported here due to space constraints) reveal that the stability region of 159 the resulting ERK method adapts to the eigenvalues of different discrete operators.

To verify the analysis in (7), we consider three different ways of computing the 161 inverse of the linear Jacobian. The first is by Gauss elimination (LU), the second 162 uses GMRES with the full Jacobian, and the third employs matrix-free GMRES it- 163 erations. In the last approach the GMRES coefficients are restricted by (7b) such 164 as to obtain a second order time discretization method. Figure 1c shows the workprecision diagram for these approaches. The Gaussian elimination and traditional 166 GMRES solutions display first order converge, while the constrained GMRES solution displays second order convergence.

5 Conclusions 169

Implicit time integration methods are becoming widely used in the the simulation of 170 time dependent PDEs, as they do not suffer from CFL stability restrictions. While 171

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implicit methods can use much larger time steps than explicit methods, their computational cost per step is also higher. The computational time is dominated by the 173 solutions of (non)linear systems of equations that define each stage of a (linearly) 174 implicit method. The implicit code is more effective only when the gains in step size 175 offset the extra cost.

To reduce the computational overhead of LU decomposition, to alleviate memory 177 requirements, and to aid parallelization, iterative Krylov space methods are used 178 to solve the large linear systems. A matrix-free implementation approximates the 179 required Jacobian vector products by finite differences.

This paper studies the effect of the matrix-free iterative solutions on the properties of the numerical integration method. The analysis reveals that matrix-free lin- 182 early implicit methods can be viewed as explicit Runge Kutta methods. Their stabil- 183 ity region is finite, and the unconditional stability property of the original implicit 184 method is lost. The equivalent Runge Kutta method is nonlinear, in the sense that 185 its weights depend on the time step and on the stage vectors. This makes the ac- 186 curacy analysis difficult. Order conditions of the equivalent explicit Runge Kutta 187 method can be fulfilled by imposing additional conditions on the Krylov solution 188 coefficients. For preconditioned matrix-free iterations the overall time stepping pro- 189 cess is equivalent to a Rosenbrock-W method, where the preconditioner determines 190 the Jacobian approximation. Future work will address the effect of a finite number 191 of Krylov iterations on the stability and accuracy of the overall scheme, in the case 192 where an analytical Jacobian is used.

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# **Augmented Interface Systems for the Darcy-Stokes Problem**

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Summary. In this paper we study interface equations associated to the Darcy-Stokes problem 8 using the classical Steklov-Poincaré approach and a new one called augmented. We compare 9 these two families of methods and characterize at the discrete level suitable preconditioners 10 with additive and multiplicative structures. Finally, we present some numerical results to assess their behavior in presence of small physical parameters.

# 1 Introduction and Problem Setting

Let  $\Omega \subset \mathbb{R}^d$  (d=2,3) be a bounded domain decomposed into two non intersecting subdomains:  $\Omega_f$ , filled by a viscous incompressible fluid, and  $\Omega_p$ , formed by a 15 porous medium, separated by an interface  $\Gamma = \bar{\Omega}_f \cap \bar{\Omega}_p$ . The fluid in  $\Omega_f$  has no free 16 surface and it can filtrate through the adjacent porous medium. The motion of the 17 fluid in  $\Omega_f$  is described by the Stokes equations:

$$- \nu \triangle \mathbf{u} + \nabla p = \mathbf{f}, \quad \text{div } \mathbf{u} = 0 \quad \text{in } \Omega_f$$
 (1)

where v > 0 is the kinematic viscosity, while **u** and p are the velocity and pressure. 19 In  $\Omega_p$  we describe the fluid motion by the equations:

$$\mathbf{u}_p = -\mathsf{K}\nabla\varphi, \quad \text{div } \mathbf{u}_p = 0 \quad \text{in } \Omega_p$$
 (2)

where  $\mathbf{u}_p$  is the fluid velocity,  $\varphi$  the piezometric head and K the hydraulic conductiv- 21 ity tensor. The first equation is Darcy's law that provides the simplest linear relation 22 between velocity and pressure in porous media. We can equivalently rewrite (2) as 23 the elliptic equation involving only the piezometric head: 24

$$-\operatorname{div}(\mathsf{K}\nabla\varphi)=0\quad\text{in }\Omega_p. \tag{3}$$

Besides suitable boundary conditions on  $\partial \Omega$ , we supplement the Darcy-Stokes 25 problem (1), (3) with the following coupling conditions on  $\Gamma$ : 26

$$-\mathsf{K}\nabla\varphi\cdot\mathbf{n} = \mathbf{u}\cdot\mathbf{n}, \quad -\mathbf{n}\cdot\mathsf{T}(\mathbf{u},p)\cdot\mathbf{n} = g\varphi, \quad -\varepsilon\boldsymbol{\tau}\cdot\mathsf{T}(\mathbf{u},p)\cdot\mathbf{n} = v\mathbf{u}\cdot\boldsymbol{\tau}, \quad (4)$$

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where  $T(\mathbf{u}, p)$  is the fluid stress tensor,  $\boldsymbol{\tau}$  denotes a set of linear independent unit 27 tangential vectors to  $\Gamma$  and  $\varepsilon$  is a coefficient related to the characteristic length of 28 the pores of the porous medium. Conditions  $(4)_1$  and  $(4)_2$  impose the continuity of 29 the normal velocity and of the normal component of the normal stress on  $\Gamma$ . The 30 so-called Beavers-Joseph-Saffman condition (4)<sub>3</sub> does not yield any coupling but 31 provides a boundary condition for the Stokes problem since it involves only quantities in the domain  $\Omega_f$ . For more details we refer to [9, 11, 12, 14].

## 2 Interface Equations Associated to the Darcy-Stokes Problem

In [7, 8], we showed that the coupled Darcy-Stokes problem can be reformulated 35 in terms of the solution of equations defined only on the interface  $\Gamma$  involving suitable Steklov-Poincaré operators associated to the subproblems in  $\Omega_f$  and  $\Omega_p$ . We 37 formally briefly review this approach referring to the cited works for more details.

If we select as interface variable  $\lambda \in H_{00}^{1/2}(\Gamma)$  to represent the normal velocity 39 across  $\Gamma$ :  $\lambda = \mathbf{u} \cdot \mathbf{n} = -\mathsf{K} \nabla \varphi \cdot \mathbf{n}$  on  $\Gamma$ , we can express the solution of the Darcy-Stokes problem in terms of the solution of the interface equation: find  $\lambda \in H^{1/2}_{00}(\Gamma)$  41 such that

$$\langle S_s \lambda, \mu \rangle + \langle S_d \lambda, \mu \rangle = \langle \chi_s, \mu \rangle + \langle \chi_d, \mu \rangle \qquad \forall \mu \in H_{00}^{1/2}(\Gamma).$$
 (5)

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Equation (5) imposes the continuity condition  $(4)_2$ . The linear continuous operators 43  $\chi_s$  and  $\chi_d$  depend on the data of the problem and  $\langle \cdot, \cdot \rangle$  denotes the duality pairing 44 between  $H_{00}^{1/2}(\Gamma)$  and its dual  $(H_{00}^{1/2}(\Gamma))'$ . Concerning  $S_s$  and  $S_d$ , we remark that

- The operator  $S_s: H^{1/2}_{00}(\Gamma) \to (H^{1/2}_{00}(\Gamma))'$  maps the space of normal velocities on 46  $\Gamma$  to the space of normal stresses on  $\Gamma$  through the solution of a Stokes problem 47 in  $\Omega_f$  with boundary condition  $\mathbf{u} \cdot \mathbf{n} = \lambda$  on  $\Gamma$ .
- $S_d$  maps the space of fluxes of  $\varphi$  on  $\Gamma$  to the space of traces of  $\varphi$  on  $\Gamma$  via the 49 solution of a Darcy problem in  $\Omega_p$  with the boundary condition  $-\mathsf{K}\nabla\varphi\cdot\mathbf{n}=\lambda_{0}$ on  $\Gamma$ . The operator  $S_d$  should be a map between  $H^{-1/2}(\Gamma)$  and  $H^{1/2}(\Gamma)$ , but in 51 (5) we are applying it to  $H_{00}^{1/2}(\Gamma)$ , a space with a higher regularity than needed 52 where we cannot guarantee the coercivity of the operator.

On the other hand, if we choose as interface unknown  $\eta \in H^{1/2}(\Gamma)$  the trace 54 of the piezometric head on  $\Gamma$ :  $\eta = g\varphi_{\Gamma} = -\mathbf{n} \cdot \mathsf{T}(\mathbf{u}, p) \cdot \mathbf{n}$  on  $\Gamma$ , the Darcy-Stokes 55 problem can be equivalently reformulated as find  $\eta \in H^{1/2}(\Gamma)$ : 56

$$\langle\!\langle S_f \eta, \mu \rangle\!\rangle + \langle\!\langle S_p \eta, \mu \rangle\!\rangle = \langle\!\langle \chi_f, \mu \rangle\!\rangle + \langle\!\langle \chi_p, \mu \rangle\!\rangle \qquad \forall \mu \in H^{1/2}(\Gamma), \tag{6}$$

where  $\chi_f$  and  $\chi_p$  are linear continuous operators depending on the data of the problem. Equation (6) imposes the coupling condition  $(4)_1$ . Here: 58

• The operator  $S_f$  maps the space of normal stresses on  $\Gamma$  to the space of normal 59 velocities on  $\Gamma$  via the solution of a Stokes problem with the boundary condition  $-\mathbf{n} \cdot \mathsf{T}(\mathbf{u}, p) \cdot \mathbf{n} = \eta$  on  $\Gamma$ . This operator would naturally be defined from 61

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 $H^{-1/2}(\Gamma)$  to  $H^{1/2}_{00}(\Gamma)$  so that in (6) we are applying it to functions with a higher 62 regularity than needed.

• The operator  $S_p: H^{1/2}(\Gamma) \to (H^{1/2}(\Gamma))'$  maps the space of traces of  $\varphi$  on  $\Gamma$  64 to the space of fluxes of  $\varphi$  on  $\Gamma$  by solving a Darcy problem in  $\Omega_p$  with the 65 Dirichlet boundary condition  $g\varphi = \eta$  on  $\Gamma$ .

## 3 Augmented Interface Equations

The classical approach summarized in Sect. 2 leads to reformulate the Darcy-Stokes 68 problem as interface equations depending on a single interface unknown: either  $\lambda$ , 69 the normal velocity across  $\Gamma$ , or  $\eta$ , the piezometric head on  $\Gamma$ . We have remarked 70 that the Steklov-Poincaré operators  $S_d$  and  $S_f$  are not acting on their natural functional spaces, but they are assigned functions with higher regularity than expected. 72 This prevents us from guaranteeing their coerciveness (see [7]). In this section we 73 present a different approach based on [3–6] consisting in writing the coupled Darcy-74 Stokes problem as a system of linear equations on  $\Gamma$  involving both variables  $\lambda$  75 and  $\eta$ .

#### 3.1 The Augmented Dirichlet-Dirichlet Problem

To obtain the augmented Dirichlet-Dirichlet (aDD) formulation assume that  $\lambda \in 78$   $H_{00}^{1/2}(\Gamma)$  is equal to the normal velocity  $\mathbf{u} \cdot \mathbf{n}$  on  $\Gamma$ , but not necessarily to the conormal derivative of  $\varphi$  on  $\Gamma$ . On the other hand, let  $\eta \in H^{1/2}(\Gamma)$  be equal to the trace of 80  $\varphi$  on  $\Gamma$  but not to the normal component of the Cauchy stress of the Stokes problem 81 on  $\Gamma$ . Then, to recover the solution of the original Darcy-Stokes problem we have to 82 impose both the continuity of normal velocity and of normal stresses: 83

$$\begin{split} &-\int_{\Gamma}\mathbf{n}\cdot \mathbf{\Gamma}(\mathbf{u}(\lambda),p(\lambda))\cdot\mathbf{n}\,\mu = \int_{\Gamma}\eta\mu &\quad \forall \mu\in H_{00}^{1/2}(\Gamma) \\ &-\int_{\Gamma}\mathsf{K}\nabla\varphi(\eta)\cdot\mathbf{n}\,\xi = \int_{\Gamma}\lambda\xi &\quad \forall \xi\in H^{1/2}(\Gamma). \end{split}$$

Using the definition of the Steklov-Poincaré operators, we can rewrite these conditions as: find  $(\lambda, \eta) \in H^{1/2}_{00}(\Gamma) \times H^{1/2}(\Gamma)$  such that

$$\langle S_s \lambda, \mu \rangle + \langle \eta, \mu \rangle = \langle \chi_s, \mu \rangle \qquad \forall \mu \in H_{00}^{1/2}(\Gamma) \langle S_p \eta, \xi \rangle - \langle \chi, \xi \rangle = \langle \chi_p, \xi \rangle \qquad \forall \xi \in H^{1/2}(\Gamma),$$
(7)

or, in operator form:

$$\begin{pmatrix} S_s & \mathscr{I} \\ -\mathscr{J} & S_p \end{pmatrix} \begin{pmatrix} \lambda \\ \eta \end{pmatrix} = \begin{pmatrix} \chi_s \\ \chi_p \end{pmatrix} \tag{8}$$

where  $\mathscr{I}: H^{1/2}(\Gamma) \to (H^{1/2}_{00}(\Gamma))'$  and  $\mathscr{J}: H^{1/2}_{00}(\Gamma) \to (H^{1/2}(\Gamma))'$  are linear continuous maps.

We call (8) *augmented Dirichlet-Dirichlet* (aDD) formulation because both functions  $\lambda$  and  $\eta$  play the role of Dirichlet boundary conditions for the Stokes and the 90 Darcy subproblems, respectively. Notice that we are imposing the equalities (8) in 91 the sense of dual spaces and that the operators  $S_s$  and  $S_p$  still act on their natural 92 functional spaces.

#### 3.2 The Augmented Neumann-Neumann Problem

We follow now a similar approach to Sect. 3.1, but we assume that  $\lambda \in H^{-1/2}(\Gamma)$  95 is equal to the conormal derivative of the piezometric head  $-\mathsf{K}\nabla\varphi\cdot\mathbf{n}$  on  $\Gamma$  and  $\eta\in$  96  $H^{-1/2}(\Gamma)$  is equal to the normal component of the fluid Cauchy stress on  $\Gamma$ . Then, 97 to recover the solution of the original problem we impose the following equalities: 98

$$\int_{\Gamma} \mathbf{u}(\eta) \cdot \mathbf{n} \mu = \int_{\Gamma} \lambda \mu \quad \forall \mu \in H^{-1/2}(\Gamma)$$

$$\int_{\Gamma} \varphi(\lambda) \xi = -\int_{\Gamma} \eta \xi \quad \forall \xi \in H^{-1/2}(\Gamma).$$

Using the definition of the Steklov-Poincaré operators, we can rewrite these conditions as: find  $(\lambda, \eta) \in H^{-1/2}(\Gamma) \times H^{-1/2}(\Gamma)$  such that

$$\langle S_f \eta, \mu \rangle_* - \langle \lambda, \mu \rangle_* = \langle \chi_f, \mu \rangle_* \qquad \forall \mu \in H^{-1/2}(\Gamma) \langle S_d \lambda, \xi \rangle_* + \langle \eta, \xi \rangle_* = \langle \chi_d, \xi \rangle_* \qquad \forall \xi \in H^{-1/2}(\Gamma),$$

$$(9)$$

corresponding to the operator form:

$$\begin{pmatrix} S_d & \mathscr{I}_* \\ -\mathscr{I}_* & S_f \end{pmatrix} \begin{pmatrix} \lambda \\ \eta \end{pmatrix} = \begin{pmatrix} \chi_d \\ \chi_f \end{pmatrix}. \tag{10}$$

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Here  $\mathscr{I}_*: H^{-1/2}(\Gamma) \to H^{1/2}(\Gamma)$  and  $\mathscr{J}_*: H^{-1/2}(\Gamma) \to H^{1/2}_{00}(\Gamma)$  are linear continuous maps, while  $\langle \cdot, \cdot \rangle_*$  and  $\langle \cdot, \cdot \rangle_*$  denote the corresponding pairing.

We call this formulation *augmented Neumann-Neumann* (aNN) because both 104 functions  $\lambda$  and  $\eta$  play the role of Neumann boundary conditions for the Darcy 105 and the Stokes subproblems, respectively.

The aNN formulation may be regarded as the "dual" of the aDD approach. Notice that the operators  $S_f$  and  $S_d$  are now acting on their natural spaces, differently form the classical setting of Sect. 2. The analysis of problems (8) and (10) can be carried out following the guidelines of [5].

# 4 Algebraic Formulation of the Interface Problems

We consider a finite element discretization of the coupled problem using conforming grids across the interface  $\Gamma$ . The discrete spaces for the Stokes problem satisfy the inf-sup condition. In this way we obtain the linear system:

$$\begin{pmatrix}
F & D & 0 & 0 \\
D^{T} A_{\Gamma\Gamma} & 0 & -M_{\Gamma} \\
0 & 0 & C_{ii} & C_{i\Gamma} \\
0 & M_{\Gamma}^{T} C_{\Gamma i} & C_{\Gamma\Gamma}
\end{pmatrix}
\begin{pmatrix}
\mathbf{u}_{i} \\
\mathbf{u}_{\Gamma} \\
\mathbf{\phi}_{i} \\
\mathbf{\phi}_{\Gamma}
\end{pmatrix} = \begin{pmatrix}
\mathbf{f}_{fi} \\
\mathbf{f}_{f\Gamma} \\
\mathbf{f}_{pi} \\
\mathbf{f}_{p\Gamma}
\end{pmatrix}$$
(11)

where  $u_{\Gamma}$  is the vector of the nodal values of the normal velocity on  $\Gamma$  while  $u_i$  is the vector of the remaining degrees of freedom (velocity and pressure) in  $\Omega_f$ . On the other hand,  $\varphi_{\Gamma}$  is the vector of the (unknown) values of  $\varphi$  on  $\Gamma$  while  $\varphi_i$  corresponds to the remaining degrees of freedom in  $\Omega_p$ .

The discrete counterpart of the Steklov-Poincaré operators can be found computing the Schur complement systems corresponding to either  $u_{\Gamma}$  or  $\varphi_{\Gamma}$ . Precisely, we 120 find: 121

 $\Sigma_{s} = A_{\Gamma\Gamma} - D^{T}F^{-1}D, \qquad \Sigma_{f} = M_{\Gamma}^{T}\Sigma_{s}^{-1}M_{\Gamma},$  $\Sigma_{p} = C_{\Gamma\Gamma} - C_{\Gamma i}C_{ii}^{-1}C_{i\Gamma}, \qquad \Sigma_{d} = M_{\Gamma}\Sigma_{p}^{-1}M_{\Gamma}^{T}.$ (12)

The characterization of these discrete operators in terms of the associated Darcy or 122 Stokes problems in  $\Omega_p$  and  $\Omega_f$  allows us to provide upper and lower bounds for 123 their eigenvalues. Assuming v and K constants in  $\Omega_f$  and  $\Omega_p$ , respectively, and the computational mesh to be uniform and regular, we can find (see [7, 13, 15]) ( $\prec$ indicates that the inequalities hold up to constants independent of h, v, K: 126

$$hv \leq \sigma(\Sigma_s) \leq v, \qquad h^2 v^{-1} \leq \sigma(\Sigma_f) \leq hv^{-1}$$
  
$$h\mathsf{K} \leq \sigma(\Sigma_p) \leq \mathsf{K}, \qquad h^2 \mathsf{K}^{-1} \leq \sigma(\Sigma_d) \leq h\mathsf{K}^{-1}$$
 (13)

The discrete counterparts of the interface problems (5), (6), (8), and (10) read: 127

• Discrete interface equation for the normal velocity: find  $u_{\Gamma}$  such that

$$\Sigma_{s} \boldsymbol{u}_{\Gamma} + \Sigma_{d} \boldsymbol{u}_{\Gamma} = \boldsymbol{\chi}_{s} + \boldsymbol{\chi}_{d}. \tag{14}$$

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• Discrete interface equation for the piezometric head: find  $\phi_{\Gamma}$  such that

$$\Sigma_f \boldsymbol{\varphi}_{\Gamma} + \Sigma_p \boldsymbol{\varphi}_{\Gamma} = \boldsymbol{\chi}_f + \boldsymbol{\chi}_p. \tag{15}$$

• Discrete aDD problem: find  $(\boldsymbol{u}_{\Gamma}, \boldsymbol{\varphi}_{\Gamma})$  such that

$$\begin{pmatrix} \Sigma_s & -M_{\Gamma} \\ M_{\Gamma}^T & \Sigma_p \end{pmatrix} \begin{pmatrix} \mathbf{u}_{\Gamma} \\ \mathbf{\varphi}_{\Gamma} \end{pmatrix} = \begin{pmatrix} \mathbf{\chi}_s \\ \mathbf{\chi}_p \end{pmatrix}. \tag{16}$$

• Discrete aNN problem: find  $(\boldsymbol{u}_{\Gamma}, \boldsymbol{\varphi}_{\Gamma})$  such that

$$\begin{pmatrix} \Sigma_d & M_{\Gamma} \\ -M_{\Gamma}^T & \Sigma_f \end{pmatrix} \begin{pmatrix} \boldsymbol{u}_{\Gamma} \\ \boldsymbol{\varphi}_{\Gamma} \end{pmatrix} = \begin{pmatrix} \boldsymbol{\chi}_d \\ \boldsymbol{\chi}_f \end{pmatrix}. \tag{17}$$

The augmented approach allows to compute both interface variable at once but it 132 requires to solve a system whose dimension is twice the one of the classical methods. 133

## 5 Iterative Solution Methods and Numerical Results

We present now some numerical methods to solve problems (14)–(17) focusing on 135 cases where the fluid viscosity v and the hydraulic conductivity K are small. These 136 are indeed situations of interest for most practical applications. In [10] a Robin- 137 Robin method was proposed to solve effectively (14). Here we adopt the generalized 138 Hermitian/skew-Hermitian splitting (GHSS) method of [2] for (14) and (15) and the 139 HSS method of [1] for (16) and (17). We start considering (14).

The matrix  $\Sigma_s + \Sigma_d$  has no skew-symmetric component being symmetric positive 141 definite, but thanks to the estimates (13) we can mimick the splitting proposed in [2] 142 considering  $\Sigma_s$  as a matrix multiplied by a coefficient (v) which may become small. 143 Thus, we can characterize the preconditioner for (14):

$$P_1 = (2\alpha_1)^{-1} (\Sigma_s + \alpha_1 I) (\Sigma_d + \alpha_1 I). \tag{18}$$

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Proceeding analogously for (15), we can characterize the preconditioner

$$P_2 = (2\alpha_2)^{-1} (\Sigma_p + \alpha_2 I)(\Sigma_f + \alpha_2 I). \tag{19}$$

Preconditioners  $P_1$  and  $P_2$  involve suitable acceleration parameters  $\alpha_1$  and  $\alpha_2$  146 and can be used within GMRES iterations. Remark that they can be regarded as 147 generalizations of the Robin-Robin method introduced in [7, 10].

On the other hand, as the matrices in (16) and (17) are positive skew-symmetric 149 with symmetric positive definite diagonal blocks, we apply the HSS splitting pro- 150 posed in [1] separating the symmetric and the skew-symmetric parts of the matrices. 151 Thus, we can characterize the following preconditioners for GMRES iterations for 152 (16) and (17), respectively, with  $\alpha_3$ ,  $\alpha_4$  suitable acceleration parameters:

$$P_3 = (2\alpha_3)^{-1} \begin{pmatrix} \Sigma_s + \alpha_3 I & 0\\ 0 & \Sigma_p + \alpha_3 I \end{pmatrix} \begin{pmatrix} \alpha_3 I - M_{\Gamma}\\ M_{\Gamma}^T & \alpha_3 I \end{pmatrix}$$
(20)

$$P_4 = (2\alpha_4)^{-1} \begin{pmatrix} \Sigma_d + \alpha_4 I & 0 \\ 0 & \Sigma_f + \alpha_4 I \end{pmatrix} \begin{pmatrix} \alpha_4 I & M_{\Gamma} \\ -M_{\Gamma}^T & \alpha_4 I \end{pmatrix}. \tag{21}$$

According to [2] these preconditioners are effective when either the skew-symmetric 155 or the symmetric part dominates. Thanks to (13) we can expect that for small  $\nu$  and 156 K the skew-symmetric part dominates in (16) and the symmetric one in (17).

All preconditioners  $P_i$  require the solution of a Stokes problem in  $\Omega_f$  and of a 158 Darcy problem in  $\Omega_p$ . However,  $P_1$  and  $P_2$  have a multiplicative structure while in 159  $P_3$  and  $P_4$  the two subproblems may be solved in a parallel fashion. They are all 160 effective when v and K become small. A thorough study of these preconditioners 161 will make the object of a future work, where also the choice of the parameters  $\alpha_i$  162 will be analyzed. For the tests reported in Table 1, following [2], we set  $\alpha_1, \alpha_3 \simeq \sqrt{\nu}$ , 163  $\alpha_2 \simeq \sqrt{K}$  and  $\alpha_4 \simeq 10^{-1}$ . However, a better characterization of such parameters is 164 necessary to have a more robust behavior of the preconditioners, independent of both 165 the mesh size and of the coefficients v and K.

In the numerical tests, both the Stokes and the Darcy subproblems are solved 167 via direct methods. The matrices in (20) and (21) involving  $M_{\Gamma}$  and I are assembled explicitly and the associated linear systems are solved using direct methods. 169 We consider  $\Omega_f = (0,1) \times (1,2)$ ,  $\Omega_p = (0,1)^2$  with interface  $\Gamma = (0,1) \times \{1\}$  and 170 the analytic solution:  $\mathbf{u} = ((y-1)^2 + (y-1) + 1, x(x-1)), \ p = 2v(x+y-1), \ 171$  $\varphi = K^{-1}(x(1-x)(y-1)+(y-1)^3/3)+2\nu x$ . A comparison with preconditioners 172  $\Sigma_s$  for (14) and  $\Sigma_p$  for (15) studied in [7] is also presented. Although such preconditioners are optimal with unitary  $\nu$  and K, they perform quite poorly when small 174 viscosities and permeabilities are considered.

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**Table 1.** Number of iterations to solve (14)-(17) using different preconditioners. Four computational meshes  $(h_i = 2^{-(j+1)})$  and several values of v and K have been considered.

#### GMRES iterations without and with preconditioner for (14) ( $tol = 10^{-7}$ ).

_	$v = 10^{-4}, K = 10^{-3}$			$v = 10^{-6}, K = 10^{-5}$			$v = 10^{-6},  K = 10^{-8}$			
-	No prec.	$\Sigma_s$	$P_1$	No prec.	$\Sigma_s$	$P_1$	No prec.	$\Sigma_s$	P <sub>1</sub>	
$\overline{h_1}$	8		$4 (\alpha_1 = 10^{-2})$		8	$3 (\alpha_1 = 10^{-3})$	8			t2.1
$h_2$	16	15	$5 (\alpha_1 = 10^{-2})$	16	15	$3 (\alpha_1 = 10^{-3})$	16	15	$3 (\alpha_1 = 10^{-3})$	t2.2
$h_3$	26	20	$7 (\alpha_1 = 10^{-3})$	26	20	$3 (\alpha_1 = 10^{-3})$	26		$3 (\alpha_1 = 10^{-3})$	
$h_4$	33	17	$7 (\alpha_1 = 10^{-3})$	33	17	$4 (\alpha_1 = 10^{-3})$	33	17	$3 (\alpha_1 = 10^{-3})$	t2.4

#### GMRES iterations without and with preconditioner for (15) ( $tol = 10^{-7}$ )

_	$v = 10^{-4}, K = 10^{-3}$			$v = 10^{-6}, K = 10^{-5}$			$v = 10^{-6}, K = 10^{-8}$			
_	No prec.	$\Sigma_p$	$P_2$	No prec.	$\Sigma_p$	$P_2$	No prec.	$\Sigma_p$	$P_2$	
$\overline{h_1}$	9		6 $(\alpha_2 = 10^{-2})$	9	9	$4 (\alpha_2 = 10^{-3})$	()- ·		$3 (\alpha_2 = 10^{-3})$	
$h_2$	17		$7 (\alpha_2 = 10^{-2})$	17	17	$4 (\alpha_2 = 10^{-3})$			$3 (\alpha_2 = 10^{-3})$	
$h_3$	32	31	$8 (\alpha_2 = 10^{-2})$	33	33	$5 (\alpha_2 = 10^{-3})$	33	33	$4 (\alpha_2 = 10^{-3})$	t4.3
$h_4$	46	42	$8 (\alpha_2 = 10^{-2})$	59	57	$5 (\alpha_2 = 10^{-3})$	63	62	$4 (\alpha_2 = 10^{-3})$	t4.4

## GMRES iterations without and with preconditioner $P_3$ for (16) ( $tol = 10^{-9}$ ).

			-			
v = 10	$0^{-4}$ , K = $10^{-3}$	v = 1	$0^{-6}$ , $K = 10^{-5}$	v = 10		
No prec.	$P_3$	No prec.	. P <sub>3</sub>	No prec.	P <sub>3</sub>	
17	14 $(\alpha_3 = 10^{-3})$	17	$7 (\alpha_3 = 10^{-3})$	17	8 $(\alpha_3 = 10^{-3})$	t6.1
33	17 $(\alpha_3 = 10^{-3})$	33	8 $(\alpha_3 = 10^{-3})$	33	10 $(\alpha_3 = 10^{-3})$	t6.2
63	22 $(\alpha_3 = 5 \cdot 10^{-4})$	) 65	$8 (\alpha_3 = 5 \cdot 10^{-4})$	65	10 $(\alpha_3 = 5 \cdot 10^{-4})$	t6.3
67	23 $(\alpha_3 = 5 \cdot 10^{-4})$	) 79	9 $(\alpha_3 = 5 \cdot 10^{-4})$	101	11 $(\alpha_3 = 5 \cdot 10^{-4})$	t6.4
	No prec. 17 33 63	17	No prec. $P_3$ No prec.       17     14 ( $\alpha_3 = 10^{-3}$ )     17       33     17 ( $\alpha_3 = 10^{-3}$ )     33       63     22 ( $\alpha_3 = 5 \cdot 10^{-4}$ )     65	No prec. $P_3$ No prec. $P_3$ 17     14 ( $\alpha_3 = 10^{-3}$ )     17     7 ( $\alpha_3 = 10^{-3}$ )       33     17 ( $\alpha_3 = 10^{-3}$ )     33     8 ( $\alpha_3 = 10^{-3}$ )       63     22 ( $\alpha_3 = 5 \cdot 10^{-4}$ )     65     8 ( $\alpha_3 = 5 \cdot 10^{-4}$ )	No prec. $P_3$ No prec. $P_3$ No prec.           17         14 ( $\alpha_3 = 10^{-3}$ )         17         7 ( $\alpha_3 = 10^{-3}$ )         17           33         17 ( $\alpha_3 = 10^{-3}$ )         33         8 ( $\alpha_3 = 10^{-3}$ )         33           63         22 ( $\alpha_3 = 5 \cdot 10^{-4}$ )         65         8 ( $\alpha_3 = 5 \cdot 10^{-4}$ )         65	No prec. $P_3$ No prec. $P_3$ No prec. $P_3$ 17         14 ( $\alpha_3 = 10^{-3}$ )         17         7 ( $\alpha_3 = 10^{-3}$ )         17         8 ( $\alpha_3 = 10^{-3}$ )           33         17 ( $\alpha_3 = 10^{-3}$ )         33         8 ( $\alpha_3 = 10^{-3}$ )         33         10 ( $\alpha_3 = 10^{-3}$ )           63         22 ( $\alpha_3 = 5 \cdot 10^{-4}$ )         65         8 ( $\alpha_3 = 5 \cdot 10^{-4}$ )         65         10 ( $\alpha_3 = 5 \cdot 10^{-4}$ )

#### GMRES iterations without and with preconditioner $P_4$ for (17) ( $tol = 10^{-9}$ ).

	$v = 10^{-4}, K = 10^{-3}$	$v = 10^{-}$	$^{6}$ , K = $10^{-5}$	$v = 10^{-6}, K = 10^{-8}$		
	No prec. $P_4$	No prec.	$P_4$	No prec.	$P_4$	
$h_1$	17 16 $(\alpha_4 = 0.1)$	16	9 $(\alpha_4 = 0.5)$	9	$8 (\alpha_4 = 1)$	
$h_2$	32 18 ( $\alpha_4 = 0.1$ )	32	8 $(\alpha_4 = 0.5)$	16	7 $(\alpha_4 = 0.5)$	
$h_3$	59 $20 \ (\alpha_4 = 5 \cdot 10^{-2})$	58	10 $(\alpha_4 = 0.1)$	30	5 $(\alpha_4 = 0.8)$	
$h_4$	82 $27 (\alpha_4 = 5 \cdot 10^{-2})$	81	8 $(\alpha_4 = 0.1)$	44	$5 (\alpha_4 = 0.8)$	

t8.1 t8.2 t8.3 t8.4

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# **Mortar Coupling for Heterogeneous Partial Differential Equations**

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1 Introduction 12

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We are interested in the approximation of 2D elliptic equations with dominated advection and featuring boundary layers. In order to reduce the computational complexity, the domain is split into two subregions, the first one far from the layer, where the can neglect the viscosity effects, and the second one next to the layer. In the latter domain the original elliptic equation is solved, while in the former one, the pure convection equation obtained by the original one by dropping the diffusive term the interface coupling is enforced by the non-conforming mortar method. We consider two different sets of interface conditions and we compare them to the what concerns both computational efficiency and stability. One of the two sets of interface conditions turns out to be very effective, especially for very small viscosity when the mortar formulation of the original elliptic problem on the global domain can fail.

# 2 The Heterogeneous Problem

We consider an open bounded domain  $\Omega\subset\mathbb{R}^2$  with Lipschitz boundary  $\partial\Omega$ , split 26 into two open subsets  $\Omega_1$  and  $\Omega_2$  such that  $\overline{\Omega}=\overline{\Omega}_1\cup\overline{\Omega}_2,\Omega_1\cap\Omega_2=\emptyset$ . Then, we 27 denote by  $\Gamma=\partial\Omega_1\cap\partial\Omega_2$ , the interface between the sub domains and we assume 28 that  $\Gamma$  is of class  $C^{1,1}$ . Given  $f\in L^2(\Omega)$ ,  $b_0\in L^\infty(\Omega)$ ,  $v\in L^\infty(\Omega_2\cup\Gamma)$  and  $\mathbf{b}\in\mathbb{C}^2$   $[W^{1,\infty}(\Omega)]^2$  satisfying the following inequalities:

$$\exists v_0 \in \mathbb{R} \text{ such that } v(\mathbf{x}) \ge v_0 > 0, \, \forall \mathbf{x} \in \Omega_2 \cup \Gamma, \\ \exists \sigma_0 \in \mathbb{R} \text{ such that } b_0(\mathbf{x}) + \frac{1}{2} \mathrm{div} \mathbf{b}(\mathbf{x}) \ge \sigma_0 > 0, \, \forall \mathbf{x} \in \Omega,$$
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we look for two functions  $u_1$  and  $u_2$  (defined in  $\overline{\Omega}_1$  and  $\overline{\Omega}_2$ , respectively) solutions 33 of the *heterogeneous problem* 34

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$$\begin{cases}
\operatorname{div}(\mathbf{b}u_{1}) + b_{0}u_{1} = f & \text{in } \Omega_{1}, \\
\operatorname{div}(-\nu\nabla u_{2} + \mathbf{b}u_{2}) + b_{0}u_{2} = f & \text{in } \Omega_{2}, \\
u_{1} = 0 & \text{on } (\partial\Omega_{1} \setminus \Gamma)^{in} \\
u_{2} = 0 & \text{on } \partial\Omega_{2} \setminus \Gamma
\end{cases} \tag{1}$$

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and satisfying the interface conditions

$$u_1 = u_2 \text{ on } \Gamma^{in}, \qquad \mathbf{b} \cdot \mathbf{n}_{\Gamma} u_1 + v \frac{\partial u_2}{\partial \mathbf{n}_{\Gamma}} - \mathbf{b} \cdot \mathbf{n}_{\Gamma} u_2 = 0, \text{ on } \Gamma.$$
 (2)

 $\mathbf{n}_{\Gamma}$  denotes the normal unit vector to  $\Gamma$  oriented from  $\Omega_1$  to  $\Omega_2$ , while for any nonempty subset  $S \subseteq \partial \Omega_1$ ,  $S^{in} = \{ \mathbf{x} \in S : \mathbf{b}(\mathbf{x}) \cdot \mathbf{n}_1(\mathbf{x}) < 0 \}$  and  $S^{out} = \{ \mathbf{x} \in S : \mathbf{b}(\mathbf{x}) \cdot \mathbf{n}_1(\mathbf{x}) < 0 \}$  $\mathbf{n}_1(\mathbf{x}) \geq 0$  are the *inflow* and the *outflow* parts of S, respectively.

Equation (2) (named IC1) express the continuity of the velocity field across the 39 inflow part of the interface and the continuity of the fluxes across the whole interface. 40 They can be equivalently expressed as (named IC2): 41

$$u_1 = u_2, \ v \frac{\partial u_2}{\partial \mathbf{n}_{\Gamma}} = 0 \ \text{on } \Gamma^{in}, \quad -\mathbf{b} \cdot \mathbf{n}_{\Gamma} u_1 = v \frac{\partial u_2}{\partial \mathbf{n}_{\Gamma}} - \mathbf{b} \cdot \mathbf{n}_{\Gamma} u_2 \ \text{on } \Gamma^{out}.$$
 (3)

Problem (1) with either interface conditions (2) or (3) is well-posed, see [5].

The heterogeneous problem (1), with either interface conditions IC1 or IC2, can 43 formally be written as an interface problem by means of Steklov-Poincaré opera- 44 tors (see, e.g., [3, 5]). Let us define the trace spaces  $\Lambda_1 = L^2_{\mathbf{h}}(\Gamma^{in}) = \{v : \Gamma^{in} \to \mathbb{R}^n\}$  $\mathbb{R}: \sqrt{|\mathbf{b} \cdot \mathbf{n}_{\Gamma}|} v \in L^{2}(\Gamma^{in})$  and  $\Lambda_{2} = H_{00}^{1/2}(\Gamma^{in}) = \{v : L^{2}(\Gamma^{in}) : \exists \tilde{v} \in H^{1/2}(\partial \Omega_{2}) : 46$  $\tilde{v}|_{\Gamma^{in}} = v, \ \tilde{v}|_{\partial\Omega_2\setminus\Gamma^{in}} = 0$ . 47 Solving (1) and (2) is equivalent to seeking  $\lambda_k \in \Lambda_k$  for k = 1, 2, such that

$$\begin{cases} \mathscr{S}_1 \lambda_1 + \mathscr{S}_2 \lambda_2 = \chi_1 + \chi_2 & \text{in } \Lambda_2', \\ \lambda_1 = \lambda_2|_{\Gamma^{in}} & \text{in } \Lambda_2, \end{cases}$$
(4)

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$$\mathscr{S}_1 \lambda_1 = -\mathbf{b} \cdot \mathbf{n}_1 u_1^{\lambda_1}, \quad \mathscr{S}_2 \lambda_2 = v \frac{\partial u_2^{\lambda_2}}{\partial \mathbf{n}_2} - \mathbf{b} \cdot \mathbf{n}_2 u_2^{\lambda_2}, \quad \text{ on } \Gamma,$$
 (5)

are the local Steklov-Poincaré operators, while  $u_1^{\lambda_1}$  and  $u_2^{\lambda}$  are the solution of 50

$$\begin{cases} \operatorname{div}(\mathbf{b}u_{1}^{\lambda_{1}}) + b_{0}u_{1}^{\lambda_{1}} = 0 & \text{in } \Omega_{1}, \\ u_{1}^{\lambda_{1}} = 0 & \text{on } (\partial \Omega_{1} \setminus \Gamma)^{in}, \quad u_{1}^{\lambda_{1}} = \lambda & \text{on } \Gamma^{in}, \end{cases}$$
(6)

and 51

$$\begin{cases} \operatorname{div}(-\nu\nabla u_2^{\lambda_2} + \mathbf{b}u_2^{\lambda_2}) + b_0 u_2^{\lambda_2} = 0 \text{ in } \Omega_2\\ u_2^{\lambda_2} = 0 \text{ on } \partial\Omega_2 \setminus \Gamma, \quad u_2^{\lambda_2} = \lambda_2 \text{ on } \Gamma, \end{cases}$$
(7)

respectively. Finally,

$$\chi_1 = \mathbf{b} \cdot \mathbf{n}_1 u_1^f, \quad \chi_2 = -v \frac{\partial u_2^f}{\partial \mathbf{n}_2} + \mathbf{b} \cdot \mathbf{n}_2 u_2^f = -v \frac{\partial u_2^f}{\partial \mathbf{n}_2}, \tag{8}$$

where  $u_1^f$  and  $u_2^f$  are the solutions of problems like (6) and (7), respectively, with null 53 trace on the interface and external load f. Note that  $\chi_1|_{\Gamma^{in}} = 0$ .

If interface conditions IC2 are considered instead of IC1, the resulting Steklov-55 Poincaré equation reads: seek  $\lambda_k \in \Lambda_k$ , for k = 1, 2 such that

$$\begin{cases} \mathscr{S}_1^0 \lambda_1 + \mathscr{S}_2^0 \lambda_2 = \chi_1 + \chi_2 & \text{in } \Lambda_2' \\ \lambda_1 = \lambda_2|_{\Gamma^{in}} & \text{in } \Lambda_2 \end{cases}$$
(9)

where 57

$$\mathcal{S}_{1}^{0}\lambda_{1} = \begin{cases} 0 & \text{on } \Gamma^{in} \\ -\mathbf{b} \cdot \mathbf{n}_{1}u_{1}^{\lambda_{1}} & \text{on } \Gamma^{out}, \end{cases} \mathcal{S}_{2}^{0}\lambda_{2} = \begin{cases} v \frac{\partial u_{2}^{\lambda_{2}}}{\partial \mathbf{n}_{2}} & \text{on } \Gamma^{in} \\ v \frac{\partial u_{2}^{\lambda_{2}}}{\partial \mathbf{n}_{2}} - \mathbf{b} \cdot \mathbf{n}_{2}u_{2}^{\lambda_{2}} & \text{on } \Gamma^{out}. \end{cases}$$
(10)

Remark 1. It is straightforward to prove that the operator  $\mathscr{S}_2^0$  is always coercive on 59  $\Lambda_2$ , whereas  $\mathscr{S}_2$  is coercive only if smallness assumption on  $\tilde{\mathbf{b}}$  is assumed. If, e.g.,

$$\|\mathbf{b}\|_{L^{\infty}(\Gamma)} \le \varepsilon_0$$
, with  $0 \le \varepsilon_0 \le 2\min\{\nu_0, \sigma_0\}/C_*^2$ , (11)

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(where  $C_*$  is the constant of the trace inequality  $\|v\|_{L^2(\partial\Omega_2)} \leq C_*\|v\|_{H^1(\Omega_2)}$ ) is sat-61 is field then  $\mathcal{S}_2$  is coercive on  $\Lambda_2$ . For this reason, the solution of problem (4) may 62 produce oscillations around  $\Gamma^{in}$  when advection dominates (i.e. the global Péclet 63 number is large), as will be shown later in our numerical results.

# 3 Mortar Coupling for Spectral Element Discretization

The discretization of the differential equation within each sub domain is performed 66 by the quadrilateral conforming Spectral Element Method (SEM). We refer to [4] 67 for a detailed description of this method. For k = 1, 2, let  $\mathcal{T}_k = \{T_{k,m}\}_{m=1}^{M_k}$  be a partition of the computational domain  $\Omega_k \subset \mathbb{R}^2$ . The SEM finite dimensional space on 69  $\overline{\Omega}_k$  is denoted by  $X_{k,\delta_k}$  and it is the set of functions in  $C^0(\overline{\Omega}_k)$  whose restriction to 70  $T_{k,m}$  is a polynomial of degree  $N_k$  in each direction.  $\delta_k$  is an abridged notation for 71 "discrete", that accounts for the local geometric sizes  $h_{k,m}$  of  $T_{k,m}$  and the local polynomial degrees  $N_k$  along each direction. Both geometric and polynomial conformity 73 is guaranteed inside  $\Omega_k$ .

The finite dimensional spaces in which we look for the SEM solution of either (4) 75 or (9) are:  $\Lambda_{1,\delta_1} \subset \Lambda_1$  and  $\Lambda_{2,\delta_2} \subset \Lambda_2$ . Their elements are globally continuous func- <sup>76</sup> tions on  $\Gamma^{in}$  and  $\Gamma$ , respectively, and local polynomials of degree  $N_k$  on each edge 77 induced by the partition  $\mathcal{T}_k$ .

For k=1,2, we denote by  $\mathcal{N}_{k,\Gamma}$  the set of nodes of  $\mathcal{T}_k \cap \Gamma$  whose cardinality is 79  $N_{k,\Gamma}$ . Similar notations are used for the nodes lying on either  $\Gamma^{in}$  or  $\Gamma^{out}$ . 80

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The finite dimensional basis  $\{\mu_1^{(i)}\}_{i=1}^{N_{1,\Gamma^{in}}}$  of  $\Lambda_{1,\delta_1}$  ( $\{\mu_2^{(i)}\}_{i=1}^{N_{2,\Gamma}}$  of  $\Lambda_{2,\delta_2}$ , resp.) is 81 composed by the characteristic Lagrange polynomials in  $\Omega_1$  ( $\Omega_2$ , resp.) associated 82 to the Legendre-Gauss-Lobatto (LGL) nodes of  $\mathcal{N}_{1,\Gamma^{in}}$  ( $\mathcal{N}_{2,\Gamma}$ , resp.). Then we set 83  $(S_{2,\delta_2})_{ij}=\int_{\Gamma}\mathscr{S}_2\mu_2^{(j)}\mu_2^{(i)}d\Gamma$  for  $i,j=0,\ldots,N_{2,\Gamma}$ , and analogous notations are used 84 to define matrices  $S^0_{2,\delta_2},S_{1,\delta_1}$  and  $S^0_{1,\delta_1}$ . Because of the high cost to compute integrals 85 exactly, all integrals are approximated by Legendre-Gauss-Lobatto (LGL) quadrature 86 rules.

We consider non-conforming couplings, i.e. we suppose that either the two partitions  $\mathcal{I}_1$  and  $\mathcal{I}_2$  do not share the same edges on  $\Gamma$  and/or the polynomial degrees 89 do not coincide in the hyperbolic domain  $\Omega_1$  and in the elliptic one  $\Omega_2$ . We adopt 90 mortar methods (see, e.g., [2]) to glue non-conforming discretization across  $\Gamma$ .

The endpoints of the edges of  $\mathscr{T}_1 \cap \Gamma^{in}$  are denoted by  $\nu_1^{(i)}$ , for  $i=1,\ldots,N_{1,\nu}$ . 92  $\tilde{\Lambda}_{1,\delta_1}$  is a suitable finite dimensional space of functions living on  $\Gamma^{in}$  and its basis 93 functions  $\psi_l$  are characterized by being  $L^2$  functions on  $\Gamma^{in}$  and local polynomials 94 of degree  $N_1-2$  on each edge of  $\mathcal{T}_1\cap\Gamma^{in}$ . Therefore, the dimension of  $\tilde{\Lambda}_{1,\delta_1}$  is 95  $N_{\tilde{\Lambda}_1} = N_{1,\Gamma^{in}} - N_{1,\nu}$ . By choosing  $\Omega_2$  as the master domain and  $\Omega_1$  as the slave, the continuity constraint  $\lambda_1 = \lambda_2|_{\Gamma^m}$  is imposed weakly, i.e. by requiring that

$$\int_{\Gamma^{in}} (\lambda_{1,\delta_1} - \lambda_{2,\delta_2}) \psi_l d\Gamma = 0 \qquad \forall \psi_l \in \tilde{\Lambda}_{1,\delta_1}, \tag{12}$$

jointly with the strong continuity at the nodes  $v_1^{(i)}$  of  $\mathcal{T}_1 \cap \Gamma^{in}$ , for  $i = 1, \dots, N_{1,v}$ . This 98 leads us to define a new set of *mortar* functions in  $\Lambda_{1,\delta_1}$ , which are denoted by  $\tilde{\mu}_1^{(k)}$  99 (for  $k = 1, ..., N_{2,\Gamma^{in}}$ ) and satisfy the constraints:

$$\begin{cases} \tilde{\mu}_{1}^{(k)}(v_{1}^{(i)}) = \mu_{2}^{(k)}(v_{1}^{(i)}), & i = 1, \dots, N_{1,\nu} \text{ and } v_{1}^{(i)} \text{ being endpoint} \\ & \text{of at least one edge of } \mathscr{T}_{1} \cap \Gamma^{in} \\ \int_{\Gamma^{in}} (\tilde{\mu}_{1}^{(k)} - \mu_{2}^{(k)}) \psi_{l} d\Gamma = 0, \ l = 1, \dots, N_{\tilde{\Lambda}_{1}} \text{ and for all } \psi_{l} \in \tilde{\Lambda}_{1,\delta_{1}}. \end{cases}$$
(13)

Remark 2. We choose  $\Omega_2$  as the master domain because the nature of the heterogeneous problem requires to work with the trace of the elliptic solution on the whole 102 interface and with the trace of the hyperbolic one only on  $\Gamma^{in}$ . Therefore it is more 103 convenient to have the master trace at disposal on the whole  $\Gamma$ , instead of on a part 104 of it.

The matrix form of system (13) reads

$$P\Xi = \Phi,\tag{14}$$

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where  $\Xi = [\xi_{jk}] \in \mathbb{R}^{N_{1,\Gamma in} \times N_{2,\Gamma in}}$  is defined by the relations 107

$$\tilde{\mu}_{1}^{(k)} = \sum_{j=1}^{N_{1,\Gamma^{in}}} \xi_{jk} \mu_{1}^{(j)}, \qquad k = 1, \dots, N_{2,\Gamma^{in}},$$
(15)

while  $P \in \mathbb{R}^{N_{1,\Gamma^{in}} \times N_{1,\Gamma^{in}}}$  and  $\Phi \in \mathbb{R}^{N_{1,\Gamma^{in}} \times N_{2,\Gamma^{in}}}$ , are defined starting from (13). The matrix P is non-singular in view of the inf-sup condition for  $\mathbb{Q}_N - \mathbb{Q}_{N-2}$  [2]. Once the discretization in  $\Omega_1$  and  $\Omega_2$  has been chosen, the matrix  $\Xi$  can be explicitly computed by solving (14).

The matrix  $\Xi$  enforces the gluing between degrees of freedom defined on  $\mathcal{N}_{2,\Gamma^{in}}$  112 and  $\mathcal{N}_{1,\Gamma^{in}}$ . Therefore, Steklov-Poincaré equations (4) and (9) can be written in a 113 nonconforming setting, by the use of matrix  $\Xi$ .

On  $\Gamma^{out}$  no continuity constraint, neither strong nor weak, is imposed, since the continuity of fluxes is a natural consequence of the interface equation. Nevertheless, 116 on  $\Gamma^{out}$  we have to compute integrals of basis functions associated to two different 117 meshes. To this aim we introduce the matrix  $Q \in \mathbb{R}^{N_2,\Gamma^{out} \times N_1,\Gamma^{out}}$  for the evaluations 118 of functions of  $\Lambda_{1,\delta_1}$  at the nodes of  $\mathscr{T}_2 \cap \Gamma$ , and the matrix  $D = M_{2,\delta_2}^{out} Q(M_{1,\delta_1}^{out})^{-1}$ , 119 where  $M_{k,\delta_k}^{out}$  are the mass matrices induced by the LGL quadrature formulas on  $\Gamma^{out}$ , 120 for k=1,2.

The nonconforming finite dimensional counterpart of (4) reads: find  $\lambda_{k,\delta_k} \in \Lambda_{k,\delta_k}$  122 for k=1,2, such that

$$\left\{ \underbrace{\begin{pmatrix} S_{2,\delta_{2}} + \begin{bmatrix} \Xi^{T} S_{1,\delta_{1}}^{in} \Xi & 0 \\ DS_{1,\delta_{1}}^{out} \Xi & 0 \end{bmatrix} \end{pmatrix}}_{S_{\delta}} \begin{bmatrix} \lambda_{2,\delta_{2}}^{in} \\ \lambda_{2,\delta_{2}}^{out} \end{bmatrix} = \begin{bmatrix} M_{2,\delta_{2}}^{in} \chi_{2,\delta_{2}}^{in} \\ M_{2,\delta_{2}}^{out} \chi_{2,\delta_{2}}^{out} + D\chi_{1,\delta_{1}}^{out} \end{bmatrix} \\
\lambda_{1,\delta} = \Xi \lambda_{2,\delta_{2}}^{in} \tag{16}$$

whereas that of (9) becomes: find  $\lambda_{k,\delta_k} \in \Lambda_{k,\delta_k}$  for k = 1, 2, such that

$$\left\{ \underbrace{\begin{pmatrix} S_{2,\delta_{2}}^{0} + \begin{bmatrix} 0 & 0 \\ DS_{1,\delta_{1}}^{out} \Xi & 0 \end{bmatrix} \end{pmatrix}}_{S_{\delta}^{0}} \begin{bmatrix} \lambda_{2,\delta_{2}}^{in} \\ \lambda_{2,\delta_{2}}^{out} \\ \lambda_{2,\delta_{2}}^{out} \end{bmatrix} = \begin{bmatrix} M_{2,\delta_{2}}^{in} \chi_{2,\delta_{2}}^{in} \\ M_{2,\delta_{2}}^{out} \chi_{2,\delta_{2}}^{out} + D\chi_{1,\delta_{1}}^{out} \end{bmatrix} \\
\lambda_{1,\delta} = \Xi \lambda_{2,\delta_{2}}^{in}.$$
(17)

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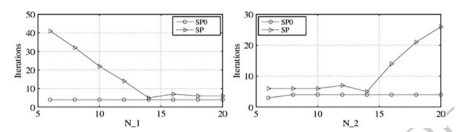
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The upper scripts in and out denote the restriction to  $\Gamma^{in}$  and  $\Gamma^{out}$ , resp.

The numerical solutions of these linear systems is carried out by preconditioned 126 Bi-CGStab iterations (see, [6]). 127

When conforming discretization is used across the interface (i.e.  $\delta_1 = \delta_2$ ), matrix 128  $\Xi$  reduces to the identity matrix. In this situation, it is well known (see, e.g. [5]) 129 that  $S_{2,\delta_2}^0$  is an optimal preconditioner for the matrix  $S_{\delta}^0$ , i.e.  $\exists C_0 > 0$  independent of 130  $\delta$  such that its spectral condition number  $\mathscr{K}((S_{2,\delta_2}^0)^{-1}S_{\delta}^0)$  is bounded by  $C_0$ . When 131  $\delta_1 = \delta_2, S_{2,\delta_2}^0$  is an optimal preconditioner also for  $S_{\delta}$  (see [3]), i.e. there exists  $C_1 > 0$  132 independent of  $\delta$  such that  $\mathscr{K}((S_{2,\delta_2}^0)^{-1}S_{\delta}) \leq C_1$ , and numerical results show that 133  $C_0 \leq C_1$ .

We extend here the use of the preconditioner  $S_{2\delta}^0$  to the non-conforming case.



**Fig. 1.** Preconditioned Bi-CGStab iterations. The viscosity is  $v = 10^{-2}$ . At left,  $N_2 = 10^{-2}$ . fixed, at right,  $N_1 = 14$  is fixed.  $4 \times 4$  equal spectral elements are taken in each  $\Omega_k$ 

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#### 4 Numerical Results

Test case: the computational domain  $\Omega = (-1,1)^2$  is split in  $\Omega_1 = (-1,0.8) \times 137$ (-1,1) and  $\Omega_2 = (0.8,1) \times (-1,1)$ . The interface is  $\Gamma = \{0.8\} \times (-1,1)$ . The data of the problem are:  $\mathbf{b} = [5y, 1-x]^t$ ,  $b_0 = 1$ , f = 1 and the inflow interface 139 is  $\Gamma^{in} = \{0.8\} \times (-1,0)$ . The imposed Dirichlet boundary conditions are:  $u_1 = 1$  140 on  $((-1,0.8)\times\{-1\})\cup(\{-1\}\times(0,1))$ ,  $u_2=0$  on  $\{1\}\times(-1,1)$ ,  $u_2=1$  on 141  $(0.8,1) \times \{-1\}$ , while the homogeneous Neumann condition  $\frac{\partial u_2}{\partial \mathbf{n}_2} = 0$  is imposed 142 on  $(0.8, 1) \times \{1\}$ .

Because of the presence of a boundary layer near the right vertical side, the mesh 144 is refined there (without losing the conformity inside  $\Omega_2$ ) to prevent the numerical 145 solution to be affected by spurious oscillations.

In Fig. 1 the number of Preconditioned Bi-CGStab (PBi-CGStab) iterations (with 147 preconditioner  $S_{2,\delta_2}$ ) required to reduce the relative norm of the residual of 12 orders 148 of magnitude is plotted versus the polynomial degrees  $N_1$  and  $N_2$  of the mortar discretization. These results refer to  $v = 10^{-2}$  and show that the Steklov-Poincaré formulation (9) performs better than (4). The analysis of this and other test cases leads 151 us to conjecture that  $\mathcal{K}((S^0_{2,\delta_0})^{-1}S^0_{\delta}) \leq C_0$  still holds for non-conforming coupling 152  $(\delta_1 \neq \delta_2)$ , while

$$\mathcal{K}((S_{2,\delta_2}^0)^{-1}S_{\delta}) \simeq C_1 \mathcal{K}(\Xi\Xi^T) \simeq C_1 \begin{cases} (N_2 - N_1 + 1)^{3/2} & \text{if } N_1 < N_2 \\ C_2 & \text{if } N_1 \ge N_2, \end{cases}$$
(18)

where  $C_1$  is the constant defined in the previous section, and  $C_2$  is another positive 154 constant independent of  $\delta$ .

Therefore, formulation (17) corresponding to IC2 is optimally preconditioned by 156  $S_{2,\delta_0}^0$  and it is better than (16) (corresponding to IC1) for what concerns the computational efficiency.

Moreover, when the viscosity vanishes (see Table 1), the performance of the SPO 159 approach (17) does not downgrade, as the number of PBi-CGStab iterations keeps 160 bounded: three or four iterations are enough to satisfy the stopping test independently 161 of both viscosity and discretization parameters.

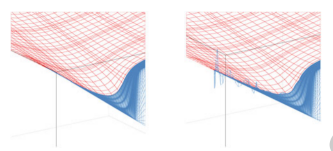


Fig. 2. Zoom on the numerical solution for  $v = 10^{-3}$  and: (9) (left), (4) (right) with  $N_1 = 8$ and  $N_2 = 24$ . The elliptic solution  $u_2$  is in front, while the hyperbolic one  $u_1$  is behind

On the contrary, the number of PBi-CGStab iterations required by SP approach 163 (16) noticeably grows up when  $v \to 0$  and behaves like  $(N_2 - N_1 + 1)^{3/4}$  when  $N_1 < 164$  $N_2$ , in agreement with (18).

The large number of PBi-CGStab iterations required by SP is due to the presence 166 of instabilities across  $\Gamma^{in}$  which develop when advection dominates and the larger 167  $N_2 - N_1$  is, the more they are pronounced.

We verified that the same instability occurs when mortar methods are applied 169 to solve the pure elliptic-elliptic couplings with dominated advection and interface 170 condition  $v \frac{\partial u_1}{\partial \mathbf{n}_{\Gamma}} - \mathbf{b} \cdot \mathbf{n}_{\Gamma} u_1 = v \frac{\partial u_2}{\partial \mathbf{n}_{\Gamma}} - \mathbf{b} \cdot \mathbf{n}_{\Gamma} u_2$  on the whole interface  $\Gamma$ . Indeed, the 171 local Steklov-Poincaré operators associated to the latter interface condition behaves 172 like operator  $\mathscr{S}_2$  introduced in (5), and they can lose the coercivity when  $\|\mathbf{b}\|_{L^{\infty}(\Omega)}$ is large. This is the subject of a work in progress. (See also [1].)

In conclusion, the heterogeneous approach (1) with interface conditions IC2 and 175 non-conforming mortar coupling turns out to be the most efficient and accurate one 176 for vanishing viscosity and it is also a valid way to overcome instabilities arising 177 from the mortar discretization of elliptic equations with dominated advection.

In Fig. 2 the heterogeneous solutions obtained by solving both (17) and (16) with 179  $v = 10^{-4}$ ,  $N_1 = 8$  and  $N_2 = 24$  are shown. The elliptic solution  $u_2$  provided by (16) 180 (Fig. 2, right) exhibits non-trivial oscillations, while that provided by (17) (Fig. 2, 181 left) does not.

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**Table 1.** PBi-CGStab iterations to solve systems SP0 (17) and SP (16) with  $P = S_{2,\delta_0}^0$  versus the viscosity. At left,  $N_1 = 8$ , at right,  $N_1 = 20$ ,  $N_2 = 24$ .  $4 \times 4$  equal spectral elements are taken in each  $\Omega_k$ .  $N_2 = 64$  along x-direction in the elements next to the layer

			$10^{-3}$				$10^{-2}$		
SP0	3	4	3	3	SP0	3	3	3	4
			262		SP	7	17	35	86

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# Heterogeneous Substructuring Methods for Coupled Surface and Subsurface Flow

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1 Introduction 9

The exchange of ground- and surface water plays a crucial role in a variety of practically relevant processes ranging from flood protection measures to preservation of ecosystem health in natural and human-impacted water resources systems.

Commonly accepted models are based on the shallow water equations for overland flow and the Richards equation for saturated—unsaturated subsurface flow with suitable coupling conditions. Continuity of mass flow across the interface is natural, because it directly follows from mass conservation. Continuity of pressure is typically imposed for simplicity. Mathematically, this makes sense for sufficiently smooth height of surface water as occurring, e.g., in filtration processes [9, 14]. Here we impose Robin-type coupling conditions modelling a thin, nearly impermeable layer at the bottom of the river bed that may cause pressure discontinuities; an effect which is known in hydrology as clogging (see [16] or [8, p. 1376]). From a mathematical perspective, clogging can be regarded as a kind of regularization, because, 22 in contrast to Dirichlet conditions, Robin conditions can be straightforwardly formulated in a weak sense.

Existence and uniqueness results for the Richards equation and the shallow water equations are rare and hard to obtain, and nothing seems to be known about 26 solvability of coupled problems. Extending the general framework of heterogeneous Steklov-Poincaré formulations and iterative substructuring [10, 13] to timedependent problems, we introduce a Robin-Neumann iteration for the continuous 29 coupled problem and motivate its feasibility by well-known existence results for the 30 linear case. As surface and subsurface flow are only weakly coupled by clogging and 31 continuity of mass flux, different discretizations with different time steps and different meshes can be used in a natural way. This is absolutely necessary, to resolve the 32 vastly different time and length scales of surface and subsurface flow. Discrete mass 34 conservation can be proved in a straightforward way.

Finally, we illustrate our considerations by coupling a finite element discretiza- 36 tion of the Richards equation based on Kirchhoff transformation [4] with a simple 37

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upwind discretization of surface flow. Numerical experiments confirm discrete mass 38 conservation and show fast convergence of the Robin-Neumann iteration for real-life 39 soil data.

## 2 Coupled Surface and Subsurface Flow

Saturated–unsaturated subsurface flow during a time interval  $(0, T_{end})$  in a porous 42 medium occupying a bounded domain  $\Omega \subset \mathbb{R}^d$ , d = 2, 3, is described by the Richards 43 equation

$$n \theta(p)_t + \operatorname{div} \mathbf{v}(p) = 0$$
,  $\mathbf{v}(p) = -\frac{K}{\mu} kr(\theta(p)) \nabla(p - \rho gz)$ . (1)

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The porosity n, permeability K, viscosity  $\mu$ , and density  $\rho$  are given parameters, and 45 g is the earth's gravitational acceleration. The unknown capillary pressure p is related 46 to saturation  $\theta(p)$  and relative permeability  $kr(\theta(p))$  by equations of state [6, 7]

$$\theta(p) = \begin{cases} \theta_m + (\theta_M - \theta_m) \left(\frac{p}{p_b}\right)^{-\lambda} & \text{for } p \leq p_b \\ \theta_M & \text{for } p \geq p_b \end{cases}$$
$$kr(\theta) = \left(\frac{\theta - \theta_m}{\theta_M - \theta_m}\right)^{3 + \frac{2}{\lambda}}, \qquad \theta \in [\theta_m, \theta_M] \subset [0, 1],$$

with residual saturation  $\theta_m$ , maximal saturation  $\theta_M$ , bubbling pressure  $p_b < 0$ , and 48 pore size distribution factor  $\lambda > 0$ . Let  $\Gamma \subset \partial \Omega$  denote the coupling boundary of the porous medium with a surface flow, and denote the outward normal vector of  $\Gamma$  by **n**. 50 We impose the coupling by Robin conditions  $p|_{\Gamma} - \alpha \mathbf{v} \cdot \mathbf{n} \in L^2((0, T_{\mathrm{end}}), H^{-1/2}(\Gamma))$  51 on  $\Gamma$  and homogeneous Neumann conditions on  $\partial\Omega\setminus\Gamma$ . With compatible initial 52 conditions  $\theta_0 \in L^1(\Omega)$  we assume that (1) admits a unique weak solution  $p \in 53$  $L^2((0,T_{\rm end}),H^1(\Omega))$ . This assumption is motivated by known existence results [1] 54 for the Kirchhoff transformed Richards equation (see also [4]) and is, obviously, sat- 55 is field in the case of saturated flow  $\theta \equiv \theta_M$ .

The surface flow on  $\Gamma$  is described by the shallow water equations

$$h_t + \operatorname{div} \mathbf{q} = r, \tag{2a}$$

$$\mathbf{q}_t + \operatorname{div} \mathbf{F}(h, \mathbf{q}) = -gh\nabla\phi$$
 (2b)

where  $\phi: \Gamma_0 \to \Gamma$  is a parametrization of the surface topography of  $\Gamma$ . The unknown 58 water height h and discharge  $\mathbf{q}$ , as well as a given mass source r are functions on 59  $(0, T_{\rm end}) \times \Gamma_0$ . For ease of presentation, we assume  $\Gamma = \Gamma_0$  so that  $\Gamma$  is an open subset of  $\mathbb{R}^{d-1}$ . For d=3, i.e.,  $\Gamma\subset\mathbb{R}^2$ , the flux function **F** takes the form 61

$$\mathbf{F} = \begin{pmatrix} \mathbf{F}_1 \\ \mathbf{F}_2 \end{pmatrix}, \qquad \mathbf{F}_1(h, \mathbf{q}) = \begin{pmatrix} q_1^2/h + \frac{1}{2}gh^2 \\ q_1q_2/h \end{pmatrix}, \quad \mathbf{F}_2(h, \mathbf{q}) = \begin{pmatrix} q_1q_2/h \\ q_2^2/h + \frac{1}{2}gh^2 \end{pmatrix}$$
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with  $\mathbf{q}=(q_1,q_2)$ . It degenerates to  $\mathbf{F}(h,\mathbf{q})=\mathbf{q}^2/h+\frac{1}{2}gh^2$  for  $\Gamma\subset\mathbb{R}$ . For suitable 63 initial conditions and inflow conditions on  $\partial\Gamma_{\mathrm{in}}\subset\partial\Gamma$  we assume that (2) has a weak 64 solution  $(h,\mathbf{q})\in L^\infty((0,T_{\mathrm{end}}),L^\infty(\Gamma))^d$  in the sense of distributions  $\mathscr{D}'((0,T_{\mathrm{end}})\times$  65  $\Gamma_{\mathrm{in}})$  where  $\Gamma_{\mathrm{in}}=\Gamma\cup\partial\Gamma_{\mathrm{in}}$ . Since regularity results for nonlinear hyperbolic systems 66 (2) do not seem to be available we note that this assumption is satisfied in the linear 67 case [15, Theorem 2.2].

Mass conservation provides the Neumann coupling condition

$$r = \mathbf{v} \cdot \mathbf{n}$$
.

Following, e.g. [16], we postulate a nearly impermeable river bed with thickness  $\varepsilon \ll 70$  1 and permeability  $K_{\varepsilon}$  (clogging). Then Darcy's law provides the flux  $\mathbf{v} = -\frac{K_{\varepsilon}}{\mu} \nabla p_{\varepsilon}$ . 71 Setting  $\nabla p_{\varepsilon} = \varepsilon^{-1} (\rho g h - p|_{\Gamma}) \mathbf{n}$ , we obtain the Robin coupling condition

$$p|_{\Gamma} - \alpha \mathbf{v} \cdot \mathbf{n} = \rho g h \tag{3}$$

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with leakage coefficient  $\alpha = \frac{\mu \varepsilon}{K_{\varepsilon}}$ . Note that (3) generally implies a pressure discontinuity across the interface  $\Gamma$  between ground and surface water.

Remark 1. In light of the above regularity assumptions on pressure p and surface 75 water height h coupling surface and subsurface flow by continuity  $p|_{\Gamma} = \rho gh$  of cap-76 illary and hydrostatic pressure is generally not possible, because there is a regularity 77 gap between the trace  $p|_{\Gamma} \in L^2((0,T_{\mathrm{end}}),H^{1/2}(\Gamma))$  and  $h \in L^{\infty}((0,T_{\mathrm{end}}),L^{\infty}(\Gamma)) \not\subset$  78  $L^2((0,T_{\mathrm{end}}),H^{1/2}(\Gamma))$  (see, e.g., [5, p. 148]) However, sufficient smoothness is available in special cases like, e.g., in- and exfiltration processes [14].

# 3 Steklov-Poincaré Formulation and Substructuring

We introduce the Robin-to-Neumann map

$$S_{\Omega}(h) = \mathbf{v}(h) \cdot \mathbf{n} = \alpha^{-1}(p|_{\Gamma} - \rho gh)$$
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for  $h \in L^{\infty}((0,T_{\mathrm{end}}),L^{\infty}(\Gamma)) \subset L^{2}((0,T_{\mathrm{end}}),H^{-1/2}(\Gamma))$ . Here, p is the solution of 84 the Richards equation (1) with Robin conditions (3). Assuming that for given 85  $h \in L^{\infty}((0,T_{\mathrm{end}}),L^{\infty}(\Gamma))$  and corresponding inflow boundary conditions, the second part (2b) of the shallow water equations has a unique weak solution  $\mathbf{q}(h) \in \mathbb{R}$   $L^{\infty}((0,T_{\mathrm{end}}),L^{\infty}(\Gamma))^{d-1}$ , we set

$$S_{\Gamma}(h) = -\operatorname{div} \mathbf{q}(h)$$
 .

The Steklov–Poincaré formulation of the coupled Richards equation and shallow 90 water equations then reads 91

$$h_t = S_{\Omega}(h) + S_{\Gamma}(h) . \tag{4}$$

Just as (2a) and (4) is understood in the sense of distributions  $\mathscr{D}'((0,T_{\mathrm{end}})\times\Gamma_{\mathrm{in}})$ .

In complete analogy to the stationary case [10, 13] we introduce a damped 93 Robin-Neumann iteration

$$h_t^{\nu+1/2} - S_{\Gamma}(h^{\nu+1/2}) = S_{\Omega}(h^{\nu}), \quad h^{\nu+1} = h^{\nu} + \omega(h^{\nu+1/2} - h^{\nu}),$$
 (5)

with a suitable damping parameter  $\omega \in (0, \infty)$  and with an initial iterate given by 95  $h^0 \in L^{\infty}((0, T_{\text{end}}), L^{\infty}(\Gamma))$ . Each step amounts to the solution of the Richards equation with Robin boundary conditions (3) to evaluate the source term  $S_O(h^V)$ , and the 97 subsequent solution of the shallow water equations (2) to evaluate  $h^{v+1/2}$ . The feasi-98 bility of (5) requires existence and uniqueness of these solutions. Note the similarity 99 to waveform relaxation methods [11].

After selecting a step size  $\Delta T = T_{\rm end}/N$  with suitable  $N \in \mathbb{N}$  and corresponding time levels  $T_k = k\Delta T$ , the Robin-Neumann iteration (5) can also be applied on 102 subintervals  $[T_{k-1}, T_k], k = 1, \dots, N$ .

#### 4 Discretization and Discrete Robin-Neumann Iteration

We first derive a discrete version of the Steklov-Poincaré formulation (4) on a fixed 105 time interval  $[T_k, T_{k+1}]$  with  $0 \le T_k < T_{k+1} = T_k + \Delta T \le T_{\text{end}}$ . To this end, we introduce intermediate time levels  $t_i = T_k + i\tau$ , i = 0, ..., M, with step size  $\tau = \Delta T/M$  and suitable  $M \in \mathbb{N}$ . Spatial discretization is based on a partition  $\mathcal{T}_{\Gamma}$  of  $\Gamma$  into simplices 108 T that is regular in the sense that the intersection of two simplices  $T, T' \in \mathcal{T}_{\Gamma}$  is either a common face, edge, vertex, or empty. We introduce the corresponding space 110 of discontinuous finite elements of order q > 0 by

$$\mathscr{V}_{\Gamma} = \{v \in L^2(\Gamma) \mid v_T \text{ is a polynomial of degree at most } q \ \forall T \in \mathscr{T}_{\Gamma} \}$$
 ,

and let  $h = (h_i)_{i=0}^M$  denote approximations  $h_i \in \mathcal{V}_{\Gamma}$  at  $t_i, i = 0, ..., M$ .

Then, utilizing the forward difference quotient  $\partial_t h_i = (h_{i+1} - h_i)/\tau$ , a discrete 114 Steklov-Poincaré formulation reads 115

$$\partial_t h_i = S_{\Gamma}(h)_i + S_{\Omega}(h)_i, \qquad i = 0, \dots, M - 1.$$
 (6)

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Here and in the rest of this section, subscripts i indicate approximations taken at time 116  $t_i$ . 117

For given  $h = (h_i)_{i=0}^M$ , the discrete surface flow 118

$$(S_{\Gamma}(h)_{i}, \nu)_{\Gamma} = \sum_{T \in \mathscr{T}_{\Gamma}} \left( (\mathbf{q}(h)_{i}, \nabla \nu)_{T} + (\mathbf{G}_{h}(h_{i}, \mathbf{q}(h)_{i}) \cdot \mathbf{n}_{T}, \nu)_{\partial T} \right) \quad \forall \nu \in \mathscr{V}_{\Gamma}$$
 (7)

results from an explicit discontinuous Galerkin discretization of (2a), characterized 119 by the discrete flux function  $G_h$ . Here,  $(\cdot,\cdot)_U$  stands for the  $L^2$  scalar product on U=120 $\Gamma$ , T,  $\partial T$ , respectively;  $\mathbf{n}_T$  is the outward normal on T, and the discrete discharge 121  $\mathbf{q}_i = \mathbf{q}(h)_i$  is obtained from an explicit discontinuous Galerkin discretization of (2b) 122

$$(\partial_t \mathbf{q}_i, v)_{\Gamma} = \sum_{T \in \mathscr{T}_{\Gamma}} \left( (\mathbf{F}(h_i, \mathbf{q}_i), \nabla v)_T + (\mathbf{G}_{\mathbf{q}}(h_i, \mathbf{q}_i) \cdot \mathbf{n}_T, v)_{\partial T} \right) \quad \forall v \in (\mathscr{V}_{\Gamma})^{d-1} . \quad (8)$$

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Since we expect the dynamics of subsurface flow to be much slower than the 123 surface water dynamics, we use the macro time step  $\Delta T$  for an implicit time discretization of  $S_{\Omega}(h)$ . The spatial discretization is based on conforming piecewise 125 linear finite elements

$$\mathscr{V}_{\Omega} = \{ v \in C(\overline{\Omega}) \mid v \mid_T \text{ is affine linear } \forall T \in \mathscr{T}_{\Omega} \}$$

with respect to a regular partition  $\mathscr{T}_{\Omega}$  of  $\Omega$ . No compatibility conditions on  $\mathscr{T}_{\Omega}$  and 128  $\mathscr{T}_{\Gamma}$  are required. For given  $p_k \in \mathscr{V}_{\Omega}$  and  $h_{k+1} \in \mathscr{V}_{\Gamma}$ , the discrete capillary pressure  $p_{k+1} \in \mathcal{V}_{\Omega}$  is then obtained from the variational equality

$$n\langle \theta_{k+1}, v \rangle_{\Omega} + \Delta T \left( (\mathbf{v}_{k+1}, \nabla v)_{\Omega} + \alpha^{-1} (\langle p_{k+1}|_{\Gamma}, v \rangle_{\Gamma} - (\rho g h_{k+1}, v)_{\Gamma}) \right) = n\langle \theta_{k}, v \rangle_{\Omega} \quad \forall v \in \mathscr{V}_{\Omega}.$$

$$(9)$$

Here  $\langle \cdot, \cdot \rangle_{\Omega}$  denotes the lumped  $L^2$  scalar product on  $\Omega$ ,  $\langle \cdot, \cdot \rangle_{\Gamma}$  is the corresponding lumped  $L^2$  scalar product on  $\Gamma$ ,  $\theta_k = \theta(p_k)$ , and  $\mathbf{v}_{k+1}$  is a discretization of the flux  $\mathbf{v}_{k+1}$ at  $T_{k+1}$ . Once  $p_{k+1} \in \mathscr{V}_{\Omega}$  is available, we set for all i = 0, ..., M

$$(S_{\Omega}(h)_{i}, v)_{\Gamma} = \alpha^{-1}(p_{k+1}|_{\Gamma} - \rho g h_{k+1}, v)_{\Gamma} \qquad \forall v \in \mathcal{V}_{\Gamma}.$$

$$(10)$$

Note that  $S_{\Omega}(h)_i$  is constant on the macro interval  $[T_k, T_{k+1}]$  and only depends on  $h_{k+1}$ . 135

Testing (6) and (9) with constant functions  $\mathbf{1} \in \mathcal{V}_{\Gamma}$  and  $\mathbf{1} \in \mathcal{V}_{\Omega}$ , respectively, and using  $\langle p_{k+1}|_{\Gamma}, \mathbf{1}\rangle_{\Gamma} = (p_{k+1}|_{\Gamma}, \mathbf{1})_{\Gamma}$  we obtain discrete mass conservation.

**Proposition 1.** The discrete Steklov–Poincaré formulation (6) with  $S_{\Gamma}$  and  $S_{\Omega}$  de-138 fined by (7) and (10) is mass conserving in the sense that 139

$$(h_{k+1},\mathbf{1})_{\Gamma} + n\langle \theta_{k+1},\mathbf{1}\rangle_{\Omega} = (h_k,\mathbf{1})_{\Gamma} + n\langle \theta_k,\mathbf{1}\rangle_{\Omega} + \tau \sum_{i=0}^{M-1} (\mathbf{G}_h(h_i,\mathbf{q}_i) \cdot \mathbf{n}_{\partial\Gamma},\mathbf{1})_{\partial\Gamma}$$
 140

holds for k = 0, 1, ..., with  $\mathbf{n}_{\partial \Gamma}$  denoting the outward normal on  $\partial \Gamma$ .

We emphasize that this result holds for arbitrary discretizations of the Richards 142 flux v. 143

The discrete Steklov–Poincaré formulation (6) gives rise to the discrete damped 144 Robin-Neumann iteration 145

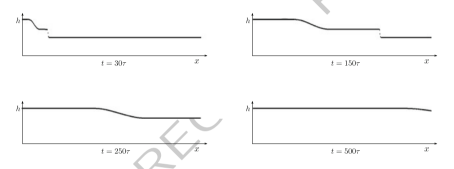
$$\partial_t h_i^{\nu+1/2} - S_{\Gamma}(h^{\nu+1/2})_i = S_{\Omega}(h^{\nu})_i , \quad h_i^{\nu+1} = h_i^{\nu} + \omega(h_i^{\nu+1/2} - h_i^{\nu}) , \quad (11)$$

with suitable damping parameter  $\omega \in (0, \infty)$ , and an initial iterate  $h_i^0 \in \mathscr{V}_{\Gamma}$  for i = 146 $0, \dots, M$ . Each step amounts to the solution of the discretized Richards equation (9) to 147 obtain  $S_{\Omega}(h^{\nu})_i$  from (10) with  $p_{k+1} = p_{k+1}^{\nu+1}$ , and to M time steps of the discontinuous Galerkin discretization of (2) described by (7) and (8) to obtain  $h_i^{v+1/2}$ ,  $i=1,\ldots,M$ . 149 For k > 0 the initial iterate  $h^0$  is the solution of the preceding time step. We emphasize that no compatibility conditions on the different meshes  $\mathscr{T}_{\Gamma}$  and  $\mathscr{T}_{\Omega}$  are necessary, 151 because only weak coupling conditions are involved. 152

## 5 Numerical Experiments

We consider a model problem on a square  $\Omega \subset \mathbb{R}^2$  of side length 10 m and select  $\Gamma$  as the upper part of its boundary. The soil parameters are n = 0.437,  $\theta_m = 0.0458$ ,  $\theta_M = 155$ 1,  $p_b = -712.2 \,\mathrm{Pa}$ ,  $\lambda = 0.694$ , and  $K = 6.66 \cdot 10^{-9} \,\mathrm{m}^2$  (sandy soil). The viscosity and density of water is  $\mu = 1$  m Pa s and  $\rho = 1,000$  kg m<sup>-3</sup>, respectively. In accordance with measurements [16] we select the leakage coefficient as  $\alpha = \rho g L^{-1}$  with L = 158 $10^{-6} \,\mathrm{s}^{-1}$  allowing for large pressure jumps across the interface.

We choose the initial conditions  $\theta_0 \equiv \theta(-20 \,\mathrm{Pa}) = 0.1401$ ,  $h(0) \equiv 1 \,\mathrm{m}$ ,  $\mathbf{q}(0) \equiv 160 \,\mathrm{m}$  $10\,\mathrm{m}^2\,\mathrm{s}^{-1}$ , and inflow boundary conditions for h(0,t) and  $\mathbf{q}(0,t)$  alternating between 161 2 and 1 m and 20 and  $10 \,\mathrm{m}^2 \,\mathrm{s}^{-1}$ , respectively, with a period of 10 s. This leads to 162 a supercritical water flow from left to right, which can result, for example, from 163 opening a flood gate.



**Fig. 1.** The water height  $h_i$  at times  $t_i = i\tau$ , i = 30, 150, 250, 500



**Fig. 2.** The pressure p at times  $T_k = k\Delta T$ , k = 200, 1,000, 2,000, 3,000

For the porous media flow on  $\Omega$  we use the uniform time step size  $\Delta T = 50$  s and a triangulation  $\mathcal{T}_O$  resulting from six uniform refinement steps applied to a partition 166 of  $\Omega$  into two triangles with hypotenuse from lower left to upper right. The Richards equation (1) is discretized by the implicit scheme based on Kirchhoff transformation 168 suggested in [4], and truncated monotone multigrid [12] is used as the algebraic 169 solver. For the surface flow we use the time step size  $\tau = \gamma \Delta T$  with  $\gamma = 3^{-1} \cdot 10^{-4}$ , 170 and the partition  $\mathcal{T}_{\Gamma}$  consists of 400 elements of equal length. Note that  $\mathcal{T}_{\Gamma}$  does not match with  $\mathcal{T}_{\Omega}|_{\Gamma}$ . The shallow water equations (2) are discretized by a discontinuous Galerkin method (7) with  $\mathcal{V}_{\Gamma}$  consisting of piecewise constant functions, and we use simple upwind flux functions  $\mathbf{G}_h$  and  $\mathbf{G}_{\mathbf{q}}$  in (7) and (8), respectively. The final time to  $T_{\mathrm{end}} = 3.5 \cdot 10^4 \, \mathrm{s}$ . For the implementation we used the DUNE libraries [2] and the domain decomposition module dune-grid-glue [3].

Figure 1 shows the evolution of the surface water height h over the first period of the boundary conditions. The porous medium flow is much slower, as expected. The Figure 2 shows the evolution of the pressure. Water enters the domain from the top, and after about 3,600 macro time steps or, equivalently, 3,000 m, the soil saturation is constant at about 75 %. Then, the domain gets fully saturated starting from the bottom. Hydrostatic pressure builds up and is fully reached at time step 4,700.

At each time step we observe discrete mass conservation up to machine precision. 183 The total relative mass loss over the entire evolution is about  $10^{-10}$ . Our numerical 184 computations thus nicely reproduce the theoretical findings of Proposition 1.

In order to investigate the convergence behavior of the Robin–Neumann iteration (11), we consider the algebraic error  $\|h_M - h_M^v\|_{L^1(\Gamma)}$  at the end of the first time interval  $[0,T_1]$  with  $T_1 = M\tau$ . It turns out that for the given leakage coefficient  $\alpha = \rho g 10^6$  s (cf. [16]), the convergence rates are in the range of  $10^{-4}$ . They remain there during the entire evolution. For each time step only two or three iterations were necessary to reduce the estimated algebraic error below the threshold  $10^{-12}$ . This is explained by the weak (in the physical sense) coupling of surface water and subsurface flow associated with large values of  $\alpha$ .

The convergence speed of (11) decreases for decreasing  $\alpha$ . This is illustrated in 194 Fig. 3 which shows convergence rates  $\rho$  of (11) for various  $\alpha$  together with the corresponding optimal damping factors  $\omega$  determined numerically. Convergence rates

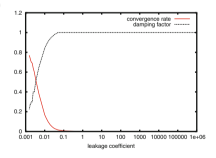


Fig. 3. Convergence rates  $\rho$  and associated optimal damping parameter  $\omega$  over leakage coefficient  $\alpha$ 

deteriorate for  $\alpha < 4 \cdot 10^{-2}$ . Moreover, for  $\alpha < 2 \cdot 10^{-3}$  ill-conditioning of the discretized Richards equation (9) leads to severe problems in the numerical solution. 198 Hence, using the Robin coupling (3) to enforce continuity of pressure by penaliza-

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tion rather than for modelling the clogging effect would require the construction of	200
suitable preconditioners and a careful selection of $\alpha$ .	201

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# An Asymptotic Approach to Compare Coupling **Mechanisms for Different Partial Differential Equations**

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1 Introduction 10

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In many applications the viscous terms become only important in parts of the computational domain. A typical example is the flow of air around the wing of an airplane. 12 It can then be desirable to use an expensive viscous model only where the viscosity is 13 essential for the solution and an inviscid one elsewhere. This leads to the interesting 14 problem of coupling partial differential equations of different types.

The purpose of this paper is to explain several coupling strategies developed over 16 the last decades, and to introduce a systematic way to compare them. We will use the 17 following simple model problem to do so:

$$\mathcal{L}_{ad}u := -vu'' + au' + cu = f \text{ in } \Omega = (-L_1, L_2),$$

$$\mathcal{B}_1u = g_1 \text{ on } x = -L_1,$$

$$\mathcal{B}_2u = g_2 \text{ on } x = L_2.$$
(1)

where v and c are strictly positive constants,  $a, g_1, g_2 \in \mathbf{R}, f \in L^2(\Omega), L_1, L_2 > 0$  19 and  $\mathcal{B}_i$ , j = 1,2 are suitable boundary operators of Dirichlet, Neumann or Robin 20 type. If in part of  $\Omega$ , the diffusion plays only a minor role, one would like to replace 21 the viscous solution u by an inviscid approximation, which leads to two separate 22 problems: a viscous problem on, say,  $\Omega^- := (-L_1, x_0 + \delta)$ , where  $\delta$  stands for the 23 size of the overlap and  $x_0$  the position of the interface,

$$\mathcal{L}_{ad}u_{ad} = f \text{ in } \Omega^{-},$$
  

$$\mathcal{B}_{1}u_{ad} = g_{1} \text{ on } x = -L_{1},$$
(2)

and a pure advection reaction problem on  $\Omega^+ := (x_0, L_2)$ ,

$$\mathcal{L}_a u_a := a u_a' + c u_a = f \qquad \text{in } \Omega^+. \tag{3}$$

Coupling conditions for (2) and (3) need then to be chosen to connect the two sub- 26 problems, and there are many coupling strategies in the literature to choose from. 27

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These strategies have been developed over the last decades for various applications, 28 and sometimes the two different models are really due to different physical phe- 29 nomena, like in fluid-structure interaction problems. In those cases, the coupling 30 conditions are given by the physics, and they are in general unique. We are how- 31 ever interested in problems where the different equations are only chosen in order to 32 achieve computational savings, as for example in [5]:

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The main goal of this paper is to present a computational method for the coupling of two distinct mathematical models describing the same physical phenomenon.

For such couplings, it is quite difficult to decide which coupling strategy from the literature to choose, since every coupling strategy leads to a different solution, and it is 38 not clear a priori which one is the best one. Furthermore, there are neither guidelines 39 nor quantitative comparisons in the literature in order to help with this decision. In 40 order to compare the quality of the various coupling strategies, we propose in this 41 paper a first very natural measure to compare different coupling strategies in such 42 situations, namely to investigate how close the coupled solution for (2) and (3) is to 43 the fully viscous solution of (1). The idea behind this quality measure is that in principle the viscosity should be taken into account everywhere, and hence it is the more 45 expensive viscous solution that we are interested in. However, for computational sav- 46 ings, one would like to use a simpler, non-viscous model whenever the viscosity does 47 not play an important role. In a more general situation, we thus would propose as a 48 natural quality measure to compare the coupled solution to the solution of the expen- 49 sive model used throughout the entire domain, and the closer the coupled solution is 50 to this expensive one, the better the coupling conditions are.

We describe in this paper in detail several coupling strategies for the viscous/in- 52 viscid coupling, and compare them by testing how close the coupled solution is to 53 the fully viscous one: in Sect. 2 we present an overlapping coupling method based on 54 optimization. In Sect. 3 we present several non-overlapping coupling strategies based 55 on coupling conditions at the interface between the two regions. In both sections, the 56 position of the interface needs to be known a priori. This is in contrast to Sect. 4, 57 where we present an adaptive coupling strategy which detects the partition into vis- 58 cous and non-viscous regions automatically. We will see that our quality measure 59 allows us to effectively compare these different strategies, and we find that the best 60 coupled solutions are obtained by judiciously chosen transmission conditions.

## 2 Methods Based on Overlap and Optimization

In this section, we present a very general overlapping coupling strategy that was proposed in [5], where the authors considered as the viscous model the incompressible 64 Navier-Stokes equations, while the inviscid model was the potential equation (the 65 assumption of a small vorticity is made).

For the model problem (1), the coupling strategy works as follows: in each subdo- 67 main, we solve the corresponding equation with a Dirichlet condition at the artificial 68

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interface, 69

$$u_{ad}(x_0 + \delta) = \lambda_1 \text{ and if } a > 0, u_a(x_0) = \lambda_2,$$

and then determine  $(\lambda_1, \lambda_2)$  to be a solution of the optimization problem

$$J(\lambda_1, \lambda_2) := \|u_{ad} - u_a\|_{L^2(x_0, x_0 + \delta)}^2 \longrightarrow \min.$$

The authors in [5] solve this optimization problem using a gradient type method, so 73 that the adjoint equation also needs to be computed. 74

This coupling strategy based on optimization has been studied mathematically in 75 [10] and [2] for our model problem in 2D, see also [6] for a complete description 76 of the algorithms for the model problem, and also for the coupling of Navier-Stokes 77 equations with a Darcy model, or the coupling of the Stokes and potential equations. 78 In [2] other cost functionals to be minimized are proposed. 79

In order to evaluate the quality of this coupling strategy, we compute numerically 80 the error between the viscous and the coupled solution as a function of the viscosity 81 for the case  $L_1 = L_2 = 1$ ,  $x_0 = -0.6$ ,  $f(x) = e^{-1,000(x+1)^2}$  and c = 1. We use a centered 82 finite difference scheme to discretize the two differential operators, with mesh size 83  $2 \times 10^{-5}$ . We consider the case of a positive velocity, a = 1, with  $g_1 = 0$ ,  $g_2 = 0$ ,  $\mathcal{B}_1 = 84$  Id and  $\mathcal{B}_2 = \partial_x - (a - \sqrt{a^2 + 4vc})/2v$  (the absorbing boundary operator) and the 85 case of a negative velocity, a = -1, with  $g_1 = 0$ ,  $g_2 = 0$ ,  $\mathcal{B}_1 = Id$  and  $\mathcal{B}_2 = Id$ . In all 86 experiments presented in this paper, the error in the advection domain  $||u - u_a||_{\Omega^+}$  is 87  $\mathcal{O}(v)$  whatever is the coupling strategy, which is natural, since the advection equation 88 is used instead of the advection-diffusion equation. The numerical error estimate for 89 this overlapping technique in the viscous domain  $\Omega^-$  is given in Table 1. We see that

**Table 1.** Overlapping coupling with optimization: numerically computed error estimate for  $\|u-u_{ad}\|_{\Omega^-}$ 

for a < 0, this coupling strategy (like most of the ones presented in this paper) gives a result  $\mathcal{O}(v)$ , since information is coming from the inviscid approximation in  $\Omega^+$  92 to  $\Omega^-$ , and in  $\Omega^+$  the error  $||u-u_a||_{\Omega^+}$  is  $\mathcal{O}(v)$ .

The non overlapping case  $\delta = 0$  is also considered in [10], namely

$$G(\lambda_1, \lambda_2) = \sigma(a)(u_{ad}(x_0) - u_a(x_0))^2 + (\phi_1 - \phi_2)^2,$$
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where  $\phi_1 = -vu'_{ad}(x_0) + au_{ad}(x_0)$  and  $\phi_2 = au_a(x_0)$  (see Sect. 3.1) and  $\sigma(a) = 1$  if 96 a > 0, 0 otherwise. Using the same numerical setting, we obtain for v small the error 97 estimates shown in Table 2.

Table 2. Non overlapping case with optimization: numerically computed error estimates for  $||u - u_{ad}||_{O^{-}}$ 

### 3 Methods Based on Coupling Conditions

From now on we assume that there is no overlap,  $\delta = 0$ . The coupling techniques in 100 this section are based on coupling conditions, and we will present three strategies: 101 the first one is based on singular perturbation, the second one on boundary layer 102 corrections, and the last one on the factorization of the operator.

### 3.1 Coupling Conditions from Singular Perturbation

In [9] the authors propose to find coupling conditions for (2) and (3) by introducing 105 a regularization of the inviscid problem using a small artificial viscosity  $\varepsilon$ . They thus 106 consider

$$-vw_{\varepsilon}'' + aw_{\varepsilon}' + cw_{\varepsilon} = f \text{ on } (-L_1, x_0),$$
  

$$-\varepsilon v_{\varepsilon}'' + av_{\varepsilon}' + cv_{\varepsilon} = f \text{ on } (x_0, L_2).$$
(4)

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This coupling problem which involves two elliptic equations needs to be completed 108 by two boundary conditions. The first one simply states continuity of the solution: 109  $w_{\varepsilon}(x_0) = v_{\varepsilon}(x_0)$ . For the second one, two choices are possible: we can impose the continuity of the normal flux,  $Vw_{\varepsilon}'(x_0) = \varepsilon v_{\varepsilon}'(x_0)$  (such boundary conditions are 111 called variational conditions) or we impose the continuity of the normal derivative, 112  $w_{\varepsilon}'(x_0) = v_{\varepsilon}'(x_0)$  (called non variational conditions). Letting  $\varepsilon$  tend to 0, it has been 113 rigorously proved in [9] that  $w_{\varepsilon}$  (resp.  $v_{\varepsilon}$ ) tends to  $u_{ad}$  (resp.  $u_a$ ). At the boundary, 114 with the variational conditions, the limiting solution satisfies

$$(-vu'_{ad} + au_{ad})(x_0) = au_a(x_0), \quad u_{ad}(x_0) = u_a(x_0) \quad \text{for} \quad a > 0,$$

$$(-vu'_{ad} + au_{ad})(x_0) = au_a(x_0), \quad \text{for} \quad a < 0,$$

$$(5)$$

while the non variational conditions lead to

$$u_{ad}(x_0) = u_a(x_0), \quad u'_{ad}(x_0) = u'_a(x_0), \quad \text{for} \qquad a > 0,$$
  
 $u_{ad}(x_0) = u_a(x_0), \quad \text{for} \quad a < 0.$  (6)

Rigorous error estimates comparing the coupled solutions obtained with these ap- 117 proaches were obtained in [7], and they are summarized in Table 3, where we ob- 118 serve that the non variational conditions lead to a better coupled solution for positive 119 advection than the variational ones, while for negative advection, again there is no 120 difference between the two approaches. Finally, it has been proved in [6] that the 121 coupling problem with variational conditions is equivalent to the problem using op- 122 timization on  $\sigma(a)(u_{ad}(0)-u_a(0))^2+(\phi_1-\phi_2)^2$ ; our observation is thus consistent. 123

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	a > 0	
	$\mathcal{O}(v^{3/2})$	
Non Variational Conditions	$\mathcal{O}(v^{5/2})$	$\mathcal{O}(v)$

Table 3. Variational versus non-variational coupling conditions: theoretical error estimates for  $||u - u_{ad}||_{Q^{-1}}$ 

### 3.2 Coupling Through Boundary Layer Correction

A different approach, only adding a correction for the boundary layer (in the case 125 a < 0), was proposed in [4]. Here, the authors define the coupled solution of interest 126 to be the solution of the regularized problem (4), and they consider the variational 127 solution obtained from (5) as a first approximation of the regularized one. More precisely the coupled solution is represented as a perturbation of the variational solution 129 in the form

$$w_{\varepsilon}(x) = u_{ad}(x) + r_{\varepsilon}(x),$$

$$v_{\varepsilon}(x) = u_{a}(x) + l_{\varepsilon}(x) + s_{\varepsilon}(x),$$
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where  $l_{\varepsilon}$  is a boundary layer function and  $r_{\varepsilon}$  and  $s_{\varepsilon}$  are the remainders of the asymptotic expansion. The boundary layer term can be computed analytically, but integrals 133 that are involved are then approximated numerically. The numerical solution does 134 not take into account the remainders  $r_{\varepsilon}$  and  $s_{\varepsilon}$  and thus, compared to the solution 135 obtained with (5), the pure advection solution in  $\Omega^+$  is the only one to be corrected. 136

#### 3.3 Coupling Conditions from Operator Factorization

A very accurate set of coupling conditions can be derived from an operator factor- 138 ization, see [7], and requires the solution of a modified advection equation: if we 139 introduce  $\lambda^{\pm} = (a \pm \sqrt{a^2 + 4vc})/2v$ , the advection diffusion equation can be factored, i.e.

$$\mathscr{L}_{ad}u = (\partial_x - \lambda^+)(\partial_x - \lambda^-)u = f,$$
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which gives after integration on  $(x_0, L_2)$ 

$$(\partial_x - \lambda^-)u(x_0) = (\partial_x - \lambda^-)u(L_2)e^{-\lambda^+L_2} + \int_{x_0}^{L_2} f(\sigma)e^{-\lambda^+\sigma}d\sigma.$$

Introducing the new advection equation  $(\partial_x - \lambda^+)\tilde{u}_a = f$ , we find that the viscous 145 solution satisfies 146

$$(\partial_x - \lambda^-)u(x_0) = \tilde{u}_a(x_0) + ((\partial_x - \lambda^-)u(L_2) - \tilde{u}_a(L_2))e^{-\lambda^+ L_2}.$$
 (7)

Solving the advection-diffusion equation in  $\Omega^-$  with the boundary condition (7) (re- 147 placing u by  $u_{ad}$  on the left hand side) would thus yield the exact coupled solution, 148 i.e.  $u_{|\Omega^-} = u_{ad}$ . However the term in  $L_2$  can not be used directly, and one chooses instead  $\tilde{u}_a(L_2)$  to be an expansion of  $(\partial_x - \lambda^-)u(L_2)$  for  $\nu$  small, so that the proposed 150 coupling condition is 151

$$(\partial_{\mathbf{x}} - \lambda^{-}) u_{ad}(\mathbf{x}_{0}) = \tilde{u}_{a}(\mathbf{x}_{0}). \tag{8}$$

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This leads to the coupling procedure

- 1. Solve the new advection equation  $(\partial_x \lambda^+)\tilde{u}_a = f$  on  $(x_0, L_2)$  with  $\tilde{u}_a(L_2) = 153$   $z_0 + z_1 v + \cdots + \mathcal{O}(v^m)$ .
- 2. Solve the advection-diffusion equation on  $(-L_1, x_0)$  with the transmission 155 condition (8).
- 3. Solve the advection equation (3) on  $(x_0, L_2)$  with the condition  $u_{ad}(x_0) = u_a(x_0)$  157 if a > 0.

For our model problem, rigorous error estimates obtained in [7] are shown in Table 4. 159 We see that this coupling strategy leads to a coupled solution which is much closer 160 to the fully viscous one than any of the other strategies. Even in the case of negative 161 advection, one can now obtain approximations more accurate than  $\mathcal{O}(v)$ . Note how- 162 ever that  $\lambda^{\pm}$  are simple constants only in the stationary one dimensional case. In the 163 case of evolution, or for higher dimensions, the  $\lambda^{\pm}$  need to be approximated (see for 164 example [8]).

## 4 The $\chi$ -Formulation

A very different approach for coupling viscous and inviscid problems is proposed in [3]: the method called  $\chi$ -formulation decides automatically where the viscous model and where the inviscid one needs to be used, and solves the equation [3]

$$-v\chi(u'') + au' + cu = f \text{ on } (-L_1, L_2),$$
  
 $u = g_1 \text{ on } x = -L_1,$   
 $\mathcal{B}u = 0 \text{ on } x = L_2,$ 

where the  $\chi$  function is defined by

$$\chi(s) = \begin{cases} 0 & 0 \le s < \delta - \sigma, \\ (s - \delta + \sigma) \frac{\delta}{\sigma} & \delta - \sigma \le s \le \delta, \\ s & s > \delta, \end{cases}$$
 172

so that the diffusion term is neglected as soon as it is small enough. This leads however to a non-linear equation, even if the underlying models are linear, which requires a Newton type algorithm.

In [3], the method is studied for the model problem at the continuous level, and well posedness is proved. Several years later, in [1] and [11], this strategy is used to solve the Navier-Stokes equations. Note that other cut-off functions can also be considered. We show in Table 5 numerically computed error estimates for the  $\chi$ - 179 formulation applied to our model problem.

	a > 0	a < 0
Factorization of the operator	$\mathscr{O}(e^{-a/v})$	$\mathcal{O}(v^m)$

**Table 4.** Coupling based on factorization: theoretical error estimates for  $||u - u_{ad}||_{\Omega^{-}}$ 

	a > 0	
$\chi$ -formulation	$\mathcal{O}(v^{5/2})$	$\mathscr{O}(v)$

**Table 5.**  $\chi$ -formulation: numerically computed error estimate for  $||u-u_{ad}||_{O^{-}}$ 

5 Conclusions 181

For a positive velocity a, among all the strategies presented in this paper, the best 182 coupling condition is provided by the factorization of the operator in the non overlapping case: the error between the corresponding coupled solution and the fully viscous 184 solution is exponentially small. Note that in the unstationary case or in higher dimensions the exponential convergence will be replaced by a polynomial one, because of 186 approximations, an issue we currently investigate. Good algebraically small errors 187 of  $\mathcal{O}(v^{5/2})$  can also be obtained using the non variational conditions (6), or with the 188  $\chi$ -formulation. The other strategies yield less accurate error estimates. When a < 0, 189 the factorization method is the only one to provide a better estimate than  $\mathcal{O}(v)$ . 190

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# **Coupling Geometrically Exact Cosserat Rods and Linear Elastic Continua**

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**Summary.** We consider the mechanical coupling of a geometrically exact Cosserat rod to a 7 linear elastic continuum. The coupling conditions are formulated in the nonlinear rod configuration space. We describe a Dirichlet–Neumann algorithm for the coupled system, and use it 9 to simulate the static stresses in a human knee joint, where the Cosserat rods are models for 10 the ligaments.

## 1 Cosserat Rods and Linear Elasticity

Cosserat rods are models for long slender objects. Let  $SE(3) = \mathbb{R}^3 \rtimes SO(3)$  be the 13 group of orientation-preserving rigid body motions of  $\mathbb{R}^3$  (the special Euclidean 14 group). A configuration of a Cosserat rod is a map  $\varphi: [0,1] \to SE(3)$ . For each 15  $s \in [0,1]$ , the value  $\varphi(s) = (\varphi_r(s), \varphi_q(s))$  is interpreted as the position  $\varphi_r(s) \in \mathbb{R}^3$  and 16 orientation  $\varphi_q(s) \in SO(3)$  of a rigid rod cross section. Strain measures  $(\mathbf{v}_{\varphi}(s), \mathbf{u}_{\varphi}(s))$  17 at  $\varphi(s)$  live in the tangent space  $T_{\varphi(s)}SE(3)$ , and are defined by

$$\mathbf{v}_{\varphi}(s) = \varphi_r'(s)$$
 and  $\varphi_q'(s) = \mathbf{u}_{\varphi}^{\times}(s)\varphi_q(s),$ 

where  $\mathbf{u}_{\varphi}^{\times}$  is the skew-symmetric matrix corresponding to  $\mathbf{u}_{\varphi}$ . On each cross section 19 s of the rod act a resultant force and torque. These are given by a tuple  $(\mathbf{n}(s), \mathbf{m}(s))$ , 20 which is an element of the cotangent space  $T_{\varphi(s)}^*\mathrm{SE}(3)$ . In the absence of external 21 forces and torques we have the equations of equilibrium [6]

$$\mathbf{m}' + \boldsymbol{\varphi}_r' \times \mathbf{n} = 0$$
 on  $[0, 1]$ ,  
 $\mathbf{n}' = 0$  on  $[0, 1]$ .

We assume there to be an energy functional W such that  $\mathbf{n} = \partial W/\partial \mathbf{v}$  and 23  $\mathbf{m} = \partial W/\partial \mathbf{u}$ . Existence of solutions for this model has been shown in [12], but 24 note that solutions may be nonunique.

We will couple the rod model to a linear elastic continuum. Let  $\Omega$  be a domain 26 in  $\mathbb{R}^3$ . Its boundary  $\partial \Omega$  is supposed to be Lipschitz and to consist of disjoint parts 27

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 $\Gamma_N$  and  $\Gamma_D$  such that  $\partial \Omega = \overline{\Gamma}_N \cup \overline{\Gamma}_D$  and  $\Gamma_D$  has positive two-dimensional measure. 28 We use  $\mathbf{v}_{\Omega}$  to denote the outward unit normal of  $\Omega$ . For any displacement function 29  $\mathbf{u} \in \mathbf{H}^1(\Omega) = (H^1(\Omega))^3$  we set  $\boldsymbol{\varepsilon} = \frac{1}{2}(\nabla \mathbf{u} + \nabla \mathbf{u}^T)$  the linear strain tensor and the 30 stress  $\boldsymbol{\sigma} = \boldsymbol{\sigma}(\boldsymbol{\varepsilon})$ , with a St. Venant–Kirchhoff-type material law

$$\boldsymbol{\sigma}(\boldsymbol{\varepsilon}) = \frac{E v}{(1+v)(1-2v)} (\operatorname{tr} \boldsymbol{\varepsilon}) \operatorname{Id} + \frac{E}{1+v} \boldsymbol{\varepsilon}.$$

The parameters E and  $\nu$  are the Young's modulus and Poisson ratio, respectively. The boundary value problem of elasticity is then

$$-\operatorname{div} \boldsymbol{\sigma}(\mathbf{u}) = \mathbf{f} \qquad \text{in } \Omega,$$

$$\mathbf{u} = 0 \qquad \text{on } \Gamma_D,$$

$$\boldsymbol{\sigma}(\mathbf{u}) \boldsymbol{v}_{\Omega} = \mathbf{t} \qquad \text{on } \Gamma_N,$$

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with volume forces  $\mathbf{f}: \Omega \to \mathbb{R}^3$  and surface force  $\mathbf{t}: \Gamma_N \to \mathbb{R}^3$ .

## 2 Coupling Conditions

We will now derive conditions for the coupling of a Cosserat rod and a linear elastic 36 three-dimensional object. The two main difficulties are the difference in dimensions 37 between the rod and the continuum, and the nonlinear nature of the rod configuration 38 space.

Previous work has mainly focused on coupling linear models of different 40 dimensions. Lagnese et al. [7] have studied the coupling of beams to plates exten- 41 sively. Modeling of 3d-2d junctions between linear elastic objects using a method of 42 asymptotic expansion has been carried out by Ciarlet et al. [4]. Monaghan et al. [8] 43 describe a 3d-1d coupling between linear elastic elements in the discrete setting. A 44 general framework which encompasses these cases is given in [3]. We are not aware 45 of previous work on the coupling of Cosserat rods.

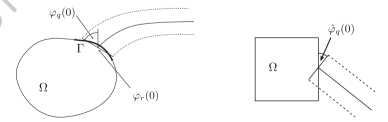


Fig. 1. Left: Coupling between a two-dimensional domain and a rod. Right: In the stress-free configuration the rod may meet the body at an arbitrary spatial angle  $\hat{\varphi}_a(0)$ 

Consider again a linear elastic continuum defined on a reference configuration 47  $\Omega$ . This time, the boundary  $\partial \Omega$  is supposed to consist of three disjoint parts  $\Gamma_D$ ,  $\Gamma_N$ , 48 and  $\Gamma$  such that  $\partial \Omega = \overline{\Gamma}_D \cup \overline{\Gamma}_N \cup \overline{\Gamma}$ . We assume that  $\Gamma_D$  and  $\Gamma$  have positive twodimensional measure. The three-dimensional object represented by  $\Omega$  will couple 50 with the rod across  $\Gamma$ , which we call the coupling boundary. The boundary of the 51 parameter domain [0,1] of a Cosserat rod consists only of the two points 0 and 1, and 52 the respective domain normals are  $\mathbf{v}_{r,0} = -1$  and  $\mathbf{v}_{r,1} = 1$ . To be specific, we pick 0 as 53 the coupling boundary. We assume a stress-free rod configuration  $\hat{\varphi}: [0,1] \to SE(3)$  54 such that  $\hat{\varphi}_r(0) = |\Gamma|^{-1} \int_{\Gamma} x \, ds$ , i.e., the coupling interface of the rod in its stress-free 55 state is placed at the center of gravity of the coupling interface of  $\Omega$ . The orientation 56  $\hat{\varphi}_a(0)$  of the stress-free state does not need to be in any relation with the shape of the 57 coupling boundary  $\Gamma$  (Fig. 1).

We define our coupling using a set of conditions for the primal variables. These 59 variables are the configuration  $\varphi$  of the rod and the displacement field **u** of the continuum. It is well known that when coupling two continuum models of the same type, 61 the solution has to be continuous [9]. Since the position  $\varphi_r(0) \in \mathbb{R}^3$  of the coupling 62 cross-section can be seen as an averaged position it is natural to couple it to the 63 averaged position of  $\Gamma$ 

$$\varphi_r(0) \stackrel{!}{=} \frac{1}{|\Gamma|} \int_{\Gamma} (\mathbf{u}(x) + x) ds. \tag{1}$$

To obtain a complete set of primal conditions we also need to relate the orientations at the interface. This requires some technical preparations. Using the deformation gradient  $F(\mathbf{u}) = \nabla(\mathbf{u} + \mathrm{Id})$  we first define the average deformation of the 67 interface boundary  $\Gamma$  as  $\mathscr{F}(\mathbf{u}) = |\Gamma|^{-1} \int_{\Gamma} \nabla(\mathbf{u}(x) + x) ds$ . If  $\mathbf{u}$  stays within the limits of linear elasticity the matrix  $\mathscr{F}(\mathbf{u})$  has a positive determinant. Using the polar 69 decomposition it can then be split into a rotation polar  $(\mathcal{F}(\mathbf{u}))$  and a stretching. We 70 define the average orientation of  $\Gamma$  induced by a deformation **u** as the rotational part 71 of  $\mathscr{F}(\mathbf{u})$ . This corresponds to the definition of the continuum rotation used in the 72 theory of Cosserat continua. In particular, if  $\mathbf{u} \equiv 0$  then polar( $\mathscr{F}(\mathbf{u})$ ) = Id.

The average orientation polar  $(\mathcal{F}(\mathbf{u}))$  can now be set in relation to  $\varphi_q(0)$ , the 74 orientation of the rod cross-section at s = 0. We require the coupling condition to be 75 fulfilled by the stress-free configuration  $\mathbf{u} = 0$ ,  $\varphi = \hat{\varphi}$ . This leads to the condition

$$\varphi_q(0) \stackrel{!}{=} \operatorname{polar}(\mathscr{F}(\mathbf{u}))\hat{\varphi}_q(0),$$
(2)

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which is an equation in the nonlinear three-dimensional space SO(3).

For ease of writing we will introduce the averaging operator Av :  $\mathbf{H}^1(\Omega) \to \mathrm{SE}(3)$  78 by setting 79

$$\operatorname{Av}(\mathbf{u}) = \left(\frac{1}{|\Gamma|} \int_{\Gamma} (\mathbf{u}(x) + x) \, ds, \operatorname{polar}(\mathscr{F}(\mathbf{u})) \, \hat{\varphi}_q(0) \right), \tag{3}$$

where we have used  $(\cdot,\cdot)$  to denote elements of the product space SE(3) =  $\mathbb{R}^3$  × SO(3). It is a nonlinear generalization of the restriction operator used in [3]. Then (1) 81 and (2) can be written concisely as 82

$$\varphi(0) \stackrel{!}{=} Av(\mathbf{u}). \tag{4}$$

Note that we do not assume that  $\Gamma$  has the same shape or area as the rod cross-section 83 at s = 0. Also, since the coupling conditions relate only finite-dimensional quantities 84 they remain the same when the subdomain problems are replaced by finite element 85 approximations.

The coupling problem is made complete by conditions for the dual variables. 87 For the continuum these variables are the normal stresses at the boundary  $\Gamma$ . For 88 the rod the dual variables are the total force  $\mathbf{n}(0)\mathbf{v}_{r0}$  and the total moment  $\mathbf{m}(0)\mathbf{v}_{r0}$  89 about  $\varphi_r(0)$  transmitted in normal direction across the cross-section at s=0. We 90 expect these to match the total force and torque exerted by the continuum across the 91 coupling boundary  $\Gamma$  in the direction of  $-\mathbf{v}_{\Omega}$ 

$$\int_{\Gamma} \boldsymbol{\sigma}(\mathbf{u}) \boldsymbol{\nu}_{\Omega} \, ds = -\mathbf{n}(0) \boldsymbol{\nu}_{r,0} \tag{5}$$

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$$\int_{\Gamma} (\mathbf{x} - \boldsymbol{\varphi}_r(0)) \times (\boldsymbol{\sigma}(\mathbf{u}) \boldsymbol{\nu}_{\Omega}) ds = -\mathbf{m}(0) \boldsymbol{\nu}_{r,0}.$$
 (6)

Together, these equations relate quantities in the six-dimensional space  $T_{\alpha(0)}^*SE(3)$ .

Remark 1. A variational formulation suggests that (5) and (6) are not the dual conditions of (4) (cf. to [3] for the linear case). Together with (10), however, they are 95 sufficient to construct a working solution algorithm. 96

## 3 A Dirichlet-Neumann Algorithm

In this section we present a Dirichlet-Neumann algorithm for the coupled problem. 98 It can be interpreted as a fixed-point iteration for an equation on the trace space of the rod configuration space at s = 0, i.e. on SE(3). Each iteration consists of three steps: 100 a Dirichlet problem for the rod, a Neumann problem for the body, and a damped 101 update along geodesics on SE(3). Let  $\lambda^0 \in SE(3)$  be the initial interface value and 102  $k \ge 0$  the iteration number. In more detail, the steps are as follows.

1. Dirichlet problem for the Cosserat rod 104 Let  $\lambda^k$ ,  $\varphi_D \in SE(3)$  be the current interface value and a Dirichlet boundary value, 105 respectively. Find a solution  $\varphi^{k+1}$  of the Dirichlet rod problem 106

$$\begin{split} (\mathbf{m}^{k+1})' + (\varphi_r^{k+1})' \times \mathbf{n}^{k+1} &= 0 & \text{on } [0,1] \\ (\mathbf{n}^{k+1})' &= 0 & \text{on } [0,1] \\ \varphi^{k+1}(0) &= \lambda^k \\ \varphi^{k+1}(1) &= \varphi_D. \end{split}$$

2. Neumann problem for the continuum

The new rod iterate  $\varphi^{k+1}$  exerts a resultant force  $\mathbf{n}^{k+1}(0)\mathbf{v}_{r,0}$  and moment 108  $\mathbf{m}^{k+1}(0)\mathbf{v}_{r,0}$  across its cross-section at s=0. Construct a Neumann data field 109  $\boldsymbol{\tau}^{k+1}: \Gamma \to \mathbb{R}^3$  such that 110

$$\int_{\Gamma} \boldsymbol{\tau}^{k+1}(x) \, ds = -\mathbf{n}^{k+1}(0) \boldsymbol{\nu}_{r,0} \tag{7}$$

and 111

$$\int_{\Gamma} (x - \varphi_r^{k+1}(0)) \times \boldsymbol{\tau}^{k+1}(x) \, ds = -\mathbf{m}^{k+1}(0) \boldsymbol{\nu}_{r,0}. \tag{8}$$

Then solve the three-dimensional linear elasticity problem with Neumann data 112  $\boldsymbol{\tau}^{k+1}$  on  $\Gamma$ 113

$$-\operatorname{div} \boldsymbol{\sigma}(\mathbf{u}^{k+1}) = \mathbf{f} \qquad \text{in } \Omega$$

$$\boldsymbol{\sigma}(\mathbf{u}^{k+1}) \boldsymbol{v}_{\Omega} = \boldsymbol{\tau}^{k+1} \qquad \text{on } \Gamma$$

$$\mathbf{u}^{k+1} = 0 \qquad \text{on } \Gamma_{D}$$

$$\boldsymbol{\sigma}(\mathbf{u}^{k+1}) \boldsymbol{v}_{\Omega} = \mathbf{t} \qquad \text{on } \Gamma_{N}.$$
(9)

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### 3. Damped geodesic update

From the solution  $\mathbf{u}^{k+1}$  compute the average interface displacement and orientation Av( $\mathbf{u}^{k+1}$ ) as defined in (3). With a damping parameter  $\theta > 0$ , the new 116 interface value  $\lambda^{k+1}$  is then computed as a geodesic combination in SE(3) of the old value  $\lambda^k$  and  $Av(\mathbf{u}^{k+1})$ ,

$$\lambda^{k+1} = \exp_{\lambda^k} \theta \left[ \exp_{\lambda^k}^{-1} \operatorname{Av}(\mathbf{u}^{k+1}) \right].$$

It remains to say how to construct suitable fields of Neumann data  $\tau^{k+1}$  that 119 satisfy the conditions (7) and (8). Let us drop the index k for simplicity. In principle, 120any function  $\tau: \Gamma \to \mathbb{R}^3$  of sufficient regularity fulfilling (7) and (8) can be used as 121 Neumann data in (9). It has been shown in [10] that such functions exist.

The theory of Cosserat rods assumes that forces and moments are transmitted 123 evenly across cross-sections. We therefore construct  $\tau$  to be 'as constant as possible'. 124 More formally, we introduce the functional 125

$$T: \mathbf{L}^2(\Gamma) \times \mathbb{R}^3 \to \mathbb{R}, \qquad T(\mathbf{h}, \mathbf{c}) = \int_{\Gamma} ||\mathbf{h}(x) - \mathbf{c}||^2 ds,$$

and construct  $\tau$  as the solution of the minimization problem

$$(\boldsymbol{\tau}, \mathbf{c}_{\boldsymbol{\tau}}) = \underset{\mathbf{h} \in \mathbf{L}^{2}(\Gamma)}{\arg \min} T(\mathbf{h}, \mathbf{c})$$
(10)

under the constraints that

$$\int_{\Gamma} \boldsymbol{\tau} ds = -\mathbf{n}(0) \boldsymbol{v}_{r,0} \quad \text{and} \quad \int_{\Gamma} (x - \varphi_r(0)) \times \boldsymbol{\tau} ds = -\mathbf{m}(0) \boldsymbol{v}_{r,0}. \quad (11)$$

Problem (10) and (11) is a convex minimization problem with linear equality 128 constraints. In [10, Lemma 5.3.4] it was shown that there exists a unique solution. In a finite element setting the problem size is given by the number of grid vertices on 130  $\Gamma$  times 3. A minimization problem of this type can be solved, e.g., with an interiorpoint method. 132 Oliver Sander

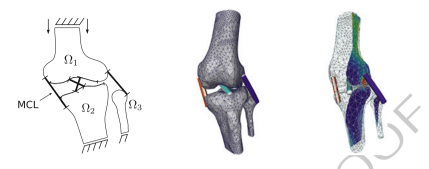


Fig. 2. Left: Problem setting. Tibia and fibula are rotated 15° in valgus direction to put additional stress on the MCL. Center: Deformed grids after two adaptive refinement steps. Right: Two sagittal cuts through the von Mises stress field

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### 4 Numerical Results

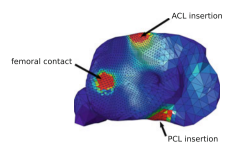
We close with a simulation result for a knee model which combines femur, tibia, and 134 fibula bones modeled as three-dimensional linear elastic objects, and the cruciate and 135 collateral ligaments, modeled as Cosserat rods. The model additionally includes the 136 contact between femur and tibia. To obtain a test case where the contact stresses do 137 not entirely predominate the stresses created in the bone by pulling ligaments, we 138 applied a valgus rotation of 15° to tibia and fibula. This leads to a high strain in 139 the medial collateral ligament (MCL) and can be interpreted as an imminent MCL 140 rupture (Fig. 2).

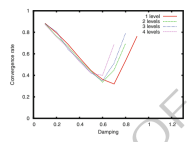
The geometry was obtained from the Visible Human data set. We modeled bone 142 with an isotropic, homogeneous, linear elastic material with E=17 GPa and v=0.3. 143 The distal horizontal sections of tibia and fibula were clamped, and a prescribed 144 downward displacement of 2 mm was applied to the upper section of the femur. We 145 used first-order finite elements for the discretization of the linear elasticity problem. 146 DUNE [2] was used for the implementation.

The four ligaments were each modeled by a single Cosserat rod with a circular 148 cross-section of radius 5 mm. The rod equations were discretized using geodesic 149 finite elements [11]. We chose a linear material law (see, e.g., [6]) with parameters 150  $E = 330 \, \text{MPa}$  and v = 0.3. On the bones, the coupling boundaries  $\Gamma$  for the different ligaments were marked by hand using a graphical editor. We modeled all ligaments 152 to be straight in their stress-free configurations and to have 8 % in situ strain.

We solved the combined problem using the Dirichlet-Neumann algorithm 154 described in Sect. 3. At each iteration, a pure Dirichlet problem had to be solved 155 for each of the rods and a contact problem with mixed Dirichlet-Neumann bound- 156 ary conditions had to be solved for the bones. The contact problem was solved using 157 the Truncated Nonsmooth Newton Multigrid (TNNMG) algorithm [5]. The TNNMG 158 method solves linear contact problems with the efficiency of linear multigrid. For the 159 ligaments we used a Riemannian trust-region solver [1, 11], and we used IPOpt [13] 160 to solve the minimization problems (10) and (11). Figure 2 shows the deformed con-

#### Coupling Geometrically Exact Cosserat Rods and Linear Elastic Continua





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Fig. 3. Left: Stress plot on the tibial plateau. Right: Convergence rates of the Dirichlet-Neumann method as a function of the damping parameter for up to four grid levels

figuration on a grid obtained by two steps of adaptive refinement and cuts through 162 the von Mises stress field. In Fig. 3, left, a caudal view onto the tibial plateau can be 163 seen, which is colored according to the von Mises stress. The peaks due to contact 164 and the pull of the cruciate ligaments can be clearly observed.

We measured the Dirichlet-Neumann convergence rates with bone grids obtained 166 by up to three steps of adaptive refinement using the hierarchical error estimator pre- 167 sented in [10]. Rod grids in turn were refined uniformly. On each new set of grids 168 we started the computation from the reference configuration. That way identical initial iterates for all grid refinement levels were obtained. Details on the measuring 170 setup can be found in [10]. Figure 3, right, shows the Dirichlet–Neumann conver- 171 gence rates plotted as a function of the damping parameter  $\theta$  for up to four levels 172 of refinement. For each further level of refinement, the optimal convergence rate 173 is slightly worse than for the previous, and obtained for a slightly lower damping 174 parameter. This behavior seems typical for Dirichlet-Neumann methods. Neverthe- 175 less the optimal convergence rates stay around 0.4. This makes the algorithm well 176 usable in practice.

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# **Parareal Schwarz Waveform Relaxation Methods**

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1 Introduction 12

Solving an evolution problem in parallel is naturally undertaken by trying to paral- 13 lelize the algorithm in space, and then still follow a time stepping method from the 14 initial time t = 0 to the final time t = T. This is especially easy to do when an explicit 15 time stepping method is used, because in that case the time step for each component 16 is only based on past, known data, and the time stepping can be performed in an 17 embarrassingly parallel way. If one uses implicit time stepping however, one obtains 18 a large system of coupled equations, and thus the linear or non-linear solver needs to be parallelized, e.g. using a domain decomposition method.

Over the last decades, people have however also tried to parallelize algorithms 21 in the time direction. One example is Womble's algorithm [22], where the systems 22 arising from an implicit time discretization are solved using an iterative method, and 23 the iteration of the next time level is started, before the iteration on the current time 24 level has converged. It is then possible to iterate several time levels simultaneously, 25 but the possible gain using a parallel computer is only small, see for example [3].

A different approach to obtain small scale parallelism in time is to use predictor- 27 corrector methods, where the prediction step and the correction step can be per- 28 formed by two (or several) processors in parallel, if organized properly. An entire 29 class of such methods has been proposed in [19], and good small scale parallelism 30 can be achieved.

A third, very different approach are the waveform relaxation algorithms, invented 32 in [15], which are based on a decomposition of the system to be solved into subsystems. An iteration is then used, which solves time dependent problems in each subsystem and communicates information at interfaces to neighboring subsystems to 35 converge to the overall solution in space-time [12, 13]. Substantial progress has been 36 made on such methods for evolution PDEs, see for example [5, 6, 14], and references 37

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therein. If a multi-grid decomposition is used, instead of a domain decomposition, 38 one obtains the so called parabolic multi-grid methods [11], which are also called 39 multi-grid waveform relaxation methods. For further results, see [17, 21].

Finally, the last class of methods, which focuses entirely on the parallelization in the time direction, are based on shooting methods in time. A first historical step in this direction is [20], and for an early analysis see [2]. The newest algorithm in this class is the parareal algorithm, invented in [16]. For a complete historical overview of such methods, further references, and a precise convergence estimate of the parareal algorithm see [4, 9].

We propose here a space time parallel algorithm for solving evolution partial 47 differential equations, and use as a model problem 48

$$\partial_{t} u = \partial_{xx} u \text{ in } \Omega = (0,1) \times (0,T), 
\mathscr{B}^{-} u(0,t) = g_{0}(t) \ t \in (0,T), 
\mathscr{B}^{+} u(1,t) = g_{1}(t) \ t \in (0,T), 
u(x,0) = u_{0}(x) \ x \in \Omega.$$
(1)

Here  $\mathscr{B}^{\pm}$  represent some boundary operators, like the identity for a Dirichlet condition, or a normal derivative for a Neumann condition. The algorithm is based on a decomposition of the space-time domain into space-time subdomains, as indicated in Fig. 1. In order to solve an evolution problem by only solving problems in small space-time domains, one has to iteratively calculate more and more accurate initial and boundary conditions for each space-time subdomain. The parareal Schwarz waveform relaxation algorithm does this by using a parareal approximation for the initial conditions, and a Schwarz waveform relaxation algorithm for the boundary conditions. For a different variant of combining a spatial and a time decomposition, 57 see [18].

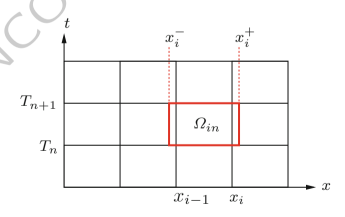


Fig. 1. Space time decomposition for the parareal Schwarz waveform relaxation algorithm

### 2 Parareal Schwarz Waveform Relaxation Algorithms

The parareal algorithm for the model problem (1) is based on a decomposition of the 60 time interval (0,T) into subintervals, given by  $0 = T_0 < T_1 < T_2 < \ldots < T_N = T$ , 61 and the algorithm is defined using two propagation operators: a coarse operator 62  $G(t_2,t_1,u_1,g_0,g_1)$  which provides a rough approximation of the solution  $u(x,t_2)$  63 of (1) with a given initial condition  $u(x,t_1)=u_1(x)$  and boundary conditions  $g_0$  64 and  $g_1$ , and a fine operator  $F(t_2,t_1,u_1,g_0,g_1)$ , which gives a more accurate ap- 65 proximation of the same solution with initial condition  $u(x,t_1) = u_1(x)$  and boundary conditions  $g_0$  and  $g_1$ . Starting with a first approximation  $U_n^0$  at the time points 67  $T_0, T_1, T_2, \dots, T_{N-1}$ , the parareal algorithm performs for  $k = 0, 1, 2, \dots$  the correction 68 iteration

$$U_{n+1}^{k+1} = F(T_{n+1}, T_n, U_n^k, g_0, g_1) + G(T_{n+1}, T_n, U_n^{k+1}, g_0, g_1) - G(T_{n+1}, T_n, U_n^k, g_0, g_1),$$
(2)

which is nothing else than a multiple shooting method with an approximate Jacobian 70 in the Newton step, see for example [9], which also contains a precise convergence 71 estimate for the case of the heat equation, or [4] for a similar precise convergence 72 estimate for the case of nonlinear problems.

In contrast to the parareal algorithm, a Schwarz waveform relaxation method 74 for the model problem (1) is based on a spatial decomposition only, in the most 75 general case into overlapping subdomains  $\Omega = \bigcup_{i=1}^{I} (x_i^-, x_i^+)$ , as shown in Fig. 1. 76 Here the boundaries  $x_i^{\pm}$  of the overlapping subdomains are constructed from a nonoverlapping decomposition given by the decomposition  $0 =: x_0 < x_1 < ... < x_I := 1$ , 78 by adding and subtracting half the overlap,  $x_i^- := x_{i-1} - \frac{L}{2}$ ,  $x_i^+ := x_i + \frac{L}{2}$ , except for 79 the first and last point,  $x_1^- = x_0$  and  $x_I^+ = x_I$ . Given an initial guess at the interfaces, 80 say  $\mathscr{B}_{i}^{\pm}u_{i}^{0}$ , the Schwarz waveform relaxation algorithm solves iteratively for k=81 $1, 2, \dots$  the subdomain problems

$$\partial_{t}u_{i}^{k} = \partial_{xx}u_{i}^{k} & \text{in } \Omega_{i} \times (0,T), \\
u_{i}^{k}(x,0) = u_{0} & \text{in } \Omega_{i}, \\
\mathscr{B}_{i}^{-}u_{i}^{k}(x_{i}^{-},t) = \mathscr{B}_{i}^{-}u_{i-1}^{k-1}(x_{i}^{-},t) \ t \in (0,T), \\
\mathscr{B}_{i}^{+}u_{i}^{k}(x_{i}^{+},t) = \mathscr{B}_{i}^{+}u_{i+1}^{k-1}(x_{i}^{+},t) \ t \in (0,T).$$
(3)

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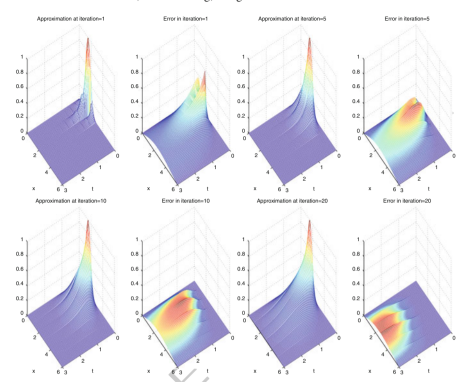
Here again, the operators  $\mathscr{B}_i^{\pm}$  are transmission operators: in the case of the identity, 83 we have the classical Schwarz waveform relaxation algorithm; for Robin or higher 84 order transmission conditions, one would obtain an optimized Schwarz waveform 85 relaxation algorithm, if the parameters in the transmission conditions are chosen to 86 optimize the convergence of the algorithm, see [1, 5].

Parareal Schwarz waveform relaxation algorithms combine the two techniques 88 for a general space-time decomposition given in Fig. 1. We propose among the many 89 possibilities the following one: given initial conditions  $u_{0,i,n}^k(x)$  and boundary conditions  $\mathscr{B}_i^- u_{i-1,n}^k(t)$  and  $\mathscr{B}_i^+ u_{i+1,n}^k(t)$  for  $i=1,2,\ldots,I$  and  $n=1,2,\ldots,N$  we compute 91

1. All accurate approximations  $u_{i,n}^{k+1}(x,t) := F_{i,n}(u_{0,i,n}^k, \mathcal{B}_i^- u_{i-1,n}^k, \mathcal{B}_i^+ u_{i+1,n}^k)$  in par- 92 allel using the more accurate evolution operator.

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**Fig. 2.** Illustration how the parareal Schwarz waveform relaxation algorithm removes the error over several iterations: each plot pair shows on the *left* the approximation and on the *right* the error (i.e. the difference between the monodomain solution and the current iterate) for k = 1, 5, 10, 20

2. For n = 0, 1, ..., new initial conditions using a parareal integration step both in space and time, 95

$$\begin{aligned} u_{0,i,n+1}^{k+1} &= u_{i,n}^{k+1}(\cdot, T_{n+1}) + G_{i,n}(u_{0,i,n}^{k+1}, \mathcal{B}_i^- u_{i-1,n}^{k+1}, \mathcal{B}_i^+ u_{i+1,n}^{k+1}) \\ &- G_{i,n}(u_{0,i,n}^k, \mathcal{B}_i^- u_{i-1,n}^k, \mathcal{B}_i^+ u_{i+1,n}^k). \end{aligned}$$

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An example on how this algorithm converges is given in Fig. 2.

We present now a first convergence result for the parareal Schwarz waveform 97 relaxation algorithm: 98

**Theorem 1 (Superlinear Convergence).** Let  $F_{i,n}$  be the exact solution,  $G_{i,n}$  be a 99 backward Euler approximation in time, and the exact solution in space, and assume 100 a decomposition of the spatial domain into two overlapping subdomains. If the algorithms uses Dirichlet transmission conditions, i.e.  $\mathcal{B}_i^{\pm} = I$ , the identity, then it 102 converges superlinearly to the solution of the underlying problem. 103

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The proof of this theorem is too long and technical for this short paper, and will 104 appear in [7]. We present however a detailed numerical study of how the algorithm 105 depends on the various parameters in the following section.

### 3 Numerical Results

In all our experiments, except otherwise mentioned, we use the domain  $\Omega = (0,6)$  108 and the time interval (0,T) with T=3, and discretize the heat equation with a centered finite difference discretization in space with  $\Delta x = \frac{1}{10}$ , and a backward Euler 110 discretization in time, with  $\Delta t = \frac{3}{100}$ , and we use a decomposition into 6 equal spatial subdomains with overlap  $2\Delta x$ .

We start with the dependence on the number of time subintervals. In Fig. 3 on the 113 left, we show the convergence of the algorithm when 1 (classical Schwarz waveform 114 relaxation), 2, 4 and 10 time subintervals are used. This shows that the algorithm 115 is quite insensitive to the number of time subintervals used. We also observe the 116 typical superlinear convergence behavior of all waveform relaxation algorithms, see 117 for example [8].

We next investigate how the convergence depends on the total time interval 119 length T. For this experiment, leaving all other parameters the same, we choose 120  $T \in \{0.1, 0.2, 0.4, 0.8, 1.6, 3.2\}, \Delta t = \frac{T}{100}$ , and ten time subintervals for each simula-121 tion. The results are shown in Fig. 3 on the right. We clearly see that convergence is 122 much faster on short time intervals, compared to long time intervals.

In order to test the dependence on the number of spatial subdomains, we use 124 again all parameters as before, but now decompose the domain into 2, 3, 6 and 125 12 spatial subdomains, and again 10 time subintervals. We see in Fig. 4 on the left 126 that using more spatial subdomains makes the algorithm converge more slowly. This 127 can however be remedied by using smaller global time intervals, as for the Schwarz 128 waveform relaxation algorithm, see [10].

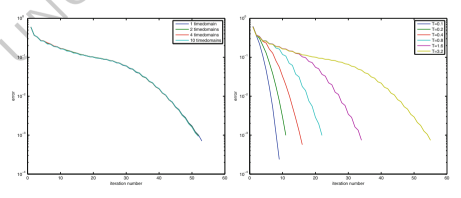


Fig. 3. Dependence of the parareal Schwarz waveform relaxation algorithm on the number of time subintervals on the *left*, and the total time window length on the *right* 

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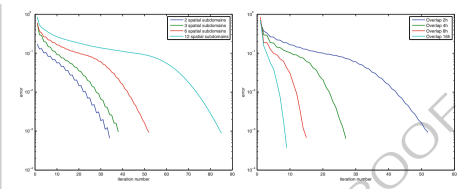


Fig. 4. Dependence of the parareal Schwarz waveform relaxation algorithm on the number of spatial subdomains on the left, and the overlap on the right

We finally test the dependence on the overlap, using  $2\Delta x$ ,  $4\Delta x$ ,  $8\Delta x$  and  $16\Delta x$  130 for the overlap. We see on the right in Fig. 4 that increasing the overlap substantially 131 improves the convergence speed of the algorithm. This increases however also the 132 cost of the method, since bigger subdomain problems need to be solved.

A better approach is to use optimized transmission conditions, see for example 134 [1, 5]. Using the same configuration as in the previous experiment, and  $2\Delta x$  overlap, 135 we obtain with first order transmission conditions and choosing the parameters p=1, 136 q = 1.75 (for terminology, see (1)) the result shown in Fig. 5. This illustrates well that using optimized transmission conditions can lead to even better performance of the 138 algorithm than very generous overlap, at no additional cost, since the subdomain size 139 and matrix sparsity is the same as for the case of Dirichlet transmission conditions. 140 In addition we observe that now the convergence has become more linear, and the

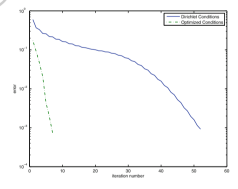


Fig. 5. Comparison of the parareal Schwarz waveform relaxation algorithm with Dirichlet and optimized transmission conditions

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algorithm does not depend significantly any more on the superlinear convergence 142 mechanism essential with Dirichlet transmission conditions. 143

4 Conclusion 144

We presented a general parareal Schwarz waveform relaxation algorithm, which is 145 based on a decomposition in space and time of a given evolution problem, in order 146 to increase parallelism. We stated a theoretical convergence result, whose proof will 147 appear elsewhere, and then illustrated the dependence of the algorithm on the space- 148 time decomposition configuration, which revealed that for fast convergence, either 149 short time intervals, large overlap, or optimized transmission conditions need to be 150 used. We are currently working on precise convergence factor estimates, a variant 151 of the algorithm which also uses a coarse spatial mesh, and the addition of a coarse 152 propagation mechanism over many spatial subdomains.

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# A Parallel Overlapping Time-Domain Decomposition **Method for ODEs**

Stefan Giittel 4

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Summary. We introduce an overlapping time-domain decomposition for linear initial-value 7 problems which gives rise to an efficient solution method for parallel computers without 8 resorting to the frequency domain. This parallel method exploits the fact that homogeneous 9 initial-value problems can be integrated much faster than inhomogeneous problems by using 10 an efficient Arnoldi approximation for the matrix exponential function.

1 Introduction 12

We are interested in the parallel solution of a linear initial-value problem

$$u'(t) = Au(t) + g(t), \quad t \in [0, T], \quad u(0) = u_0,$$
 (1)

where  $A \in \mathbb{R}^{N \times N}$  is a possibly large (and sparse) matrix and  $u, g : t \mapsto \mathbb{R}^{N}$ . Throughout this paper we assume that the function g(t) is a source term which is difficult 15 to integrate numerically (e.g., highly oscillating or given by a slow computer subroutine). For example, if (1) arises from the space discretization of a heat-diffusion 17 problem, then A represents a diffusion operator and g(t) is a time-dependent heat 18 source.

Problems of the above form arise often in scientific computing, and various solu- 20 tion methods for parallel computers have been proposed in the literature. A popular 21 approach (see, e.g., [1, 8]) is based on the Laplace-transformed equation

$$s\widehat{u}(s) - u_0 = A\widehat{u}(s) + \widehat{g}(s)$$

and the contour integral representation of the inverse transformation

$$u(t) = \frac{1}{2\pi i} \int_{\Gamma} e^{ts} \widehat{u}(s) \, \mathrm{d}s,$$
 25

with a suitable contour  $\Gamma$  surrounding the singularities of  $\widehat{u}(s)$  (which are the eigenvalues of A and all singularities of  $\widehat{g}(s)$ ). Discretization of this integral by a quadra- 27 ture formula with complex nodes  $s_i$  and weights  $w_i$  yields 28

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$$u(t) \approx \sum_{j=1}^{p} w_j \widehat{u}(s_j) = \sum_{j=1}^{p} w_j (s_j I - A)^{-1} (u_0 + \widehat{g}(s_j)).$$
 29

This method is suitable for parallel computation because the p complex shifted linear 30 systems are decoupled. On the other hand, there are obvious drawbacks such as the 31 introduction of complex arithmetic into a real problem and the need for calculating 32  $\widehat{g}(s_i)$ . Moreover, many nodes  $s_i$  may be required to represent a stiff source g(t) to 33 prescribed accuracy.

Another approach, perhaps closest in spirit to the method described here, is 35 known as exponential quadrature. It is based on the variation-of-constants formula

$$u(t) = e^{tA}u_0 + \int_0^t e^{(t-\tau)A}g(\tau) d\tau$$
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and the approximation of the integrand by a quadrature rule in nodes  $\tau_1, \ldots, \tau_p$ . This 38 yields p + 1 independent matrix exponentials

$$e^{tA}u_0$$
 and  $e^{(t- au_j)A}g( au_j)$  for  $j=1,\ldots,p,$ 

each of which may be approximated efficiently by a Krylov method (see the discus- 41 sion in Sect. 3). However, exponential quadrature is impractical if the source term 42 g(t) is stiff enough so that too many quadrature nodes are needed.

To overcome the problems mentioned above, we propose in Sect. 2 a decom- 44 position of (1) into subproblems on overlapping time intervals. These subproblems 45 are decoupled and can be assigned to independent processors. Our method requires 46 almost no communication or synchronization between the processors, except a summation step at the end of the algorithm. Another advantage of our method is its 48 ease of implementation; any available serial integrator for (1) can be used in black- 49 box fashion. Because the efficiency of our method relies on the fast integration of 50 homogeneous linear initial-value problems, Sect. 3 contains a brief discussion of the 51 Arnoldi method for computing the matrix exponential function. In Sect. 4 we discuss 52 the error control and parallel efficiency of our method. In Sect. 5 we present results 53 of a numerical experiment.

## 2 Overlapping Time-Domain Decomposition

On a time grid  $\{T_j = jT/p : j = 0,...,p\}$  we decompose (1) into the following 56 subproblems of two types.

Type 1 : For 
$$j = 1, ..., p$$
 solve

$$v_i'(t) = Av_i(t) + g(t), \quad v_i(T_{i-1}) = 0, \quad t \in [T_{i-1}, T_i],$$
 59

using some serial integrator.

Type 2: For j = 1, ..., p solve

$$w'_{i}(t) = Aw_{j}(t), \quad w_{j}(T_{j-1}) = v_{j-1}(T_{j-1}), \quad t \in [T_{j-1}, T],$$
 62

$$W_j(t) = AW_j(t), \quad W_j(I_{j-1}) = V_{j-1}(I_{j-1}), \quad t \in [I_{j-1}, I],$$

using exponential propagation (we set  $v_0(T_0) := u_0$ ).

Note that the p subproblems of Type 1 are completely decoupled due to the 64 homogeneous initial values. The same is true for each subproblem of Type 2, the 65 exact solution of which can be computed as

$$w_j(t) = e^{(t-T_{j-1})A} v_{j-1}(T_{j-1})$$
(2)

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as soon as the initial value  $v_{j-1}(T_{j-1})$  is available. Therefore it is natural to assign the 67 integrations for  $v_{i-1}$  and  $w_i$  to the same processor so that there is no need for communication and synchronization between the two types of subproblems. Note that the time intervals  $[T_{i-1}, T]$  for the  $w_i$  are overlapping (see also Fig. 1). By superposition, 70 the solution of (1) is

$$u(t) = v_k(t) + \sum_{j=1}^k w_j(t)$$
 with  $k$  such that  $t \in [T_{k-1}, T_k]$ .

Only the computation of this sum requires communication between the processors. 73 Our parallel algorithm is given by simultaneously integrating the subproblems of 74 Type 1 and Type 2, and finally forming the sum for u(t) at the required time points t. 75

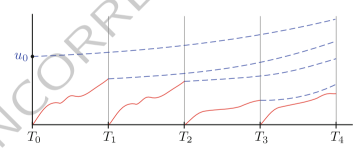


Fig. 1. Time-domain decomposition of an initial-value problem into inhomogeneous subproblems with zero initial value (Type 1, solid red curves) and overlapping homogeneous subproblems (Type 2, dashed blue curves). The solution is obtained as the sum of all curves

# 3 Computing the Matrix Exponential

The overlapping propagation of the linear homogeneous subproblems of Type 2 is 77 clearly redundant. To obtain an efficient parallel method, we require that the computation of the matrix exponentials in (2) is fast compared to the integration of the 79 subproblems of Type 1. 80

For scalar problems (N = 1) the computation of the exponential is a trivial task. 81 For computing the exponential of small to medium-sized dense matrices ( $N \lesssim 500$ ) 82 there are various methods available, see the review [5] and the monograph [4].

The computations become more challenging when the problem size N gets large, 84 in which case the matrix A should be sparse. Then one has to make use of the 85 fact that not the matrix exponential  $\exp(tA)$  itself is required, but only the prod- 86 uct  $\exp(tA)v_0$  with a vector  $v_0$ , by using a polynomial or rational Krylov method 87 (see [3] and the references therein). For brevity we will only describe a variant of 88 the restricted-denominator Arnoldi method described in [6] (see also [9]), which 89 extracts an approximation  $f_n(t) \approx \exp(tA)v_0$  from a Krylov space built with the 90 matrix  $S = (I - A/\sigma)^{-1}A$ ,

$$\mathcal{K}_n(S, v_0) = \text{span}\{v_0, Sv_0, \dots, S^{n-1}v_0\},$$
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the choice of the parameter  $\sigma \in (\mathbb{R} \cup \{\infty\}) \setminus (\Lambda(A) \cup \{0\})$  being dependent on the 93 spectral properties of A. For  $\sigma = \infty$  we obtain a standard Krylov space with the matrix A, i.e.,  $\mathcal{K}_n(S, v_0) = \mathcal{K}_n(A, v_0)$ . If  $\mathcal{K}_n(S, v_0)$  is of full dimension n, as we assume 95 in the following, we can compute an orthonormal basis  $V_n = [v_1, v_2, \dots, v_n]$  by using 96 the well-known Arnoldi orthogonalization process (see, e.g., [2, Sect. 9.3.5]). The 97 Arnoldi approximation of  $\exp(tA)v_0$  is then defined as

$$f_n(t) := V_n \exp(t (S_n^{-1} + I_n/\sigma)^{-1}) V_n^* v_0, \quad S_n := V_n^* S V_n.$$
 99

Provided that n is small, the computation of  $f_n(t)$  requires the evaluation of a  $n \times n_{-100}$ matrix function which is small compared to the original  $N \times N$  matrix exponential. 101 Moreover, the matrix  $S_n$  can be constructed without explicit projection from quantities computed in the Arnoldi process.

In Fig. 2 we show the error norm  $\|\exp(A)v_0 - f_n(1)\|_2$  of the Arnoldi approximations with parameters  $\sigma = \infty$  and  $\sigma = 40$  (a rather arbitrary choice) as a function 105 of n, for the matrices

$$A_1 = \operatorname{tridiag}(30, -40, 10) \in \mathbb{R}^{199 \times 199}, A_2 = \operatorname{tridiag}(60, -90, 30) \in \mathbb{R}^{299 \times 299}$$

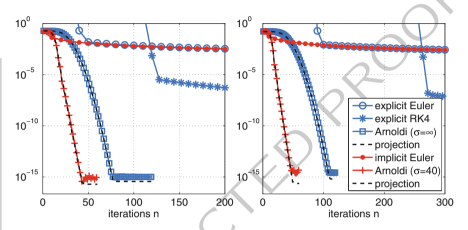
arising from the finite-difference discretization of the same 1D advection-diffusion 108 problem, and a random vector  $v_0$ . We have also plotted the error of orthogonal projection of the exact solution onto the space  $\mathcal{K}_n(S, v_0)$ , namely  $V_n V_n^* e^A v_0$ , and observe 110 that the Arnoldi method is capable of extracting an approximation nearby this projection. For comparison we show the error of the result produced by n steps of various 112 explicit and implicit integrators for the initial-value problem v' = Av,  $v(0) = v_0$ , integrated to t = 1. For this linear homogeneous problem all integrators actually compute 114 approximations from some Krylov space  $\mathcal{K}_n(S, v_0)$  (for the explicit integrators with 115 shift  $\sigma = \infty$  and for implicit Euler with  $\sigma = n$ ), but the Arnoldi methods extract much better approximations in the same number of iterations. Note also that the Arnoldi 117 method with finite shift  $\sigma = 40$  converges almost independently of the problem size 118 N, a property often referred to as mesh-independence.

Because the error of Arnoldi approximations decays usually very fast (i.e., 120  $\|e^{tA}v_0 - f_{n+1}(t)\|$  is considerably smaller than  $\|e^{tA}v_0 - f_n(t)\|$ , it is often sufficient 121

to use the difference of two consecutive iterates as an estimate for the approximation 122 error:

$$||e^{tA}v_0 - f_n(t)|| \le ||e^{tA}v_0 - f_{n+1}(t)|| + ||f_{n+1}(t) - f_n(t)||$$

$$\approx ||f_{n+1}(t) - f_n(t)||.$$
(3)



**Fig. 2.** Error (2-norm) of various time-stepping methods and Krylov methods for a linear homogeneous advection–diffusion problem v' = Av,  $v(0) = v_0$ , of size N = 199 (*left*) and N = 299 (*right*) as a function of time steps or Krylov space dimension n, respectively

# 4 Error Control and Parallel Efficiency

Many ODE solvers, for example those of MATLAB, use an error control criterion like 125

$$\|e(t)\|_{\infty} \leq \max\{\text{reltol} \cdot \|\widetilde{u}(t)\|_{\infty}, \text{abstol}\}, \quad t \in [0, T],$$

124

where  $e(t)=u(t)-\widetilde{u}(t)$  is the (estimated) error of the computed solution  $\widetilde{u}(t)$ . 127 Because the inhomogeneous subproblems of Type 1 for  $v_j(t)$  are solved with zero 128 initial guess, it is not advisable to use an error criterion which is relative to the norm 129 of the solution. Hence we assume that all of these subproblems are solved with an 130 absolute error  $\|e_j(t)\|_{\infty} \leq \text{abstol}/p$  over the time interval  $[T_{j-1},T_j]$ . This error is 131 then propagated exponentially over the remaining interval  $[T_j,T]$ , hence we have to 132 study the transient behavior of

$$||e^{tA}e_i(T_i)||_{\infty} \le ||e^{tA}||_{\infty} \operatorname{abstol}/p \tag{4}$$

for  $t \in [0, T - T_j]$ . It is well known that for a *stable* matrix A (i.e., all eigenvalues lie 134 in the left complex half-plane) the limit  $\lim_{t\to\infty} \|e^{tA}\|_{\infty}$  is finite. Unfortunately, the 135

norm may initially grow arbitrarily large before convergence sets in, a phenomenon 136 usually referred to as hump (see [5]). However, for a diagonally dominant matrix  $A = (a_{ij})$  with  $a_{ii} \leq 0$  this cannot happen, as one can show as follows (cf. [7]): Define  $\rho = \max_i \{a_{ii} + \sum_{j \neq i} |a_{ij}|\} \le 0$ . By the formula  $\exp(tA) = \lim_{k \to \infty} (I + tA/k)^k$ we have  $||e^{tA}||_{\infty} \leq \lim_{k \to \infty} ||I + tA/k||_{\infty}^{k}$ . For k sufficiently large we have 140

$$||I + tA/k||_{\infty} = \max_{i} \left\{ 1 + t \left( a_{ii} + \sum_{j \neq i} |a_{ij}| \right) / k \right\} = 1 + t\rho/k,$$
 141

hence

$$||e^{tA}||_{\infty} \le \lim_{k \to \infty} (1 + t\rho/k)^k = e^{t\rho} \le 1 \quad \text{for all } t \ge 0.$$

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Of course, it is possible to estimate the behavior of  $||e^{tA}||$  for general matrices and in other norms (see, e.g., [10]), but for brevity we will only consider a diagonally dominant A. In this case the errors  $e_i(t)$  of the subproblem solutions  $v_i(t)$  (j = 1, ..., p) 146 are non-increasing when being exponentially propagated, and if we assume that 147 the subproblems of Type 2 are solved exactly (or with sufficiently high accuracy), 148 then the overall error e(t) is bounded by the sum of subproblem errors (4), hence 149  $||e(t)||_{\infty} \leq$  abstol. If the integrator is a time-stepping method of order q, it is reasonable to assume that the computation time for one subproblem of Type 1 is at 151 most  $\tau_1(p) = (\tau_0 \cdot p^{1/q})/p$ , where  $\tau_0$  is the computation time for serial integration 152 over [0,T]. If each subproblem of Type 2 takes at most  $\tau_2$  units of computation time, 153 the expected efficiency of our parallel algorithm is at least

efficiency = 
$$\frac{\text{speedup}}{p} = \frac{1}{p} \cdot \frac{\tau_0}{\tau_1(p) + \tau_2} = \left(p^{1/q} + \frac{p \cdot \tau_2}{\tau_0}\right)^{-1}.$$
 (5)

The efficiency becomes large if the serial computation time  $\tau_0$  is long compared to 155  $p \cdot \tau_2$ , and if the integration order q is high. 156

# 5 Numerical Example

As a simple model problem we consider the 1D heat equation

$$\partial_t u(t,x) = \alpha \, \partial_{xx} u(t,x) + g(t,x)$$
 on  $x \in (0,1)$ ,  
 $u(t,0) = u(t,1) = 0$ ,  
 $u(0,x) = u_0(x) = 4x(1-x)$ ,  
 $g(t,x) = e \max\{1 - |c-x|/d,0\}$ , where  $c = .5 + (.5-d)\sin(2\pi ft)$ .

The source term g(t,x) is a hat function centered at c with half-width d=0.05 and 159 height  $e = 100 \cdot \alpha^{1/2}$ , oscillating with frequency f. Finite-difference discretization 160

This worst-case bound is sharp only if all errors  $e_j$  are collinear, which is rather unlikely. Probabilistic error estimation would give  $\|e(t)\|_{\infty} \lesssim \mathtt{abstol}/\sqrt{p}$ . This explains why the observed parallel efficiency of our algorithm is usually better than predicted by (5). We plan to investigate this in a sequel.

at N=100 points  $x_j=j/(N+1)$   $(j=1,\ldots,N)$  yields an initial-value problem (1), 161 where  $A = \alpha (N+1)^2$  tridiag $(1, -2, 1) \in \mathbb{R}^{N \times N}$ . This problem is integrated over the time interval [0, T = 1]. For the serial integration we have used the classical Runge-Kutta method of order q = 4 (implemented in MATLAB) with constant step size

$$h_0 = \min\{5 \cdot 10^{-5}/\alpha, 10^{-2}/f\},$$

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chosen to avoid instability of the time-stepping method caused by the stiff linear 166 term Au(t) and to capture the oscillations of g(t). As shown in Table 1, the absolute error ( $\infty$ -norm) is at most  $5 \cdot 10^{-4}$  for all diffusion coefficients  $\alpha = 0.01, 0.1, 1$  and 168 frequencies f = 1, 10, 100. These parameters determine the stiffness of Au(t) and 169 g(t), respectively. We have also tabulated the serial integration times  $\tau_0$ . As expected, 170 these are roughly proportional to  $h_0^{-1}$ .

For our parallel algorithm we have partitioned the interval [0, T] in p = 4 subintervals, and computed the solution u(t) at all time points  $T_j = jT/p$  (j = 1, ..., p). 173 The subproblems of Type 1 are integrated with step size  $h_1 = h_0 / \sqrt{p^{1/q}}$  (based on a 174 probabilistic error assumption, see the footnote on p. 6). In Table 1 we list the maximal computation time  $\tau_1$  for all subproblems of Type 1 among all processors.

For the subproblems of Type 2 we have used the Arnoldi method described in 177 Sect. 3 with shift  $\sigma = 5.3$ , in combination with the  $\infty$ -norm error estimate (3) for an 178 accuracy of  $10^{-4}$  (for more details on the selection of  $\sigma$  we refer to [9]). In Table 1 179 we list the maximal computation time  $\tau_2$  for all subproblems of Type 2 among all 180 processors.

The errors of the final solutions computed with our parallel algorithm are shown 182 in the second-last column, and they are all below the errors obtained by sequential 183 integration. This indicates that our choice for the step size  $h_1$  is reasonable. The parallel efficiency of our algorithm is above 50 % for all nine tests, and it increases with 185 frequency f because smaller time steps are required to integrate the inhomogene- 186 ity accurately. We finally note that for large-scale computations our algorithm could 187 also be used to further speed up a saturated space parallelization (e.g., by domain 188 decomposition).

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**Table 1.** Serial and parallel performance with p = 4 processors for a heat equation with diffusion coefficient  $\alpha$  and source-term frequency f.

α	f	serial			effi-	t1.1		
		$ au_0$	error	$ au_1$	$ au_2$	error	ciency	t1.2
0.01	1	4.97e-02	3.01e-04	1.58e-02	9.30e-03	2.17e-04	50 %	t1.3
0.01	10	2.43e-01	4.14e-04	7.27e-02	9.28e-03	1.94e-04	74 %	t1.4
0.01	100	2.43e+00	1.73e-04	7.19e-01	9.26e-03	5.68e-05	83 %	t1.5
0.1	1	4.85e-01	2.24e-05	1.45e-01	9.31e-03	5.34e-06	79 %	t1.6
0.1	10	4.86e-01	1.03e-04	1.45e-01	9.32e-03	9.68e-05	79 %	t1.7
0.1	100	2.42e+00	1.29e-04	7.21e-01	9.24e-03	7.66e-05	83 %	t1.8
1	1	4.86e+00	7.65e-08	1.45e+00	9.34e-03	1.78e-08	83 %	t1.9
1	10	4.85e+00	8.15e-06	1.45e+00	9.33e-03	5.40e-07	83 %	t1.10
1	100	4.85e+00	3.26e-05	1.44e+00	9.34e-03	2.02e-05	84 %	t1.11

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# Two-Grid LNKSz for Distributed Control of Unsteady Incompressible Flows

3

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Summary. The distributed control of unsteady incompressible flows has been the focus of 9 intense research in scientific computing in the past few years. Most of the existing approaches 10 for distributed control problems are based on the so-called reduced space method which is 11 easier to implement but may have convergence issues in some situations. In this paper we 12 investigate some fully coupled parallel two-grid Lagrange-Newton-Krylov-Schwarz (LNKSz) 13 algorithms for the implicit solution of distributed control problems. In the full space approach 14 we couple the control variables, the state variables and the adjoint variables in a single large 15 system of nonlinear equations. Numerical experiments are presented to show the efficiency 16 and scalability of the algorithm on supercomputers with more than one thousand processors. 17

1 Introduction 18

Flow optimal control problems have many important applications in science and engineering and many attempts have been made in the past few years to mathematically 20 understand and numerically solve flow control problems in various forms; see e.g., 21 [3, 6]. Popular approaches for solving unsteady flow control problems are explicit or 22 semi-implicit methods, both are limited by a Courant-Friedrichs-Lewy (CFL) condi- 23 tion. Recently, the class of full space Lagrange-Newton-Krylov-Schwarz (LNKSz) 24 algorithms was introduced for solving the steady state flow control problem [4, 5]. 25 The methods include two parts: a Lagrange-Newton method for the nonlinear sys- 26 tem obtained from the optimization problem and a Krylov subspace method for the 27 Jacobian system arising from the Newton method. In this paper we propose a class 28 of fully coupled parallel two-grid Lagrange-Newton-Krylov-Schwarz (LNKSz) alg-29 orithms for the distributed control of unsteady incompressible flows. Since we use a 30 fully implicit scheme, the CFL condition can be completely relaxed. We show numerically that the proposed LNKSz is stable and converges well with relatively large 32 times steps, and it is robust with respect to some of the physical parameters, such as 33 the Reynolds number.

The rest of the paper is organized as follows. In Sect. 2, we present the unsteady 35 distributed control problems and introduce a fully implicit discretization scheme. 36

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Section 3 includes the main components and features of LNKSz. Some numerical results are given in Sect. 4. We end the paper with some concluding remarks 38 in Sect. 5.

## 2 Mathematical Model and Discretization

We consider the two-dimensional unsteady incompressible Navier-Stokes equations 41 in the velocity-vorticity formulation:

$$\begin{cases}
-\Delta v_1 - \frac{\partial \omega}{\partial y} &= 0 \text{ in } [0, T] \times \Omega, \\
-\Delta v_2 + \frac{\partial \omega}{\partial x} &= 0 \text{ in } [0, T] \times \Omega, \\
\frac{\partial \omega}{\partial t} - \frac{1}{Re} \Delta \omega + v_1 \frac{\partial \omega}{\partial x} + v_2 \frac{\partial \omega}{\partial y} - \text{curl } \mathbf{f} = 0 \text{ in } [0, T] \times \Omega,
\end{cases}$$
(1)

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where  $\Omega$  is the computational domain and [0,T] is the time interval. In the above 43 equations the velocity field  $\mathbf{v}=(v_1,v_2)$  and the vorticity  $\omega$  are the state variables, 44  $\mathbf{f}=(f_1,f_2)$  is the external force, curl  $\mathbf{f}=-\partial f_1/\partial y+\partial f_2/\partial x$ , and Re is the Reynolds 45 number.

In the distributed control problem we try to find an external force  ${\bf f}$  over the 47 control domain  $\Omega_f\subseteq\Omega$  in order to achieve the goal 48

$$\min \mathscr{F}(\mathbf{v}, \boldsymbol{\omega}, \mathbf{f}) = \frac{1}{2} \int_0^T \mathscr{G}(\mathbf{v}, \boldsymbol{\omega}) dt + \frac{\gamma}{2} \int_0^T \int_{\Omega_f} ||\mathbf{f}||_2^2 d\Omega dt$$
 (2)

subject to the constraints (1) with some initial and boundary conditions. Here, 49  $\mathcal{G}(\mathbf{v}, \omega)$  is the objective function of the optimal control problem,  $\gamma > 0$  is a regularization parameter used to restrict the magnitude of the external force so that it is 51 not unrealistically large.

For solving unsteady distributed control problems, it typically requires a combination of a discretization in space and time with an optimization method. In this paper we follow the discretize-then-optimize approach with a finite difference method for the space discretization and a second-order backward differentiation formula for the time discretization. The original full-time-interval problem is too expensive to solve even on the latest supercomputers, we therefore replace it by a sequence of suboptimal problems, which are similar to the original problem but only defined on the time interval  $[t^{(k-1)}, t^{(k)}], k = 1, 2, \dots, k_{\max}$ , with  $t^{(0)} = 0$  and  $t^{(k_{\max})} = T$ . Let  $\mathbf{x} = (\mathbf{v}, \boldsymbol{\omega}, \mathbf{f})$ . 60 Then on each time interval we write the discrete suboptimization problem as follows:

$$\begin{cases} \min \mathcal{F}_h^{(k)}(\mathbf{x}) \\ \text{s.t. } \mathbf{C}_h^{(k)}(\mathbf{x}) = \mathbf{0}, \end{cases}$$
 (3)

where  $\mathscr{F}_h^{(k)}(\mathbf{x})$  is the restriction of  $\mathscr{F}$  on the interval  $[t^{(k-1)},t^{(k)}]$ , and  $\mathbf{C}_h^{(k)}(\mathbf{x})$  are the 62 constraints defined on the time interval  $[t^{(k-1)},t^{(k)}]$ .

By introducing the Lagrange multipliers  $\lambda$  with respect to the state and control 64 variables, we define the following Lagrangian functional 65

$$\mathcal{L}^{(k)}(\mathbf{x},\lambda) \equiv \mathcal{F}_h^{(k)}(\mathbf{x}) + (\lambda, \mathbf{C}_h^{(k)}(\mathbf{x})). \tag{4}$$

Let  $X \equiv (\mathbf{x}, \lambda)$ . Then, for  $k = 1, 2, ..., k_{\text{max}}$ , the KKT system obtained by differentiating (4) becomes

$$G^{(k)}(X) = \begin{pmatrix} \nabla_{\mathbf{x}} \mathcal{L}^{(k)}(\mathbf{x}, \lambda) \\ \nabla_{\lambda} \mathcal{L}^{(k)}(\mathbf{x}, \lambda) \end{pmatrix} = 0.$$
 (5)

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The optimality system (5) is a large, nonlinear, coupled, and muti-components 68 system. Moreover, the corresponding Jacobian matrix is indefinite and very ill-69 conditioned. Hence, a good preconditioner is essential to solve the optimality system 70 efficiently.

## 3 Two-Grid Newton Method and Schwarz Preconditioners

The class of full space LNKSz method includes the following steps: the Lagrangian 73 functional is formed and differentiated to obtain the KKT system; then the inexact 74 Newton method with line search is applied; and at each Newton iteration the linear 75 system is solved with a one-level or two-level Schwarz preconditioned Krylov sub-76 space method. We refer to LNKSz combined with the one-level (two-level) Schwarz 77 preconditioner as one-level (two-level) LNKSz method.

When using Newton's method to solve the nonlinear system (5) on a grid, one 79 of the major problems is the deterioration of the convergence rate when the grid is 80 refined, specially for the first time step, since in this case the initial guess is not good 81 enough for the Newton iterations. After many experiments, we find that a solution to 82 the problem is "grid-sequencing", which is quite effective in keeping the number of 83 nonlinear iterations small. In order to use grid-sequencing, we assume there are two 84 grids covering  $\Omega$ , a coarse grid of size H and a fine grid of size h. We first use the 85 one-level method to solve the nonlinear problem on the coarse grid with the initial guess obtained as a restriction of the fine grid solution from the previous timestep. 87 Of course, at the first time step, we choose the initial condition as the initial guess. 88 Then, we interpolate the solution to the fine grid and use it as an initial guess for the 89 nonlinear problem on the fine grid. We refer to this LNKSz method combined with 90 the grid-sequencing technique as the two-grid LNKSz method in which the same 91 coarse grid is also used to build the two-level Schwarz preconditioner for solving the 92 Jacobian problem.

We assume that  $\Omega$  is covered by a non-overlapping and an overlapping partition 94 as in [2]. Let J be the Jacobian matrix of the nonlinear problem (5) on the fine grid 95 and let  $R_i^{\delta}$  and  $R_i^{0}$  be the restriction operator from  $\Omega$  to its overlapping and non-overlapping subdomains, respectively. Here  $\delta$  is the size of the overlap. Then the 97 one-level restricted additive Schwarz (RAS) preconditioner [2] is defined as

$$M_{RAS}^{-1} = \sum_{i=1}^{N_p} (R_i^0)^T J_i^{-1} R_i^{\delta}.$$
 (6)

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with  $J_i = R_i^{\delta} J(R_i^{\delta})^T$  and  $N_p$  is the number of subdomains, which is the same as 99 the number of processors. Let  $J_c$  be the Jacobian matrix on the coarse grid and  $I_h^H$  a 100 restriction operator from the fine grid to the coarse grid. Then a multiplicative type two-level Schwarz preconditioner [8, 9] is defined as 102

$$M^{-1} = \left(I - (I - M_{RAS}^{-1}J)(I - M_c^{-1}J)(I - M_{RAS}^{-1}J)\right)J^{-1}$$
 (7)

with  $M_c^{-1} = (I_h^H)^T J_c^{-1} I_h^H$  and I is the identity matrix.

# **4 Numerical Experiments**

Our algorithms are implemented based on the Portable Extensible Toolkit for Scien- 105 tific computing (PETSc) [1]. All computations are performed on an IBM BlueGene/L 106 supercomputer.

In the following, we describe a backward-facing step flow control problem [7]. 108 Let  $\Omega = (0,6) \times (0,1)$ ,  $\Omega_f = (0,1) \times (0,0.5)$ , T = 1,  $\Gamma$  be the boundary of the domain  $\Omega$ ,  $\Gamma_2 = \{(x,y) \in \Gamma: 0 < y < 1, x = 6\}$ ,  $\Gamma_4 = \{(x,y) \in \Gamma: 0 < y < 1, x = 0\}$ , 110 and  $\Gamma_{4,a} = \{(x,y) \in \Gamma_4: 0.5 \le y < 1\}$ . Then the backward-facing step control problem 111 consists of finding  $(v_1, v_2, \omega, f_1, f_2)$  such that the minimization 112

$$\min \mathscr{F}(\boldsymbol{\omega}, \mathbf{f}) = \frac{1}{2} \int_0^T \int_{\Omega} \omega^2 \, d\Omega \, dt + \frac{\gamma}{2} \int_0^T \int_{\Omega_f} \|\mathbf{f}\|_2^2 \, d\Omega \, dt$$
 (8)

is achieved subject to the constraints (1) with the following boundary conditions: 113

$$\begin{cases}
v_{1} & = v_{in} \text{ on } [0, T] \times \Gamma_{4,a}, \\
v_{1} & = v_{out} \text{ on } [0, T] \times \Gamma_{2}, \\
v_{1} & = 0 \text{ on } [0, T] \times \Gamma_{u}, \\
v_{2} & = 0 \text{ on } [0, T] \times \Gamma, \\
\omega + \frac{\partial v_{1}}{\partial y} - \frac{\partial v_{2}}{\partial x} & = 0 \text{ on } [0, T] \times \Gamma, \\
\mathbf{v}(0, x, y) - \mathbf{v}_{0} & = \mathbf{0} \text{ in } \overline{\Omega}, \\
\omega(0, x, y) + \frac{\partial v_{0,1}}{\partial y} - \frac{\partial v_{0,2}}{\partial x} & = 0 \text{ in } \overline{\Omega},
\end{cases}$$
(9)

where  $\Gamma_u = \Gamma \setminus (\Gamma_{4,a} \cup \Gamma_2)$ . At the inflow boundary, a parabolic velocity profile  $v_{in} = 114$  $8(1-y)(y-\frac{1}{2})cos(t)$  is imposed. At the outflow boundary,  $v_{out}=y(1-y)cos(t)$  is 115 applied. The following initial velocity is defined by  $\mathbf{v}_0 = (v_{0.1}, v_{0.2})$  with 116

$$v_{0,1} = \begin{cases} y(1-y) + \frac{1}{16}y & \text{if } 0 \le y \le \frac{1}{2}, \\ y(1-y) + \frac{1}{16}(1-y) & \text{if } \frac{1}{2} \le y \le 1, \end{cases}$$

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and  $v_{0,2}(x,y) = 0$ . The parameter  $\gamma = 0.1$ .

In the experiments, we compare the following algorithms which are introduced 118 in Sect. 3:

- One-level LNKSz: one-level additive Schwarz is used as the Jacobian solve, and 120 inexact Newton is carried out on the fine grid;
- Two-level LNKSz: two-level multiplicative Schwarz is used as the Jacobian 122 solve, and inexact Newton is carried out on the fine grid;
- Two-grid LNKSz: two-level multiplicative Schwarz is used as the Jacobian solve on the fine grid, inexact Newton is used on the coarse grid to generate the initial 125 guess for the inexact Newton on the fine grid.

In all the experiments, all Jacobian matrices are constructed approximately using a 127 multi-colored finite difference method. The size of the coarse grid H is taken as 4h, 128 where h is the size of the fine grid. GMRES(90) and FGMRES(90) are used to solve  $^{129}$ the linear system at each Newton step on the coarse and the fine grids, respectively. 130 In the one-level method, the overlapping size is  $\delta = 6$ . In the two-level and two-grid 131 methods, the overlapping sizes of the coarse grid and the fine grids are  $\delta_c = 4$  and 132  $\delta=6$ , respectively. There are several nested iterative procedures in the proposed 133 algorithms, and each requires a proper stopping condition. We use  $10^{-10}$  ( $10^{-6}$ ) as 134 the absolute (relative) condition for all linear and nonlinear solves, except for the 135 linear coarse solve of the two-level preconditioner, for which we use  $10^{-4}$  ( $10^{-2}$ ) as 136 the absolute (relative) condition. The subdomain problems are solved with a sparse 137 LU factorization.

Next, we present results for the test problem and discuss some details of the 139 two-grid LNKSz. First, we compare the three methods in Table 1. Note that, the onelevel method doesn't converge when  $N_p = 1,024$ , which is caused by the divergence 141 of GMRES. Moreover, we note that: (1) for the linear solver, the number of GMRES 142 iterations for the one-level LNKSz is much larger than that for the two-level and twogrid methods; (2) for the nonlinear solver, the numbers of Newton iterations for the 144 one-level and two-level methods are also larger than that for the two-grid method; 145 and (3) compared with the one-level and two-level methods, the total computing time 146 for the two-grid method is much smaller. When the Reynolds number increases from 147 200 to 400, for one-level and two-level methods, the average number of Newton iter- 148 ations and the total computing time become larger. With the help of grid-sequencing, 149 the convergence of the two-grid method is less sensitive to the Reynolds number. 150 Based on the results of Table 1, it is clear that the two-grid method is better than the 151 others.

An important implementation detail to consider in designing two-grid LNKSz is 153 to balance the quality of the initial guess for the fine grid Newton iterations and the 154 computing time on the coarse solver. In Table 2, we present a comparison of the computing time for the two-level and two-grid methods. In this table, we report the total 156 time spent on the Newton iterations at some time steps, the time spent on the Newton 157 iterations on the coarse solver, and the percentage between these two computational 158 costs. We observe that the cost of Newton iterations on the coarse grid is very small 159 compared with the total computational cost. It is important to note that the coarse 160

**Table 1.** A comparison of three methods.  $768 \times 128$  grid, and  $\Delta t = 0.1$  (i.e., there are 10 time steps). "N<sub>p</sub>" stands for the number of processors which is the same as the number of subdomains, "IN" is the average number of inexact Newton iterations per time step on the fine grid, "RAS" is the average number of RAS preconditioned GMRES iterations per Newton iteration, and "Time" is the total computing time in seconds. "\*\*" means the divergence of GMRES.

$\overline{N_p}$	Method	IN	RAS	Time	IN	RAS	Time	
			Re=2	00		Re=400		
64	One-level	3.2	165.4	1370.4	3.7	158.9	1557.5	
64	Two-level	3.2	20.4	1342.8	3.7	19.2	1528.0	
64	Two-grid	2.1	18.7	898.2	2.0	18.0	836.4	
256	One-level	3.2	531.3	795.5	3.7	632.9	1052.3	
256	Two-level	3.2	27.4	479.9	3.7	27.1	560.1	
256	Two-grid	2.1	25.5	317.5	2.0	26.1	313.2	
1024	One-level		**			**		
1024	Two-level	3.2	66.3	314.3	3.7	67.9	376.9	
1024	Two-grid	2.1	64.2	208.5	2.0	68.5	209.8	

grid has to be sufficiently fine so that the coarse solution has a reasonable accuracy, 161 otherwise, it won't be able to provide a good initial guess for the fine grid nonlinear 162 solver.

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Table 2. A comparison of the computing time for the test problem at several different time steps.  $Re = 400, 768 \times 128$  grid, and  $\Delta t = 0.1$  (i.e., there are 10 time steps). The heading "Timestep(k)" represents the time step k, "Time" is the total time spent on the Newton iterations at the time step k, "Coarse\_time" is the time spent on the Newton iterations on the coarse solver at the time step k, and "Percent(%)" is ("Coarse\_time"/"Time").

$\overline{N_p}$	Timestep(k	Time				
>		Two-grid			Two-level	
64	k = 1	110.0	3.87	3.52%	458.9	
64	k = 2	80.0	2.39	2.99%	117.0	
64	k = 5	82.5	2.50	3.03%	118.0	
64	k = 10	84.7	2.51	2.96%	119.0	
256	k = 1	38.6	1.71	4.43%	172.8	
256	k = 2	29.7	0.99	3.33%	41.4	
256	k = 5	30.0	1.04	3.43%	41.6	
256	k = 10	30.8	1.06	3.44%	42.3	
1024	k = 1	23.3	1.37	5.88%	115.1	
1024	k = 2	20.6	0.68	3.30%	28.1	
1024	k = 5	21.2	0.72	3.39%	28.4	
1024	k = 10	21.5	0.74	3.44%	30.8	

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One of the difficulties in the nonlinear solver is the choice of the initial guess. 164 In Fig. 1, we show the nonlinear residual history by using three different methods at 165 the first time step (i.e., k = 1). One can see that the nonlinear system is difficult to 166 solve by using one-level or two-level method. In fact, it takes 11 iterations for the 167 one-level or two-level method to converge. By using the two-grid method only three 168 Newton iterations are required to satisfy the desired stopping condition.

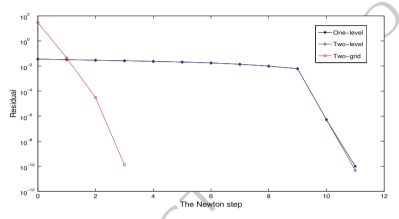


Fig. 1. Nonlinear residual history by using three different methods at the first time step, for  $Re = 200, 768 \times 128$  grid and 64 processors, and  $\Delta t = 0.1$ 

5 Conclusions 170

In this paper, we developed a family of two-grid algorithms for distributed control 171 of unsteady incompressible flows. With the help of the two-grid Newton method and 172 the two-level Schwarz preconditioner, we showed numerically that these strategies 173 provide substantial improvement of the overall method in terms of the total computing time, the number of linear iterations, and the number of Newton iterations, 175 especially when the number of processors is large.

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# On the Applicability of Lions' Energy Estimates in the <sup>2</sup> Analysis of Discrete Optimized Schwarz Methods with <sup>3</sup> Cross Points

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1 Introduction 8

For a bounded open subset  $\Omega \subset \mathbb{R}^2$ , suppose we want to solve

$$(\eta - \Delta)u = f$$
 on  $\Omega$ ,  $u = g$  on  $\partial \Omega$ , (1)

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for  $\eta \ge 0$  using the optimized Schwarz method (OSM)

$$(\eta - \Delta)u_i^k = f|_{\Omega_i} \text{ on } \Omega_i, \qquad u_i^k = g|_{\partial\Omega_i} \text{ on } \partial\Omega_i \cap \partial\Omega,$$

$$\frac{\partial u_i^k}{\partial n_i} + p_{ij}u_i^k = \frac{\partial u_j^{k-1}}{\partial n_i} + p_{ij}u_j^{k-1} \qquad \text{on } \Gamma_{ij} \text{ for all } \Gamma_{ij} \neq \emptyset,$$
(2)

for  $k=1,2,\ldots$  and  $i=1,\ldots,n$ , where  $\Omega_i\subset\Omega$  are non-overlapping subdomains, 11  $\Gamma_{ij}=\partial\Omega_i\cap\overline{\Omega_j}$  is the interface between  $\Omega_i$  and an adjacent subdomain  $\Omega_j,\ j\neq i$ , 12 and  $p_{ij}>0$  are Robin parameters along  $\Gamma_{ij}$ . In [7], the powerful technique of energy estimates is used to show convergence of (2) for  $\eta=0$  under very general 14 conditions. Similar techniques have been used to prove convergence results for other 15 types of equations, cf. [2] for the Helmholtz equation and [5] for the time-dependent 16 wave equation. While one often assumes that the proof carries over trivially to finite-element discretizations, it has been reported in the literature (cf. [8, 9]) that discrete 18 OSMs can diverge when the domain decomposition contains cross points, i.e., when 19 more than two subdomains share a common point. This is in apparent contradiction 20 to Lions' proof, and such difficulties contribute to the limited use of OSMs in practice. The goal of this paper is to explain why the presence of cross points makes 22 it possible for the discrete OSM to diverge despite the proof of convergence at the 23 continuous level, and why this difference in behavior is generally unavoidable.

The remainder of the paper proceeds as follows. In Sect. 2, we recall Lions' energy estimate argument. In Sect. 3, we explain why it is impossible to convert the 26 continuous energy estimate into a discrete one in a generic way, without sacrificing 27 continuity of the solutions across subdomain boundaries. In Sect. 4, we show two 28 modifications that preserve continuity of the discrete solutions, but both must be used 29

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with Krylov methods to avoid divergent iterations. Finally, we show in Sect. 5 30 that a Lions-type discrete estimate can only hold under very stringent conditions; 31 thus, continuous estimates generally do not predict the behavior of discrete OSMs. 32

# 2 Continuous Energy Estimates

We briefly recall the argument in [7] proving the convergence of (2). We assume  $p_{ij} = p_{ji}$  to be a positive function that is bounded away from zero and defined on  $\tau_{ij} = \tau_{ji}$ . To show that (2) converges for all initial guesses, we first write the error equations

$$(\eta - \Delta)e_i^k = 0$$
 on  $\Omega_i$ ,  $e_i^k = 0$  on  $\partial \Omega \cap \partial \Omega_i$ ,

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$$\frac{\partial e_i^k}{\partial n_i} + p_{ij}e_i^k = \frac{\partial e_j^{k-1}}{\partial n_i} + p_{ij}e_j^{k-1} \quad \text{on } \Gamma_{ij} \text{ for all } \Gamma_{ij} \neq \emptyset,$$
(3)

where  $e_i = u_i^k - u|_{\Omega_i}$  with u being the exact solution to (1). We then multiply the first sequation in (3) by  $e_i^k$  and integrate to get

$$0 = a_i(e_i^k, e_i^k) - \int_{\partial \Omega_i} e_i^k \frac{\partial e_i^k}{\partial n_i} = a_i(e_i^k, e_i^k) - \sum_{(i,j) \in E} \int_{\Gamma_{ij}} e_i^k \frac{\partial e_i^k}{\partial n_i},$$

where the last sum is over all pairs of subdomains (i, j) that share an interface, and  $a_i(u_i, v_i) = \int_{\Omega_i} (\nabla u \cdot \nabla v + \eta u v) dx$  is the energy bilinear form defined on subdomain  $\Omega_i$ , so that  $a_i(e_i^k, e_i^k) = \int_{\Omega_i} \eta |e_i^k|^2 + |\nabla e_i^k|^2 dx \ge 0$  is the energy of the error on subdomain  $\Omega_i$ . We now rewrite the product term as

$$e_i^k \frac{\partial e_i^k}{\partial n_i} = \frac{1}{4p_{ij}} \left[ \left( \frac{\partial e_i^k}{\partial n_i} + p_{ij} e_i^k \right)^2 - \left( -\frac{\partial e_i^k}{\partial n_i} + p_{ij} e_i^k \right)^2 \right] =: \left( T_{+ij}^k \right)^2 - \left( T_{-ij}^k \right)^2, \qquad ^{48}$$

where  $T_{\pm ij}^k = \frac{1}{\sqrt{4p_{ij}}} (\pm \frac{\partial e_i^k}{\partial n_i} + p_{ij}e_i^k)$ . Since  $\frac{\partial e_j^k}{\partial n_i} = -\frac{\partial e_j^k}{\partial n_j}$  on  $\Gamma_{ij}$ , the interface condition 50 in (3) can be written as  $T_{+ij}^k = T_{-ii}^{k-1}$ , which means

$$a_i(e_i^k, e_i^k) = \sum_{(i,j) \in E} \int_{\Gamma_{ij}} \left[ \left( T_{+ij}^k \right)^2 - \left( T_{-ij}^k \right)^2 \right] ds = \sum_{(i,j) \in E} \int_{\Gamma_{ij}} \left[ \left( T_{-ji}^{k-1} \right)^2 - \left( T_{-ij}^k \right)^2 \right] ds.$$
 52

Thus,

$$a_{i}(e_{i}^{k}, e_{i}^{k}) + \sum_{(i,j)\in E} \int_{\Gamma_{ij}} (T_{-ij}^{k})^{2} ds = \sum_{(i,j)\in E} \int_{\Gamma_{ij}} (T_{-ji}^{k-1})^{2} ds.$$
 (4) 54

If we sum (4) through all subdomains i, we get

$$\sum_{i=1}^{N} a_i(e_i^k, e_i^k) + \sum_{i=1}^{N} \sum_{(i,j) \in E} \int_{\Gamma_{ij}} \left( T_{-ij}^k \right)^2 ds = \sum_{i=1}^{N} \sum_{(i,j) \in E} \int_{\Gamma_{ij}} \left( T_{-ji}^{k-1} \right)^2 ds. \tag{5}$$

We can now sum (5) over k and simplify to get

$$\sum_{k=0}^{K} \sum_{i=1}^{N} a_i(e_i^k, e_i^k) + B^K = B^0, \tag{6}$$

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where  $B^k := \sum_{i=1}^N \sum_{(i,j) \in E} \int_{\Gamma_{ij}} \left(T_{-ij}^k\right)^2 ds \ge 0$ . Since  $B^K \ge 0$  and each  $a_i(e_i^k, e_i^k) \ge 0$ , 57 we see that  $\sum_{k=0}^K a_i(e_i^k, e_i^k) \le B^0$  for all i and all K; hence  $a_i(e_i^k, e_i^k) \to 0$  as  $k \to \infty$  58 for all i. This implies that  $\|e_i^k\|_{H^1(\Omega_i)} \to 0$  when  $\eta > 0$ , so  $u_i \to u|_{\Omega_i}$  in the  $H^1$  norm. 59 A similar argument holds for  $\eta = 0$ . Note that the possible presence of cross points 60 does not cause any difficulty in the proof, since they form a subset of measure zero 61 in  $\partial \Omega_i$  and thus do not contribute to the boundary terms when integrating by parts, 62

#### 3 Finite Element Discretization

We now try to mimic Lions' proof in the finite element case. The finite element 64 method uses the weak form of (2), i.e., we must multiply the PDE by a test function 65  $\phi$  and integrate by parts. The problem becomes 66

Find 
$$u_i \in V^h \subset H^1(\Omega_i)$$
 s.t. for all  $\phi \in W^h \subset H^1_0(\Omega) \cap H^1(\Omega_i)$ ,

$$\int_{\Omega_i} (\nabla \phi \cdot \nabla u_i^k + \eta \phi u_i^k) - \int_{\partial \Omega_i} \phi \frac{\partial u_i^k}{\partial n_i} = \int_{\Omega_i} \phi f. \tag{7}$$

We now suppose that  $\phi$  is a basis function corresponding to a degree of freedom 70 along  $\Gamma_{ij}$ , whose support does not contain any cross points, see Fig. 1a To obtain an 71 expression for  $\int_{\partial \Omega_i} \phi \frac{\partial u_i^k}{\partial n_i}$ , we multiply the interface condition by  $\phi$  and integrate to 72 get

$$\int_{\Gamma_{ij}} \phi\left(\frac{\partial u_i^k}{\partial n_i} + p u_i^k\right) = \int_{\Gamma_{ij}} \phi\left(\frac{\partial u_j^{k-1}}{\partial n_i} + p u_j^{k-1}\right). \tag{8}$$

Substituting into (7) gives

$$a_i(\phi, u_i^k) + \int_{\Gamma_{ii}} \phi p u_i^k - \int_{\Gamma_{ii}} \phi \frac{\partial u_j^{k-1}}{\partial n_i} = \int_{\Omega_i} \phi f + \int_{\Gamma_{ii}} \phi p u_j^{k-1}. \tag{9}$$

Thus, we are faced with the same problem of finding an expression for  $\int_{\Gamma_{ij}} \phi \frac{\partial u_i^{k-1}}{\partial n_i}$ . 76 Fortunately, we can use the weak form of the PDE from  $\Omega_j$ 

$$a_j(\phi, u_j^{k-1}) - \int_{\partial \Omega_j} \phi \frac{\partial u_j^{k-1}}{\partial n_j} = \int_{\Omega_j} \phi f.$$
 (10)

Since  $n_i = -n_j$  on  $\Gamma_{ij}$ , adding (9) and (10) and rearranging gives

$$a_{i}(\phi, u_{i}^{k}) + \int_{\Gamma_{ij}} \phi p u_{i}^{k} = \int_{\Omega_{i}} \phi f - a_{j}(\phi, u_{j}^{k-1}) + \int_{\Gamma_{ij}} \phi p u_{j}^{k-1}, \tag{11}$$

which is just the usual block-Jacobi splitting of the stiffness matrix along  $\Gamma_{ij}$ .

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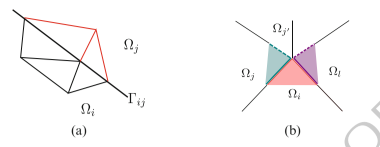


Fig. 1. Finite element discretization (a) without cross points and (b) with a cross point

Now assume that the support of  $\phi$  contains cross points, see Fig. 1b. Here  $\Omega_i$  is 81 adjacent to two distinct subdomains  $\Omega_j$  and  $\Omega_l$ ,  $j \neq l$ , and  $\phi$  is non-zero on all three 82 subdomains. Since the two parts of the interface,  $\Gamma_{ij}$  and  $\Gamma_{il}$ , must satisfy different 83 interface conditions, we must separate  $\int_{\partial\Omega_i}\phi\,\frac{\partial u_i^k}{\partial n}$  into contributions along  $\Gamma_{ij}$  and  $\Gamma_{il}$ , 84

$$a_i(\phi,u_i^k)-\int_{\Gamma_{ii}}\phi\,rac{\partial u_i^k}{\partial n_i}-\int_{\Gamma_{ii}}\phi\,rac{\partial u_i^k}{\partial n_i}=\int_{\Omega_i}\phi\,f.$$
 as

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The boundary term along  $\Gamma_{ij}$  can be replaced by the interface condition

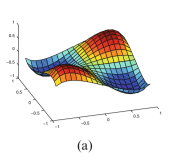
$$\int_{\Gamma_{ij}} \phi(\frac{\partial u_i^k}{\partial n_i} + pu_i^k) = \int_{\Gamma_{ij}} \phi(\frac{\partial u_j^{k-1}}{\partial n_i} + pu_j^{k-1}),$$
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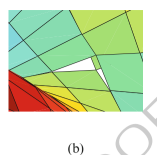
but now if we try to use the PDE on  $\Omega_j$  to eliminate the term  $\int_{\Gamma_{ij}} \phi \frac{\partial u_j^{k-1}}{\partial n_i}$ , we would 89 get get

 $\int_{\Gamma_{i}} \phi \frac{\partial u_{j}^{k-1}}{\partial n_{i}} = a_{j}(\phi, u_{j}^{k-1}) - \int_{\Gamma_{i}, j} \phi \frac{\partial u_{j}^{k-1}}{\partial n_{i}} - \int_{\Omega_{i}} \phi f,$ 

so we get a new term representing the trace along  $\Gamma_{ii'}$ , where  $\Omega_{i'}$  is another subdomain adjacent to j (see Fig. 1b). The same problem occurs when we try to eliminate 93 the trace along  $\Gamma_{il}$ . Note that, in the discrete FEM setting, the Robin traces are integrated along a subset of  $\partial \Omega_i$  of non-zero measure straddling both interfaces  $\Gamma_{ij}$  95 and  $\Gamma_{il}$ , and piecewise interface quantities are not available. Thus, the traces cannot be transmitted separately along  $\Gamma_{ij}$  and  $\Gamma_{il}$ , unlike in the continuous case; one must 97 introduce extra unknowns to represent the piecewise Robin traces (integrated against 98 a test function) for each subdomain at the cross point.

One way of circumventing the problem is to use mortar methods [1, 6], which 100 are designed for non-conforming grids. In these methods, the interface conditions 101 are imposed using mortar functions, which have one degree of freedom less at the 102 ends of intervals. Thus, there is no equation at the cross point, and the problem of 103 unavailable Robin traces goes away. However, since the interface conditions are only 104 enforced weakly, the method does not generally converge to the exact solution of the 105 global FEM problem, but rather to a discontinuous solution (Fig. 2) that is  $O(h^p)$ - 106 accurate, where p is the order of the finite element method.





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**Fig. 2.** (a) The solution of  $-\Delta u = f$  with four subdomains on  $\Omega = [-1, 1]^2$ , with right-hand side  $f(x, y) = \sin(xy)$ . The interface conditions are imposed using a mortar space. (b) Discontinuity of the composite solution near the origin

# 4 Two Lagrange Multiplier and Primal-Dual Methods

If we want to formulate subdomain problems that are equivalent to the discrete global 109 FEM problem, we need to introduce extra variables to represent the total Robin 110 traces. Thus, at the cross point, we impose for each  $\Omega_i$ 

$$a_i(\phi, u_i^k) + \int_{\partial \Omega_i} p\phi \cdot u_i^k + \lambda_i^k = \int_{\Omega_i} \phi f, \tag{12}$$

where  $\lambda_i^k$  are Lagrange multipliers for ensuring consistency with the global problem. 112 A cross point touching r subdomains requires r such Lagrange multipliers, so we also need r constraints to be satisfied at convergence: 114

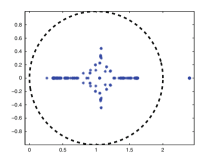
- Continuity constraints (r-1 equations): at the cross point, we must have  $u_1 = 115$  $u_2 = \cdots = u_r$ . 116
- PDE constraint (1 equation): if we sum (12) over the r subdomains and then 117 subtract the global PDE  $\sum_{i=1}^{r} a_i(\phi, u_i) = \int_{\Omega} \phi f$  from the result, we get 118

$$\sum_{i=1}^N \int_{\partial \Omega_i} p \phi u_i + \sum_{i=1}^N \lambda_i = 0.$$
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This gives two types of algorithms:

- 1. Primal-Dual methods: the continuity constraints are enforced for every iteration. 121 Thus, it suffices to introduce one extra variable (typically a coarse-grid basis 122 function that has the value one at the cross point), and the PDE constraint is 123 used as part of the coarse problem. This approach is similar to FETI-DP [3], 124 except it is usually formulated with Neumann rather than Robin traces.
- 2. Two-Lagrange Multiplier methods: the  $\lambda_i^k$  are retained, but the  $u_i^k$  are eliminated 126 using the PDE in the interior of the subdomains. This leads to a substructured 127 problem formulated on the interface, which is then solved using a preconditioned 128 Krylov method such as GMRES. This is known as the Two-Lagrange Multiplier 129 (2LM) method and has been studied in detail in [8].

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**Fig. 3.** Eigenvalues of the 2LM-preconditioned system for Poisson's equation ( $\eta = 0$ ), using a 4  $\times$  4 decomposition of the unit square with mesh size h = 1/64 and Robin parameter p = $C/\sqrt{h}$  for all interface nodes

Note that neither formulation is an exact discretization of (2) at cross points; thus, 131 Lions' convergence analysis does not apply there. In fact, one can show [4] that the 132 eigenvalues of the iteration matrix of the 2LM method may lie outside the unit disc 133 when cross points are present, as seen in the  $4 \times 4$  example shown in Fig. 3. In such 134 cases, the method diverges. However, convergence can be restored if one uses Robin 135 parameters with a different scaling at the cross points [4].

# **5** Conditions for Existence of Discrete Energy Estimates

To see what conditions are needed for Lions' estimates to hold in the discrete case, 138 let us consider solving  $-\Delta u = f$  on  $\Omega = [-1,1]^2$  using  $P^1$  finite elements on a 139 structured triangular mesh. This yields the system Au = f, where A is identical to the 140 matrix obtained from finite differences. If we now divide  $\Omega$  into four subdomains 141 corresponding to the four quadrants of the plane, then an optimized Schwarz method 142 must solve

143  $(A_i + L_i)u_i^k = g_i^k$  on each  $\Omega_i$ .

Here,  $A_i$  is the partially assembled stiffness matrix for  $\Omega_i$ ,  $L_i$  corresponds to transmission conditions, and  $g_i^k$  is a function of f and  $u_j^{k-1}$  for  $j \neq i$ . To define the *discrete* 146 error function, let us write  $u_i^* = u^*|_{\Omega_i}$ , where  $u^*$  is the exact solution to Au = f. Then 147 the error on  $\Omega_i$  is  $e_i^k = u_i^k - u_i^*$ , with discrete energy  $a_i(e_i^k, e_i^k) = (e_i^k)^T A_i e_i^k > 0$  when- 148 ever  $e_i^k \neq 0$ , since each subdomain touches a Dirichlet boundary. Now observe that 149

$$A_i e_i^k = A_i u_i^k - A_i u_i^* = A_i u_i^k - f_i$$
 at interior nodes.

Since the stencils of  $A_i$  and A coincide at interior nodes, we see that  $A_i e_i^k$  must be zero away from the interfaces. Thus, we in fact have 152

$$a_i(e_i^k,e_i^k) = \sum_{v \in \partial \Omega_i \backslash \partial \Omega} e_i^k(v) \cdot (A_i e_i^k)(v) = \sum_{v \in \partial \Omega_i \backslash \partial \Omega} [(T_{+i}^k(v))^2 - (T_{-i}^k(v))^2], \qquad \text{153}$$
 where  $T_{\pm i}^k(v)$  are the "Robin traces" at an interface point  $v$ :

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$$T_{+i}^k(v) = \frac{1}{\sqrt{4p}} \left[ (A_i e_i^k)(v) + p e_i^k(v) \right], \qquad T_{-i}^k(v) = \frac{1}{\sqrt{4p}} \left[ -(A_i e_i^k)(v) + p e_i^k(v) \right].$$
 156

Hence, if we let  $T_{+i}^k(v) = T_{-j}^{k-1}(v)$  at every point v on the interface, then the energy 157 estimate holds exactly the same way as in the continuous case, and we have convergence of the method. This allows us to deduce the correct interface conditions for v 160 away from the cross point. Using the definition  $e_i^k = u_i^k - u_i^*$ , we have

$$(A_i(u_i^k - u_i^*))(v) + p(u_i^k(v) - u_i^*(v)) = -(A_j(u_j^{k-1} - u_j^*))(v) + p(u_j^{k-1}(v) - u_j^*(v)).$$
(13)

But since

$$(A_i u_i^*)(v) + (A_i u_i^*)(v) = f(v), \tag{14}$$

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we can simplify (13) to get

$$(A_i u_i^k)(v) + p u_i^k(v) = f(v) - (A_j u_j^{k-1})(v) + p u_j^{k-1}(v).$$
 164

In other words, we need

(
$$L_i u_i^k$$
) $(v) = p u_i^k(v), \qquad g_i^k(v) = f(v) - (A_j u_j^{k-1})(v) + p u_j^{k-1}(v).$  166

On the other hand, if v is a cross point, then (14) is no longer valid, since f(v) is the sum of many subdomain contributions. Thus, it is in general impossible to find  $L_i$ and  $g_i^k$  such that the relation  $T_{-i}^k(v) = T_{-i}^{k-1}(v)$  holds at the cross point for some j. In our model problem, however, the stencil at the cross point has a special form for the 170 first and third quadrant:

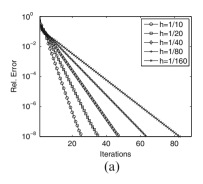
$$(A_1u_1^*)(0,0) = u^*(0,0) - \frac{1}{2}u^*(0,h) - \frac{1}{2}u^*(h,0),$$
  

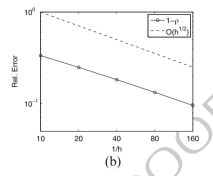
$$(A_3u_3^*)(0,0) = u^*(0,0) - \frac{1}{2}u^*(0,-h) - \frac{1}{2}u^*(-h,0).$$

Thus, we actually have  $(A_1u_1^*)(0,0) + (A_3u_3^*)(0,0) = \frac{1}{2}f(0,0)$ , a known quantity! A 172 similar relation holds between  $\Omega_2$  and  $\Omega_4$ , so it is actually possible to find transmission conditions at the cross point that satisfy the discrete energy estimate. For  $\Omega_1$ , 174 this reads

$$(A_1 u_1^k)(v) + p u_1^k(v) = \frac{1}{2} f(v) - (A_3 u_2^{k-1})(v) + p u_2^{k-1}(v).$$
 178

Figure 4 shows the convergence of the method for  $p = \frac{\pi}{2\sqrt{h}}$ , which gives the optimal 178 contraction factor  $\rho = 1 - O(\sqrt{h})$ , just as in the two-subdomain case. Since the discrete energy estimate holds, the converged subdomain solutions always coincide with 180 the exact discrete solution  $u^*$ , unlike in the mortar case. In general, discrete energy 181 estimates can only be derived if for every cross point v, its set of neighbors can be 182 partitioned into disjoint pairs (i, j) such that  $(A_i u_i^*)(v) + (A_j u_i^*)(v) = f_{ij}(v)$  can be 183 calculated without knowing u\*. For cross points with wide stencils or an odd number 184 of neighbors, this is not possible. In such cases, the methods in Sect. 4 are still excel- 185 lent choices in practice, but one cannot use Lions' estimates to deduce convergence 186 for arbitrary positive Robin parameters p.





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**Fig. 4.** (a) Convergence for different grid spacing h; (b) Contraction rate versus h

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# Non Shape Regular Domain Decompositions: An Analysis Using a Stable Decomposition in $H_0^1$

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Summary. In this paper, we establish the existence of a stable decomposition in the Sobolev 9 space  $H_0^1$  for domain decompositions which are not shape regular in the usual sense. In particular, we consider domain decompositions where the largest subdomain is significantly larger 11 than the smallest subdomain. We provide an explicit upper bound for the stable decomposition 12 that is independent of the ratio between the diameter of the largest and the smallest subdomain. 13

1 Introduction 14

One of the great success stories in domain decomposition methods is the invention 15 and analysis of the additive Schwarz method by Dryja and Widlund in [2]. Even 16 before the series of international conferences on domain decomposition methods 17 started, Dryja and Widlund presented a variant of the historical alternating Schwarz 18 method invented by Schwarz in [5] to prove the Dirichlet principle on general 19 domains. This variant, called the additive Schwarz method, has the advantage of 20 being symmetric for symmetric problems, and it also contains a coarse space compo- 21 nent. In a fully discrete analysis in [2], Dryja and Widlund proved, based on a stable 22 decomposition result for shape regular decompositions, that the condition number of 23 the preconditioned operator with a decomposition into many subdomains only grows 24 linearly as a function of  $\frac{H}{\delta}$ , where H is the subdomain diameter, and  $\delta$  is the over- 25 lap between subdomains. This analysis inspired a generation of numerical analysts, 26 who used these techniques in order to analyze many other domain decomposition 27 methods, see the reference books [4, 6, 7], or the monographs [1, 8], and references 28 therein.

The key assumption that the decomposition is shape regular is, however, often 30 not satisfied in practice: because of load balancing, highly refined subdomains are 31 often physically much smaller than subdomains containing less refined elements, 32 and it is therefore of interest to consider domain decompositions that are only 33 locally shape regular, i.e., domain decompositions where the largest subdomain can 34 be considerably larger than the smallest subdomain, and therefore the subdomain 35

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diameter and overlap parameters depend strongly on the subdomain index. In such 36 a domain decomposition, the generic ratio  $\frac{H}{\delta}$  from the classical convergence result 37 of the additive Schwarz method can be given at least two different meanings: let  $H_i$  38 refer to the diameter of subdomain number i and  $\delta_i$  refer to the width of the overlap around subdomain number i. Then in the classical convergence result from [2], 40 one could replace the generic ratio  $\frac{H}{\delta}$  by  $\frac{\max_i(H_i)}{\min_i(\delta_i)}$ , but this is likely to lead to a very 41 pessimistic estimate for the condition number growth. The general analysis of the 42 additive Schwarz method based on a shape regular decomposition does unfortunately 43 not permit to answer the question if the condition number growth for a locally shape 44 regular decomposition is in fact only linear in the quantity  $\max_i(\frac{H_i}{\delta_i})$ , which is much 45 smaller than  $\frac{\max_i(H_i)}{\min_i(\delta_i)}$  in the case of subdomains and overlaps of widely different sizes, 46 a case of great interest in applications.

In [3], we established the existence of a stable decomposition in the continuous 48 setting with an explicit upper bound and a quantitative definition of shape regular- 49 ity in two spatial dimensions. The explicit upper bound is also linear in the generic 50 quantity  $\frac{H}{\delta}$ , and the result is limited to shape regular domain decompositions where 51 all subdomains have similar size and where the overlap width is uniform over all 52 subdomains. Having explicit upper bounds, however, allows us now, using simi- 53 lar techniques, to establish the existence of a stable decomposition in the continuous setting with explicit upper bounds when  $\max_i(H_i) \gg \min_i(H_i)$ , and we provide 55 an explicit upper bound which is linear in  $\max_i(H_i/\delta_i)$  for problems in two spatial 56 dimensions. To get this result, only a few of the inequalities established in [3] need to 57 be reworked, and it would be very difficult to obtain such a result without the explicit 58 upper bounds from the continuous analysis in [3].

We state first in Sect. 2 our main theorem along with the assumptions we make on 60 the domain decomposition. We then prove the main theorem in Sect. 3 in two steps: 61 first, we show in Lemma 1 how to construct the fine component in Sect. 3.1, which 62 is an extension of the result [3, Theorem 4.6] for the case where subdomain sizes 63  $H_i$  and overlaps  $\delta_i$  can strongly depend on the subdomain index i. The major contribution is however in the second step, presented in Lemma 2 in Sect. 3.2, where we 65 show how to construct the coarse component in the case of strongly varying  $H_i$  and  $\delta_i$  66 between subdomains. This result is a substantial generalization of [3, Lemma 5.7]. 67 Using these two new results, and the remaining estimates from [3] which are still 68 valid, we can prove our main theorem. We finally summarize our results in the conclusions in Sect. 4.

#### 2 Geometric Parameters and Main Theorem

In the remainder of this paper, we always consider a domain decomposition that has 72 the following properties:

•  $\Omega$  is a bounded domain of  $\mathbb{R}^2$ .

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- The  $(U_i)_{1 \le i \le N}$  are a non-overlapping domain decomposition of  $\Omega$ , i.e., satisfy 75  $\bigcup_{i=1}^N \overline{U}_i = \overline{\Omega}$  and  $U_i \cap U_j = \emptyset$  when  $i \neq j$ . The  $U_i$  are bounded connected open 76 sets of  $\mathbb{R}^2$  and for all subdomains  $U_i$  the measure of  $\overline{U}_i \setminus U_i$  is zero.
- We set  $H_i := diam(U_i)$ .
- Two distinct subdomains  $U_i$  and  $U_j$  are said to be neighbors if  $\overline{U}_i \cap \overline{U}_j \neq \emptyset$ .
- For each subdomain  $U_i$ , let  $\delta_i > 0$  be such that  $2\delta_i \leq \min_{j,\overline{U}_i \cap \overline{U}_j = \emptyset} (\operatorname{dist}(U_i,U_j))$ . 80 We set  $\Omega_i := \{ \mathbf{x} \in \Omega, \operatorname{dist}(\mathbf{x}, U_i) < \delta_i \}$ . The  $\Omega_i$  form an overlapping domain 81 decomposition of  $\Omega$ . When subdomains  $U_i$  and  $U_j$  are neighbors, then the overlap between  $\Omega_i$  and  $\Omega_j$  is  $\delta_i + \delta_j$  wide. The intersection  $\Omega_i \cap \Omega_j$  is empty if and 83 only if the distance between  $U_i$  and  $U_j$  is positive.
- We set  $\delta_i^s = \min_{j \neq i, \overline{U}_i \cap \overline{U}_i \neq \emptyset} \delta_j$  and  $\delta_i^l = \max_{j \neq i, \overline{U}_i \cap \overline{U}_i \neq \emptyset} \delta_j$ .
- The domain decomposition has  $N_c$  colors: there exists a partition of  $\mathbb{N} \cap [1,N]$ into  $N_c$  sets  $I_k$  such that  $\Omega_i \cap \Omega_j$  is empty whenever  $i \neq j$  and i and j belong to 87 the same color  $I_k$ .
- $\mathcal{T}$  is a coarse triangular mesh of  $\Omega$ : one node  $x_i$  per subdomain  $\Omega_i$  (not counting 89 the nodes located on  $\partial\Omega$ ). By  $P_1(\mathcal{T})$ , we denote the standard finite element 90 space of continuous functions that are piecewise linear over each triangular cell 91 of  $\mathcal{T}$ .
- Let  $\theta_{min}$  be the minimum of all angles of mesh  $\mathcal{T}$ .
- No node (including the nodes located on  $\partial\Omega$ ) of the coarse mesh has more than 94 K neighbors. 95
- Let  $d_i$  be the length of the largest edge originating from node  $x_i$  in the mesh  $\mathcal{T}$ . 96
- Let  $H_{h,i}$  be the length of the shortest height through  $x_i$  of any triangle in the 97 coarse mesh  $\mathcal{T}$  that connects to  $\mathbf{x}_i$ . We also set  $H'_{h,i}$  as the minimum of  $H_{h,j}$  over 98 i and its direct neighbors in mesh  $\mathcal{I}$ .
- We suppose that for each subdomain  $U_i$ , there exists  $r_i > 0$  such that  $U_i$  is starshaped with respect to any point in the ball  $B(\mathbf{x}_i, r_i)$ . We also suppose  $r_i \leq \frac{H_{h,i}}{4K+1}$ and  $r_i \leq H'_{h,i}/2$ .
- We also assume the existence of both a pseudo normal  $X_i$  and of a pseudo curvature radius  $\tilde{R}_i$  for the domain  $U_i$ , i.e., we suppose that for each  $U_i$  there exists 104 an open layer  $L_i$  containing  $\partial U_i$ , a vector field  $\mathbf{X}_i$  continuous on  $L_i \cap \overline{U}_i$ ,  $\mathscr{C}^{\infty}$  on 105  $L_i \cap U_i$  such that  $D\mathbf{X}_i(\mathbf{x})(\mathbf{X}_i(\mathbf{x})) = 0$ ,  $\|\mathbf{X}_i(\mathbf{x})\| = 1$ , and  $\varepsilon_0 > 0$  such that for all 106 positive  $\varepsilon < \varepsilon_0$  and for all  $\hat{x}$  in  $\partial U_i$ ,  $\hat{x} + \varepsilon X_i(\hat{x}) \in U_i$  and  $\hat{x} - \varepsilon X_i(\hat{x}) \notin U_i$ . We set, 107 for all positive  $\delta'$ ,  $U_i^{\delta'} = \{ \mathbf{x} \in U_i, \operatorname{dist}(\mathbf{x}, \partial U_i) < \delta' \}$ , and  $V_i^{\delta'} = \{ \hat{\mathbf{x}} + s \mathbf{X}_i(\hat{\mathbf{x}}), \hat{\mathbf{x}} \in \mathcal{X}_i(\hat{\mathbf{x}}) \}$  $\partial U_i, 0 < s < \delta'$ . We assume there exist  $\hat{R}_i > 0$ ,  $\theta_{\mathbf{X}}$ ,  $0 < \theta_{\mathbf{X}} \leq \pi/2$ , and  $\delta_{0i}$ , 109  $0 < \delta_{0i} \le \hat{R}_i \sin \theta_{\mathbf{X}}$  such that  $V_i^{\hat{R}} \subset L_i \cap U_i$  and  $U_i^{\delta'} \subset V^{\delta'/\sin \theta_{\mathbf{X}}}$  for all positive  $\delta' \leq \delta_{0i}$ . Set  $\tilde{R}_i := 1/\|\text{div}\boldsymbol{X}_i\|_{\infty}$ . We suppose  $\delta_{0i} > \delta_i^l$ .

We finally define, for all i, the linear form on  $H_0^1(\Omega)$  by

$$\ell_i(u) := \frac{1}{\pi r_i^2} \int_{B(\boldsymbol{x}_i, r_i)} u(\boldsymbol{x}) d\boldsymbol{x} = \frac{1}{\pi} \int_{B(\boldsymbol{0}, 1)} u(\boldsymbol{x}_i + r_i \boldsymbol{y}) d\boldsymbol{y}.$$

We can now state our main theorem, namely the existence of a stable decomposition of  $H^1_0(\Omega)$  whose upper bound is independent of  $\frac{\max_i(H_i)}{\min_i(H_i)}$ . This theorem therefore leads to a substantially sharper condition number estimate in the important case 115 of an only locally shape regular decomposition, and is a major improvement of [3, 116 Theorem 5.12], which only considered shape regular decompositions, albeit at the 117 continuous level, in contrast to [2].

**Theorem 1.** For u in  $H_0^1(\Omega)$ , there exists a stable decomposition  $(u_i)_{0 \le i \le N}$  of u, i.e., 119  $u = \sum_{i=0}^N u_i$ ,  $u_0$  in  $P_1(\mathcal{T}) \cap H_0^1(\Omega)$  and  $u_i \in H_0^1(\Omega_i)$  such that

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$$\sum_{i=0}^{N} \|\nabla u_i\|_{L^2(\Omega_i)}^2 \le C \|\nabla u\|_{L^2(\Omega)}^2,$$

where  $C = 2C_1 + 2(1 + C_1)C_2$  and

$$\begin{split} C_1 &= \frac{1}{\tan \theta_{min}} \frac{\left(1 + 2 \max_i (\frac{r_i}{H_{h,i}})\right) K(\frac{25}{6\pi} \max_i (\frac{d_i}{r_i}) + 2\pi)}{1 - \left((2K+1) + (4K+1) \max_i (\frac{r_i}{H_{h,i}})\right) \max_i (\frac{r_i}{H_{h,i}})}, \\ C_2 &= 2 + 8\lambda_2^2 (N_c - 1)^2 (1 + \max_i \frac{\hat{R}_i}{\tilde{R}_i}) \max_i \frac{\delta_i^l}{\delta_i^s} \max_i \frac{\hat{R}_i}{\delta_i^s \sin \theta_{\mathbf{X}}} \\ &+ \frac{8}{3}\lambda_2^2 (N_c - 1)^2 (1 + \max_i \frac{\hat{R}_i}{\tilde{R}_i}) \max_i \frac{\delta_i^l}{\delta_i^s} \max_i \frac{r_i^2}{\delta_i^s \hat{R}_i \sin \theta_{\mathbf{X}}} \times \\ &\times \max_i \left( \left( \left( \frac{H_i^2}{r_i^2} + \frac{1}{2} \right)^{\frac{1}{4}} + \frac{H_i}{\sqrt[4]{2}r_i} \right)^4 - \frac{1}{2} - \frac{H_i^2}{r_i^2} - \frac{H_i^4}{2r_i^4} \right), \end{split}$$

with  $\lambda_2$  a universal constant depending only on the dimension, and being smaller 122 than 6 in the two dimensional case we consider here.

Note that the condition  $r_i \le \frac{H_{h,i}}{4K+1}$  implies that the denominator of  $C_1$  is positive. The value of  $C_2$  is also always positive.

# 3 Proof of Theorem 1

The proof is based on the continuous analysis in [3], but two results must be 127 adapted to the situation of only locally shape regular decompositions: we first show 128 in Sect. 3.1 how to construct the fine component, which is a technical extension of 129 the result [3, Theorem 4.6] for the case where subdomain sizes  $H_i$  and overlaps  $\delta_i$  can 130 strongly depend on the subdomain index i. Second, we explain in Sect. 3.2 the con-131 struction of the coarse component in the case of strongly varying  $H_i$  and  $\delta_i$  between 132 subdomains, which is a non-trivial generalization of [3, Lemma 5.7]. With these two 133 new results, and the remaining estimates from [3], the proof can be completed.

#### 3.1 Constructing the Fine Component

We begin by establishing a stable decomposition when there is no coarse mesh.

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**Lemma 1.** Let u be in  $H_0^1(\Omega)$ . Then, there exist  $(u_i)_{1\leq i\leq N}$ ,  $u_i$  in  $H_0^1(\Omega_i)$  such that  $u = \sum_{i=1}^{N} u_i$ , and

$$\sum_{i=1}^{N} \|\nabla u_{i}\|_{L^{2}(\Omega)}^{2} \leq 2 \|\nabla u\|_{L^{2}(\Omega)}^{2} + 8\lambda_{2}^{2} (N_{c} - 1)^{2} \left( \sum_{i=1}^{N} (1 + \frac{\hat{R}_{i}}{\tilde{R}_{i}}) \frac{\delta_{i}^{l}}{\delta_{i}^{s}} \frac{\hat{R}_{i}}{\delta_{i}^{s}} \|\nabla u\|_{L^{2}(U_{i})}^{2} \right) + 8\lambda_{2}^{2} (N_{c} - 1)^{2} \left( \sum_{i=1}^{N} (1 + \frac{\hat{R}_{i}}{\tilde{R}_{i}}) \frac{\delta_{i}^{l}}{\delta_{i}^{s}} \frac{1}{\delta_{i}^{s} \hat{R}_{i} \sin \theta_{\mathbf{X}}} \|u\|_{L^{2}(U_{i})}^{2} \right), \tag{1}$$

where  $\lambda_2$  is the universal constant of Theorem 1. We further have, for all  $\eta > 0$ , 139

$$\begin{split} \sum_{i=1}^{N} \|\nabla u_{i}\|_{L^{2}(\Omega)}^{2} &\leq 2 \|\nabla u\|_{L^{2}(\Omega)}^{2} + 8\lambda_{2}^{2}(N_{c} - 1)^{2} \sum_{i=1}^{N} (1 + \frac{\hat{R}_{i}}{\tilde{R}_{i}}) \frac{\delta_{i}^{l}}{\delta_{i}^{s}} \frac{\hat{R}_{i}}{\delta_{i}^{s}} \|\nabla u\|_{L^{2}(U_{i})}^{2} \\ &+ \frac{8(1 + \eta)}{3} \lambda_{2}^{2}(N_{c} - 1)^{2} \sum_{i=1}^{N} (1 + \frac{\hat{R}_{i}}{\tilde{R}_{i}}) \frac{\delta_{i}^{l}}{\delta_{i}^{s}} \frac{r_{i}^{2}}{\delta_{i}^{s} \hat{R}_{i} \sin \theta_{\mathbf{X}}} \times \\ &\times \left( \left( \left( \frac{H_{i}^{2}}{r_{i}^{2}} + \frac{1}{2} \right)^{\frac{1}{4}} + \frac{H_{i}}{\sqrt[4]{2}r_{i}} \right)^{4} - \frac{1}{2} - \frac{H_{i}^{2}}{r_{i}^{2}} - \frac{H_{i}^{4}}{2r_{i}^{4}} \right) \|\nabla u\|_{L^{2}(U_{i})}^{2} \\ &+ 8(1 + \frac{1}{\eta}) \pi \lambda_{2}^{2}(N_{c} - 1)^{2} \sum_{i=1}^{N} (1 + \frac{\hat{R}_{i}}{\tilde{R}_{i}}) \frac{\delta_{i}^{l}}{\delta_{i}^{s}} \frac{H_{i}^{2}}{\delta_{i}^{s} \hat{R}_{i} \sin \theta_{\mathbf{X}}} |\ell_{i}(u)|^{2}. \end{split}$$

*Proof.* We follow the proof of [3, Theorem 4.6]. Let  $\rho$  be a  $\mathscr{C}^{\infty}$  non-negative function whose support is included in the closed unit ball of  $\mathbb{R}^2$  and whose  $L^1$  norm 141 is 1. Let  $\rho_{\varepsilon}(\mathbf{x}) = \rho(\mathbf{x}/\varepsilon)/\varepsilon^2$  for all  $\varepsilon > 0$ . Let  $h_i$  be the characteristic function of the set  $\{x \in \mathbb{R}^2, \operatorname{dist}(x, U_i) < \delta_i/2\}$ . Let  $\phi_i = \rho_{\delta_i/2} * h_i$ . The function  $\phi_i$  is equal 143 to 1 inside  $U_i$ , vanishes outside of  $\{x \in \mathbb{R}^2, \operatorname{dist}(x, U_i) < \delta_i\}$ , and  $\|\nabla \phi_i\|_{L^{\infty}(\mathbb{R}^2)} \le$  $2\|\nabla\rho\|_{L^1(\mathbb{R}^2;(\mathbb{R}^2,\|\cdot\|_2))}/\delta_i. \text{ Here, } \|\nabla\rho\|_{L^1(\mathbb{R}^2;(\mathbb{R}^2,\|\cdot\|_2))} \text{ means } \int_{\mathbb{R}^2} \sqrt{\sum_{i=1}^2 |\partial_i\rho|^2} d\mathbf{x}.$ 145 For i in  $\mathbb{N} \cap [1, N]$ , let  $\psi_i = \phi_i \prod_{k=1}^{i-1} (1 - \phi_k)$ . We have  $0 \le \psi_i \le 1$ ,  $\psi_i$  zero in  $\Omega \setminus \Omega_i$ and  $\sum_{i} \psi_{i} = 1$  in  $\Omega$ . Set  $u_{i} = \psi_{i}u$ . The function  $u_{i}$  is in  $H_{0}^{1}(\Omega_{i})$  and  $u = \sum_{i} u_{i}$ . Following the proof of [3, Lemma 4.3], we get  $\sum_{i=1}^{N} \|\nabla \psi_{i}(\mathbf{x})\|_{2}^{2} \leq 2(N_{C}-1)\sum_{i=1}^{N} \|\nabla \phi_{i}(\mathbf{x})\|_{2}^{2}$ . Therefore, for all  $\boldsymbol{x}$  in  $\Omega$ ,

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$$\sum_{i=1}^{N} \|\nabla \psi_i(\mathbf{x})\|_2^2 \leq 8(N_c - 1) \|\nabla \rho\|_{L^1(\mathbb{R}^2; (\mathbb{R}^2, \|\cdot\|_2))}^2 \sum_{i=1}^{N} \frac{\mathbb{1}_{\Omega_i \setminus U_i}(\mathbf{x})}{\delta_i^2},$$

where  $\mathbb{1}_{\mathscr{O}}$  is the indicator function for the set  $\mathscr{O}$ . Since  $\sum_{i} \|\nabla u_{i}\|_{L^{2}(\Omega)}^{2} \leq 2 \|\nabla u\|_{L^{2}(\Omega)}^{2} +$  $2\int_{\Omega} |u(\mathbf{x})|^2 \sum_i |\nabla \psi_i(\mathbf{x})|^2 d\mathbf{x}$ , we get

$$\sum_{i=1}^{N} \|\nabla u_i\|_{L^2(\Omega)}^2 \leq 2 \|\nabla u\|_{L^2(\Omega)}^2 + 4\lambda_2^2 (N_c - 1)^2 \sum_{i=1}^{N} \int_{U_i} \mathbb{1}_{\{\text{dist}(\boldsymbol{x}, \partial U_i) < \delta_i^l\}} \frac{|u(\boldsymbol{x})|^2}{(\delta_i^s)^2} d\boldsymbol{x},$$

with  $\lambda_2 := 2\|\nabla \rho\|_{L^1(\mathbb{R}^2;(\mathbb{R}^2,\|\cdot\|_2))}$ . Using the  $W^{1,1}(\mathbb{R}^2)$  function  $\rho(\mathbf{x}) = 1 - \|\mathbf{x}\|_2$ , we obtain the estimate  $\lambda_2 = 6$ . To get (1), we apply Lemma 4.5 in [3] to each  $U_i$ , and to obtain (2), we apply Lemma 5.10 from the same reference.

To obtain a stable decomposition with a coarse component, we want to construct  $u_0$  in  $P_1(\mathcal{T})$  such that for all i,  $\ell_i(u_0) = \ell_i(u)$ .

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#### 3.2 Constructing the Coarse Component

To construct  $u_0$ , we follow the ideas of [3, Sect. 5.2]. First, we define a special norm. 155

**Definition 1.** Let  $\mathcal{T}$  be the coarse mesh of the domain  $\Omega$ . Let  $\mathcal{B}'$  be the set of indices 157 of the nodes of  $\mathcal{T}$  located on the boundary  $\partial \Omega$ . Let  $\partial \Omega$  be the set of the indices of 158 the nodes that are neighbors to the nodes with index in  $\partial \Omega$ . Let  $\partial \Omega$  be the set of pairs 159 of indices of neighboring nodes in  $\partial \Omega$  which are not on  $\partial \Omega$ . We define 160

$$\begin{aligned} \|\cdot\|_{\mathscr{V},\mathscr{B}} : \mathbb{R}^N &\to \mathbb{R}^+, \\ \mathbf{y} &\mapsto \sqrt{\sum_{(i,j) \in \mathscr{V}} |y_i - y_j|^2 + \sum_{i \in \mathscr{B}} |y_i|^2}. \end{aligned}$$

When u is in  $P_1(\mathcal{T}) \cap H_0^1(\Omega)$ , set  $\|u\|_{\mathscr{V},\mathscr{B}} := \|(u(\mathbf{x}_i))_{1 \le i \le N}\|_{\mathscr{V},\mathscr{B}}$ , where the  $\mathbf{x}_i$  are 161 the interior nodes of the mesh  $\mathscr{T}$ .

**Lemma 2.** For u in  $H_0^1(\Omega)$ , there exists  $u_0$  in  $P_1(\mathcal{T}) \cap H_0^1(\Omega)$  such that, for all i in 163  $\{1,\ldots,N\}$ ,  $\ell_i(u_0)=\ell_i(u)$  and

$$\|\nabla u_0\|_{L^2(\Omega)}^2 \leq \frac{1}{\tan \theta_{min}} \frac{\left(1 + 2\max_i(\frac{r_i}{H_{h,i}})\right) K\left(\frac{25}{6\pi} \max_i(\frac{d_i}{r_i}) + 2\pi\right)}{1 - \left((2K+1) + (4K+1)\max_i(\frac{r_i}{H_{h,i}})\right) \max_i(\frac{r_i}{H_{h,i}})}.$$

*Proof.* The results of [3, Lemmas 5.6 and 5.8] stand without modifications. Therefore  $u_0$  exists, and we have

$$\|\nabla u_0\|_{L^2(\Omega)}^2 \leq \frac{1}{\tan \theta_{min}} \frac{1 + 2\max_i(\frac{r_i}{H_{h,i}})}{1 - \left((2K+1) + (4K+1)\max_i(\frac{r_i}{H_{h,i}})\right)\max_i(\frac{r_i}{H_{h,i}})} \|u\|_{\mathscr{V},\mathscr{B}}^2.$$

Note that the condition  $r_i \le \frac{H_{h,i}}{4K+1}$  implies the second denominator in the above equation positive.

It remains to compare  $\|u\|_{\mathscr{V},\mathscr{B}}^2$  and  $\|\nabla u\|_{L^2(\Omega)}^2$ . We need to adapt the proof of [3, 169 Lemma 5.7]. We can suppose without any loss of generality that u is in  $\mathscr{C}^{\infty}(\overline{\Omega})$ . 170 Let i,j in  $\{1,\ldots,N\}$  be indices of neighboring nodes of  $\mathscr{T}$ . Let  $d_{ij}=x_i-x_j$ , and 171  $d_{ij}=\|d_{ij}\|$ . We have for all  $(i,j)\in\mathscr{V}$ 

<sup>&</sup>lt;sup>4</sup> Because of the homogenous Dirichlet condition on the boundary  $\partial\Omega$ , the nodes whose indices are in  $\mathscr{B}'$  are not associated to a degree of freedom, therefore  $\mathscr{B}'$  and  $\{1,\ldots,N\}$  have empty intersection.

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$$\begin{aligned} |\ell_{i}(u) - \ell_{j}(u)|^{2} &= \frac{1}{\pi^{2}} \left( \int_{B(\mathbf{0},1)} (u(\mathbf{x}_{i} + r_{i}\mathbf{y}) - u(\mathbf{x}_{j} + r_{j}\mathbf{y})) d\mathbf{y} \right)^{2} \\ &\leq \frac{1}{\pi} \int_{B(\mathbf{0},1)} \int_{0}^{1} \|\nabla u(t(\mathbf{x}_{i} + r_{i}\mathbf{y}) + (1-t)(\mathbf{x}_{j} + r_{j}\mathbf{y}))\|_{2}^{2} \|\mathbf{x}_{i} - \mathbf{x}_{j} + (r_{i} - r_{j})\mathbf{y}\|_{2}^{2} dt d\mathbf{y} \\ &\leq \frac{(d_{ij} + |r_{i} - r_{j}|)^{2}}{\pi} \int_{B(\mathbf{0},1)} \int_{0}^{1} \|\nabla u(t(\mathbf{x}_{i} + r_{i}\mathbf{y}) + (1-t)(\mathbf{x}_{j} + r_{j}\mathbf{y}))\|_{2}^{2} dt d\mathbf{y} \\ &\leq \frac{(d_{ij} + |r_{i} - r_{j}|)^{2}}{\pi} \int_{T_{i,j}} \|\nabla u(\mathbf{y}')\|_{2}^{2} \int_{0}^{1} \frac{1}{8} \frac{\|\mathbf{y}' - t\mathbf{x}_{i} - (1-t)\mathbf{x}_{j}\| \leq tr_{i} + (1-t)r_{j}}{(tr_{i} + (1-t)r_{j})^{2}} dt d\mathbf{y}', \end{aligned}$$

where the tube  $T_{i,j}$  is the convex hull of  $B(\mathbf{x}_i, r_i) \cup B(\mathbf{x}_i, r_i)$ . We get

$$\begin{aligned} \max_{\mathbf{y}' \in \mathbb{R}^2} \int_0^1 \frac{\mathbb{1}_{\{\|\mathbf{y}' - t\mathbf{x}_i - (1 - t)\mathbf{x}_j\| \le tr_i + (1 - t)r_j\}}}{(tr_i + (1 - t)r_j)^2} \mathrm{d}t \\ &= \max_{(s, s') \in \mathbb{R}^2} \int_0^1 \frac{\mathbb{1}_{\{\sqrt{(s - td_{ij})^2 + s'^2} \le tr_i + (1 - t)r_j\}}}{(tr_i + (1 - t)r_j)^2} \mathrm{d}t \\ &= \max_{s \in [-r_j, d_{ij} + r_i]} \int_0^1 \frac{\mathbb{1}_{\{|s - td_{ij}| \le tr_i + (1 - t)r_j\}}}{(tr_i + (1 - t)r_j)^2} \mathrm{d}t \\ &\le \max_{s \in [-r_j, d_{ij} + r_i]} \int_{\frac{s - r_j}{d_{ij} - (r_i - r_j)}}^{\frac{s + r_j}{d_{ij} - (r_i - r_j)}} \frac{1}{(tr_i + (1 - t)r_j)^2} \mathrm{d}t \\ &= \max_{s \in [-r_j, d_{ij} + r_i]} - \frac{1}{r_i - r_j} \left[ \frac{1}{(tr_i + (1 - t)r_j)} \right]_{\frac{s - r_j}{d_{ij} + (r_i - r_j)}}^{\frac{s + r_j}{d_{ij} + (r_i - r_j)}} \\ &= \max_{s \in [-r_j, d_{ij} + r_i]} \left( \frac{2}{d_{ij}r_j + s(r_i - r_j)} \right) \\ &= \frac{2}{\min(r_i, r_j)(d_{ij} - |r_i - r_j|)}. \end{aligned}$$

Since  $d_{i,i} \ge H_{h,i} \ge 4 \max(r_i, r_i)$ , we have

$$|\ell_i(u) - \ell_j(u)|^2 \le \frac{25d_{ij}}{6\pi \min(r_i, r_j)} \|\nabla u\|_{L^2(T_{ij})}^2.$$
(3)

If i is in the boundary set of the coarse mesh, then the node  $x_i$  is neighbor to a 175 node  $x_{i'}$  located on  $\partial \Omega$ . Note that i' lies outside of the range  $\{1,\ldots,N\}$ . Using [3, 176 Eqs. (5.7) and (5.9)], we get

$$\sum_{i \in \mathscr{B}} |\ell_i(u)|^2 \le \left(\sum_{i \in \mathscr{B}} \frac{4\|\boldsymbol{x}_i - \boldsymbol{x}_{i'}\|}{\pi r_i} \int_{T_i'} \|\nabla u(\boldsymbol{x})\|^2 d\boldsymbol{x}\right) + 2K\pi \|\nabla u\|_{L^2(\Omega)}^2, \tag{4}$$

where  $T_i'$  is the convex hull of  $B(\mathbf{x}_i, r_i) \cup B(\mathbf{x}_{i'}, r_i)$ . We sum inequality (3) over all i, j in the neighbor set and combine the resulting inequality with Eq. (4). Since

 $\max(r_i, r_j) \le H'_{h,i}/2 \le \min(H_{h,i}, H_{h,j}))/2$ , no point can belong to more than K tubes  $T_{i,j}$  or  $T_i'$ . Therefore,  $\|u\|_{\mathcal{V},\mathcal{B}}^2 \leq K(25 \max_i (d_i/r_i)/(6\pi) + 2\pi) \|\nabla u\|_{L^2(\Omega)}^2$ . This concludes the proof. AO1 To prove Theorem 1, we use Lemma 2 to construct the coarse component  $u_0$ . We then 178 apply Lemma 1 to  $u - u_0$  to get the fine components  $u_i$ . The terms in  $\ell_i(u)$  vanish. 4 Conclusion 180 We have proved the existence of a stable decomposition of the Sobolev space  $H_0^1(\Omega)$  181 in the presence of a coarse mesh when the domain decomposition is only guaran- 182 teed to be locally shape regular. We provided an explicit upper bound for the stable 183 decomposition that depends neither on  $\max_i(H_i)/\min_i(H_i)$ , nor on the number of 184 subdomains. This would not have been possible without the explicit upper bounds 185

sitions with local refinement.

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provided in [3]. This shows that deriving such explicit upper bounds can be important 186 for problems arising naturally in applications, e.g., load balanced domain decompo- 187

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AQ1. Please provide opening parenthesis in the sentence starting "Since  $\max(r_i, r_j) \le H'_{h,i}/2 \le \min(H_{h,i}, H_{h,j}))/2$ ."

# **Overlapping Domain Decomposition: Convergence Proofs**

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1 Introduction 7

During the last two decades many domain decomposition algorithms have been constructed and lot of techniques have been developed to prove the convergence of the algorithms at the continuous level. Among the techniques used to prove the convergence of classical Schwarz algorithms, the first technique is the maximum principle 11 used by Schwarz. Adopting this technique M. Gander and H. Zhao proved a conver- 12 gence result for n-dimensional linear heat equation in [4]. The second technique is 13 that of the orthogonal projections, used by P. L. Lions in [7], and his convergence 14 results are for linear Laplace equation and linear Stokes equation. In the same pa- 15 per, P. L. Lions also proved that the Schwarz sequences for linear elliptic equations 16 are related to classical minimization methods over product spaces and this technique 17 was then used by L. Badea in [1] for nonlinear monotone elliptic problems. Another 18 technique is the Fourier and Laplace transforms used in the papers [3, 5] for some 19 1-dimensional evolution equations, with constant coefficients. In [10, 11], S. H. Lui 20 used the idea of upper-lower solutions methods to study the convergence problem for 21 some PDEs, with initial guess to be an upper or lower solution of the equations and 22 monotone iterations. For nonoverlapping optimized Schwarz methods, P. L. Lions 23 in [8] proposed to use an energy estimate argument to study the convergence of the 24 algorithm. The energy estimate technique was then developed in [2] for Helmholtz 25 equation and it has then become a very powerful tool to study nonoverlapping prob- 26 lems. J.-H. Kimn in [6] proved the convergence of an overlapping optimized Schwarz 27 method for Poisson's equation with Robin boundary data and S. Loisel and D. B. 28 Szyld in [9] extended the technique of J.-H. Kimn to linear symmetric elliptic equa- 29 tion. Another technique is to use semiclassical analysis, which works for overlapping 30 optimized Schwarz methods with rectangle subdomains, linear advection diffusion 31 equations on the half plane (see [12]). This paper is devoted to the study of the convergence of Schwarz methods at the continuous level. We give a sketch of the proof 33 of the convergence of optimized Schwarz methods for semilinear parabolic equa- 34 tions, with multiple subdomains. Complete convergence proofs for both classical 35

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and optimized Schwarz methods, both semilinear parabolic and elliptic equations, 36 with multiple subdomains could be found in [13].

## 2 Convergence for Semilinear Parabolic Equations

Consider the following parabolic equation

$$\begin{cases} \frac{\partial u}{\partial t}(x,t) - \sum_{i,j=1}^{n} a_{i,j}(x) \frac{\partial^{2} u}{\partial x_{i} \partial x_{j}}(x,t) + \sum_{i=1}^{n} b_{i}(x) \frac{\partial u}{\partial x_{i}}(x,t) \\ + c(x)u(x,t) = F(x,t,u(x,t)), \text{ in } \Omega \times (0,\infty), \\ u(x,t) = g(x,t), \text{ on } \partial \Omega \times (0,\infty), \\ u(x,0) = g(x,0), \text{ on } \Omega, \end{cases}$$

$$(1)$$

where  $\Omega$  is a bounded and smooth enough domain in  $\mathbb{R}^n$ . The following conditions 40 are imposed on 1).

(A1) For all i, j in  $\{1, ..., I\}$ ,  $a_{i,j}(x) = a_{j,i}(x)$ . There exist strictly positive numbers 42  $\lambda$ ,  $\Lambda$  such that  $A = (a_{i,j}(x)) \ge \lambda I$  in the sense of symmetric positive definite matrices 43 and  $a_{i,j}(x) < \Lambda$  in  $\Omega$ . 44

(A2) The functions  $a_{i,j}$ ,  $b_i$ , c are in  $C^{\infty}(\mathbb{R}^n)$  and g is in  $C^{\infty}(\mathbb{R}^{n+1})$ .

(A3) There exists C > 0, such that  $\forall t \in \mathbb{R}, \forall x \in \mathbb{R}^n, |F(x,t,z) - F(x,t,z')| \le$ 46  $C|z-z'|, \forall z, z' \in \mathbb{R}$ . We now describe the way that we decompose the domain  $\Omega$ : 47 The domain  $\Omega$  is divided into I smooth overlapping subdomains  $\{\Omega_l\}_{l\in\{1,I\}}$ : 48

$$(\partial \Omega_l \backslash \partial \Omega) \cap (\partial \Omega_{l'} \backslash \partial \Omega) = , \quad \forall \quad l, l' \in \{1, \dots, l\}, \quad l \neq l';$$

$$\forall l \in \{1, \dots, l\}, \forall l', l'' \in J_l, l'' \neq l', \quad \Omega_{l'} \cap \Omega_{l''} = ,$$
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where

$$J_l = \{l' | \Omega_{l'} \cap \Omega_l \neq \};$$
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$$\cup_{l=1}^n \Omega_l = \Omega.$$
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This decomposition means that we do not consider cross-points in this paper. 56 Denote by  $\Gamma_{l,l'}$ , for  $l' \in J_l$ , the set  $(\partial \Omega_l \setminus \partial \Omega) \cap \overline{\Omega}_{l'}$ . The transmission operator  $\mathfrak{B}_{l,l'}$  57 is of Robin type  $\mathfrak{B}_{l,l'}v=\sum_{i,j=1}^n a_{i,j}\frac{\partial v}{\partial x_i}n_{l,l',j}+p_{l,l'}v$  and  $n_{l,l',j}$  is the j-th component 58 of the outward unit normal vector of  $\Gamma_{l,l'}$ ;  $p_{l,l'}$  is positive and belongs to  $L^{\infty}(\Gamma_{l,l'})$ . 59 The iterate #k in the l-th domain, denoted by  $u_l^k$  of the Schwarz waveform relaxation 60 algorithm is defined by: 61

$$\begin{cases} \frac{\partial u_l^k}{\partial t} - \sum_{i,j=1}^n a_{i,j} \frac{\partial^2 u_l^k}{\partial x_i \partial x_j} + \sum_{i=1}^n b_i \frac{\partial u_l^k}{\partial x_i} + c u_l^k = F(t,x,u_l^k), \text{ in } \Omega_l \times (0,\infty), \\ \mathfrak{B}_{l,l'} u_l^k = \mathfrak{B}_{l,l'} u_{l'}^{k-1}, \text{ on } \Gamma_{l,l'} \times (0,\infty), \forall l' \in J_l, \end{cases}$$
 (2)

where 62

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$$u_l^k(x,t)=g(x,t) \text{ on } (\partial\Omega_l\cap\partial\Omega)\times(0,\infty), \quad u_l^k(x,0)=g(x,0) \text{ in } \Omega_l.$$

The initial guess  $u^0$  is bounded in  $C^{\infty}(\overline{\Omega \times (0,\infty)})$ ; and at step 0, the Eq. (2) is solved 64 with boundary data 65

$$\mathfrak{B}_{l,l'}u_l^1(x,t) = u^0(x,t) \text{ on } \Gamma_{l,l'} \times (0,\infty), \forall l' \in J_l.$$

A compatibility condition on  $u^0(x,t)$  is also assumed

$$\mathfrak{B}_{l,l'}g(x,0) = u^0(x,0)$$
 on  $\Gamma_{l,l'}, \forall l' \in J_l$ .

By an induction argument, the algorithm is well-posed. Let  $e_l^k$  be  $u_l^k - u$ 67

$$\begin{cases}
\frac{\partial e_{l}^{k}}{\partial t} - \sum_{i,j=1}^{n} a_{i,j}(x) \frac{\partial^{2} e_{l}^{k}}{\partial x_{i} \partial x_{j}} + \sum_{i=1}^{n} b_{i}(x) \frac{\partial e_{l}^{k}}{\partial x_{i}} \\
+ c(x) e_{l}^{k} = F(t, x, u_{l}^{k}) - F(t, x, u), & \text{in } \Omega_{l} \times (0, \infty), \\
\mathfrak{B}_{l,l'} e_{l}^{k}(x, t) = \mathfrak{B}_{l,l'} e_{l'}^{k-1}(x, t), & \text{on } \Gamma_{l,l'} \times (0, \infty), \forall l' \in J_{l}.
\end{cases}$$
(3)

Moreover, 68

$$e_l^k(x,t)=0 ext{ on } (\partial\Omega_l\cap\partial\Omega) imes(0,\infty), \quad e_l^k(x,0)=0 ext{ in } \Omega_l.$$

For any function f in  $L^2(0,\infty)$ , define

$$\int_0^\infty f(x) \exp(-yx) dx.$$

For any fixed positive number  $\alpha$ , define

$$|f|_{\alpha} = \sup_{\alpha' > \alpha} \left[ \int_{\alpha'}^{\alpha' + 1} \left( \int_{0}^{\infty} f(x) \exp(-yx) dx \right)^{2} dy \right]^{\frac{1}{2}},$$

$$\mathbb{L}^{2}_{\alpha}(0,\infty) = \{ f : f \in L^{2}(0,\infty), |f|_{\alpha} < \infty \}.$$

Thus  $(\mathbb{L}^2_{\alpha}(0,\infty),|.|_{\alpha})$  is a normed subspace of  $L^2(0,\infty)$ .

**Theorem 1.** Consider the Schwarz algorithm with Robin transmission conditions 75 and the initial guess  $u^0$  in  $C_c^{\infty}(\overline{\Omega \times (0,\infty)})$ . There exists a constant  $\alpha$  large enough 76 such that 77

$$\lim_{k\to\infty}\sum_{l=1}^{I}\int_{\Omega_l}|e_l^k|_{\alpha}^2dx=0.$$

*Proof.* Let  $g_l$  be a function bounded and greater than 1 in  $C^{\infty}(\mathbb{R}^n,\mathbb{R})$ ,  $\alpha$  be a positive 78 constant, we define

$$\Phi_l^k(x) := \left( \int_0^\infty e_l^k \exp(-\alpha t) dt \right) g_l(x),$$

then  $\Phi_l^k(x)$  belongs to  $H^1(\Omega_l)$ . Let  $B_l^l$  and  $C^l$  be functions in  $L^{\infty}(\mathbb{R}^n)$  defined by

$$B_i^l := b_i + \sum_{j=1}^n \left( a_{i,j} \frac{\partial_j g_l}{g_l} \right),$$
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$$C^l = \left[\frac{\alpha}{2} + \sum_{i,j=1}^n \left(-a_{i,j} \frac{2\partial_i g_l \partial_j g_l}{(g_l)^2} - \partial_j a_{i,j} \frac{\partial_i g}{g} + a_{i,j} \frac{\partial_{i,j} g_l}{g_l}\right) - \sum_{i=1}^n b_i \frac{\partial_i g_l}{g_l}\right].$$
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Define

$$\begin{split} \mathfrak{L}_{lR}(\boldsymbol{\Phi}_{l}^{k}) &= -\sum_{i,j=1}^{n} \partial_{j}(a_{i,j}\partial_{i}\boldsymbol{\Phi}_{l}^{k}) + \sum_{i=1}^{n} B_{l}^{l}\partial_{i}\boldsymbol{\Phi}_{l}^{k} + C^{l}\boldsymbol{\Phi}_{l}^{k} \\ &+ \left\{ \int_{0}^{\infty} \left[ \left( \frac{\alpha}{2} + c \right) e_{l}^{k} - F(u_{l}^{k}) + F(u) \right] \exp(-\alpha t) dt \right\} g_{l}. \end{split}$$

It is possible to suppose  $\alpha$  to be large such that  $C^l$  belongs to  $(\frac{\alpha}{4}, \alpha)$ .

**Lemma 1.** Choose  $g_l$ ,  $g_{l'}$  such that  $\nabla g_l = \nabla g_{l'} = 0$  on  $\Gamma_{l,l'}$  and  $\frac{g_{l'}}{g_l} > 1$  on  $\Gamma_{l,l'}$ , for all 86 l' in  $J_l$ .  $\Phi_l^k$  is then a solution of the following equation 87

$$\begin{cases} \mathfrak{L}_{lR}(\boldsymbol{\Phi}_{l}^{k}) = 0, & \text{in } \Omega_{l} \times (0, \infty), \\ \beta_{l} \mathfrak{B}_{l,l'}(\boldsymbol{\Phi}_{l}^{k}) = \mathfrak{B}_{l,l'}(\boldsymbol{\Phi}_{l'}^{k-1}) & \text{on } \Gamma_{l,l'} \times (0, \infty), \forall l' \in J_{l}. \end{cases}$$

$$(4)$$

where  $eta_l = rac{g_{l'}}{g_l}$  on  $\Gamma_{l,l'}$ , for all l' in  $J_l$ .

For all l in  $\{1,I\}$ , denote by  $\tilde{\Omega}_l$  the open set  $\Omega_l \setminus \overline{\bigcup_{l' \in J_l} \Omega_{l'}}$ . For all l in I such that 89  $\varphi_l^{k+1} = \varphi_{l'}^k$  on  $\Gamma_{l,l'}$  for all l' in  $J_l$ , let  $\varphi_l^k$  and  $\varphi_l^{k+1}$  be functions in  $H^1(\tilde{\Omega}_l)$  and  $H^1(\Omega_l)$ . 90 Use the test functions  $\varphi_l^{k+1}$  and  $\varphi_l^k$ , and take the sum (with respect to l in  $\{1,I\}$ ) of 91  $\int_{\tilde{\Omega}_l} \mathfrak{L}_{lR}(\Phi_l^{k+1}) \varphi_l^{k+1}$  and  $\int_{\tilde{\Omega}_l} \mathfrak{L}_{lR}(\Phi_l^k) \varphi_l^k$  to get

$$-\sum_{l=1}^{I} \left\{ \int_{\tilde{\Omega}_{l}} C^{l} \Phi_{l}^{k} \varphi_{l}^{k} dx + \right.$$

$$+ \int_{\tilde{\Omega}_{l}} \sum_{i,j=1}^{n} a_{i,j} \partial_{i} \Phi_{l}^{k} \partial_{j} \varphi_{l}^{k} dx + \sum_{i=1}^{n} \int_{\tilde{\Omega}_{l}} B_{i}^{l} \partial_{i} \Phi_{l}^{k} \varphi_{l}^{k} dx - \sum_{l' \in J_{l}} \int_{\Gamma_{l',l}} p_{l',l} \Phi_{l}^{k} \varphi_{l}^{k} d\sigma$$

$$+ \int_{\tilde{\Omega}_{l}} \left\{ \int_{0}^{\infty} \left[ \left( \frac{\alpha}{2} + c \right) e_{l}^{k} - F(u_{l}^{k}) + F(u) \right] \exp(-\alpha t) dt \right\} g_{l} \varphi_{l}^{k} dx \right\}$$

$$= \sum_{l=1}^{I} \beta_{l} \left\{ \int_{\Omega_{l}} C^{l} \Phi_{l}^{k+1} \varphi_{l}^{k+1} dx + \sum_{l' \in J_{l}} \int_{\Gamma_{l,l'}} p_{l,l'} \Phi_{l}^{k+1} \varphi_{l}^{k+1} d\sigma$$

$$+ \int_{\Omega_{l}} \sum_{i,j=1}^{n} a_{i,j} \partial_{i} \Phi_{l}^{k+1} \partial_{j} \varphi_{l}^{k+1} dx + \sum_{l' \in J_{l}} \int_{\Gamma_{l,l'}} p_{l,l'} \Phi_{l}^{k+1} \varphi_{l}^{k+1} d\sigma$$

$$+ \int_{\Omega_{l}} \sum_{i=1}^{n} B_{i}^{l} \partial_{i} \Phi_{l}^{k+1} \varphi_{l}^{k+1} dx +$$

$$+ \int_{\Omega_{l}} \left\{ \int_{0}^{\infty} \left[ \left( \frac{\alpha}{2} + c \right) e_{l}^{k+1} - F(u_{l}^{k+1}) + F(u) \right] \exp(-\alpha t) dt \right\} g_{l} \varphi_{l}^{k+1} dx \right\}.$$

In (5), choose  $\varphi_l^{k+1}$  to be  $\Phi_l^{k+1}$ , then there exists  $\varphi_l^k$ , such that for all l' in  $J_l$   $\varphi_l^k = \varphi_{l'}^{k+1}$  so on  $\Gamma_{l,l'}$  and

$$||\varphi_l^k||_{H^1(\Omega_l)} \leq C \sum_{l' \in J_l} ||\varphi_{l'}^{k+1}||_{H^1(\Omega_{l'})} \text{ and } ||\varphi_l^k||_{L^2(\Omega_l)} \leq C \sum_{l' \in J_l} ||\varphi_{l'}^{k+1}||_{L^2(\Omega_{l'})}, \qquad \text{95}$$

where *C* is a positive constant.

The right hand side of (5) is then greater than or equal to

$$\sum_{l=1}^{I} \beta_{l} \left\{ \int_{\Omega_{l}} \lambda |\nabla \Phi_{l}^{k+1}|^{2} dx - \sum_{i=1}^{n} \int_{\Omega_{l}} ||B_{i}^{l}||_{L^{\infty}(\Omega_{l})} \left| \partial_{i} \Phi_{l}^{k+1} \right| |\Phi_{l}^{k+1}| dx \right\}.$$

$$\geq \sum_{l=1}^{I} \beta_{l} \left\{ \int_{\Omega_{l}} \frac{\lambda}{2} |\nabla \Phi_{l}^{k+1}|^{2} dx + \frac{\alpha}{8} \int_{\Omega_{l}} |\Phi_{l}^{k+1}|^{2} \right\}. \tag{6}$$

Similarly, the left hand side of (5) is less than or equal to

$$\sum_{l=1}^{I} \left\{ \int_{\tilde{\Omega}_{l}} \Lambda |\nabla \Phi_{l}^{k}| |\nabla \varphi_{l}^{k}| dx + \sum_{i=1}^{n} \int_{\tilde{\Omega}_{l}} ||B_{i}^{l}||_{L^{\infty}(\tilde{\Omega}_{l})} \left| \partial_{i} \Phi_{l}^{k} \right| |\varphi_{l}^{k}| dx \right. \\
\left. + \sum_{l' \in J_{l}} ||p_{l',l}||_{L^{\infty}(\Gamma_{l',l})} (||\Phi_{l}^{k}||_{H^{1}(\tilde{\Omega}_{l})}^{2} + ||\varphi_{l}^{k}||_{H^{1}(\tilde{\Omega}_{l})}^{2}) \right\} \\
\leq \sum_{l=1}^{I} M_{1} \left\{ \frac{1}{2} (||\nabla \Phi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2} + (\max_{i \in \{1,l\}} ||B_{i}^{l}||_{L^{\infty}(\tilde{\Omega}_{l})})^{2} ||\varphi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2}) \right. \\
\left. + \int_{\tilde{\Omega}_{l}} 2\alpha |\Phi_{l}^{k}| |\varphi_{l}^{k}| dx + \sum_{l' \in J_{l}} \int_{\Gamma_{l',l}} p_{l',l} |\Phi_{l}^{k}| |\varphi_{l}^{k}| d\sigma \right. \tag{7} \\
\left. + \Lambda \left( ||\nabla \Phi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2} + ||\nabla \varphi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2} \right) + \frac{\alpha}{2} ||\Phi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2} + \frac{\alpha}{2} ||\varphi_{l}^{k}||_{L^{2}(\tilde{\Omega}_{l})}^{2} \right\},$$

where  $M_1$  depends only on  $\{\Omega_l\}_{l\in\{1,I\}}$  and the Eq. (3). Choose  $\alpha$  such that  $\alpha>99$   $(\max_{i\in\{1,I\}}||B_i^l||_{L^\infty(\tilde{\Omega}_l)})^2$ , there exists  $M_2$  positive, depending only on  $\{\Omega_l\}_{l\in\{1,I\}}$  100 and (3) such that the right hand side of (7) is dominated by

$$\sum_{l=1}^{I} M_{2} \left\{ \int_{\tilde{\Omega}_{l}} \left( \frac{\lambda}{2} |\nabla \Phi_{l}^{k}|^{2} dx + \frac{\alpha}{8} |\Phi_{l}^{k}|^{2} + \frac{\lambda}{2} |\nabla \Phi_{l}^{k+1}|^{2} + \frac{\alpha}{8} |\Phi_{l}^{k+1}|^{2} \right) dx \right\}$$

$$\leq \sum_{l=1}^{I} M_{2} \left( \frac{\lambda}{2} ||\nabla \Phi_{l}^{k}||_{L^{2}(\Omega_{l})}^{2} + \frac{\alpha}{8} ||\Phi_{l}^{k}||_{L^{2}(\Omega_{l})}^{2} + \frac{\lambda}{2} ||\nabla \Phi_{l}^{k+1}||_{L^{2}(\Omega_{l})}^{2} + \frac{\alpha}{8} ||\Phi_{l}^{k+1}||_{L^{2}(\Omega_{l})}^{2} \right).$$
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Define 103

$$E_k := \sum_{l=1}^{I} \left( \frac{\lambda}{2} ||\nabla \Phi_l^k||_{L^2(\Omega_l)}^2 + \frac{\alpha}{8} ||\Phi_l^k||_{L^2(\Omega_l)}^2 \right), \tag{9}$$

then (6), (7), and (8) imply

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$$(\beta - M_2)E_{k+1} < M_2 E_k, \tag{10}$$

where  $\beta = \min\{\beta_1, \dots, \beta_I\}$ .

Since  $M_2$  depends only on  $\{\Omega_l\}_{l\in\{1,l\}}$  and (3),  $\beta$  can be chosen such that

$$M_3 := \frac{M_2}{\beta - M_2} < 1.$$

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We get

$$\begin{split} E_k &\leq M_3^k E_0 \\ &\leq M_3^k \sum_{l=1}^I \left( \frac{\lambda}{2} ||\nabla \Phi_l^0||_{L^2(\Omega_l)}^2 + \frac{\alpha}{8} ||\Phi_l^0||_{L^2(\Omega_l)}^2 \right). \end{split}$$

That deduces 109

$$||\Phi_l^k||_{L^2(\Omega_l)}^2 \le M_3^k \sum_{l=1}^I \left( \frac{4\lambda}{\alpha} ||\nabla \Phi_l^0||_{L^2(\Omega_l)}^2 + ||\Phi_l^0||_{L^2(\Omega_l)}^2 \right). \tag{11}$$

Since (11) still holds if  $M_3$  and  $\lambda$  are fixed, and  $\alpha$  is replaced by  $y > \alpha$ , then

$$\sum_{l=1}^{I} \int_{\Omega_{l}} \left( \int_{0}^{\infty} e_{l}^{k} \exp(-yt) dt g_{l} \right)^{2} dx \qquad (12)$$

$$\leq M_{3}^{k} \left[ \frac{4\lambda}{y} \sum_{l=1}^{I} \int_{\Omega_{l}} \left( \int_{0}^{\infty} |\nabla e_{l}^{0}| \exp(-yt) dt \right)^{2} g_{l}^{2} dx \right]$$

$$+ \frac{4\lambda}{y} \sum_{l=1}^{I} \int_{\Omega_{l}} \left( \int_{0}^{\infty} e_{l}^{0} \exp(-yt) dt \right)^{2} |\nabla g_{l}|^{2} dx$$

$$+ \sum_{l=1}^{I} \int_{\Omega_{l}} \left( \int_{0}^{\infty} e_{l}^{0} \exp(-yt) dt \right)^{2} g_{l}^{2} dx$$

Let  $\alpha'$  be a constant larger than or equal to  $\alpha$ , (12) implies

$$\sum_{l=1}^{I} \int_{\Omega_{l}} \int_{\alpha'}^{\alpha'+1} \left( \int_{0}^{\infty} e_{l}^{k} \exp(-yt) dt \right)^{2} g_{l}^{2} dy dx \tag{13}$$

$$\leq M_{3}^{k} \left[ \sum_{l=1}^{I} \int_{\Omega_{l}} \int_{\alpha'}^{\alpha'+1} \frac{4\lambda}{y} \left( \int_{0}^{\infty} |\nabla e_{l}^{0}| \exp(-yt) dt \right)^{2} g_{l}^{2} dy dx + \sum_{l=1}^{I} \int_{\Omega_{l}} \int_{\alpha'}^{\alpha'+1} \frac{4\lambda}{y} \left( \int_{0}^{\infty} e_{l}^{0} \exp(-yt) dt \right)^{2} |\nabla g_{l}|^{2} dy dx + \sum_{l=1}^{I} \int_{\Omega_{l}} \int_{\alpha'}^{\alpha'+1} \left( \int_{0}^{\infty} e_{l}^{0} \exp(-yt) dt \right)^{2} g_{l}^{2} dy dx \right].$$

Since  $u^0$  belongs to  $C_c^{\infty}(\overline{\Omega \times (0,\infty)})$ , the right hand side of (13) is bounded by a constant  $M_3^k M_4(\alpha)$ . The fact that  $g_l$  is greater than 1 implies

$$\sum_{l=1}^{I} \int_{\Omega_l} \int_{\alpha'}^{\alpha'+1} \left( \int_0^{\infty} e_l^k \exp(-yt) dt \right)^2 dy dx \le M_3^k M_4(\alpha). \tag{14}$$

Inequality (14) deduces

 $\lim_{k\to\infty}\sum_{l=1}^{I}\int_{\Omega_{l}}|e_{l}^{k}|_{\alpha}^{2}dx=0.$ (15) 114

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## FETI Methods for the Simulation of Biological Tissues 2

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**Summary.** In this paper we describe the application of finite element tearing and interconnecting methods for the simulation of biological tissues, as a particular application we consider the myocardium. As most other tissues, this material is characterized by anisotropic and 8 nonlinear behavior.

#### 1 Modeling Biological Tissues

In this paper we consider the numerical simulation of biological tissues, that can be described by the stationary equilibrium equations 12

$$\operatorname{div} \sigma(u, x) + f(x) = 0 \quad \text{for } x \in \Omega \subset \mathbb{R}^3, \tag{1}$$

to find a displacement field u where we have to incorporate boundary conditions to describe the displacements or the boundary stresses on  $\Gamma = \partial \Omega$ .

In the case of biological tissues the material is assumed to be hyperelastic, i.e. we 15 have to incorporate large deformations and a non-linear stress-strain relation. For the 16 derivation of the constitutive equation we introduce the strain energy function  $\Psi(C)$  17 which represents the elastic stored energy per unit reference volume. From this we 18 obtain the constitutive equation as in [1]

$$\sigma = J^{-1} \mathsf{F} \frac{\partial \Psi(\mathsf{C})}{\partial C} \mathsf{F}^{\top},$$

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where  $J = \det F$  is the Jacobian of the deformation gradient  $F = \nabla \varphi$ , and  $C = F^{\top} F$  is 21 the right Cauchy-Green tensor. In what follows we make use of the Rivlin-Ericksen 22 representation theorem to find a representation of the strain energy function  $\Psi$  in 23 terms of the principal invariants of  $C = F^{\top} F$ .

The cardiac muscle, the so-called *myocardium*, is the most significant layer for 25 the modeling of the elastic behavior of the heart wall. Muscle fibers are arranged in 26 parallel, in different sheets within the tissue. Although this fiber type is predominant, 27 we have also collagen that is arranged in a spatial network connecting the muscle 28

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fibers. We denote by  $\mathbf{f}_0$  the fiber axis which is referred to as the main direction of 29 the cardiac muscle fibers. The *sheet axis*  $\mathbf{s}_0$  is defined to be perpendicular to  $\mathbf{f}_0$  in 30 the plane of the layer. This direction coincides with the collagen fiber orientation. As 31 many other biological tissues we treat the myocardium as a nearly incompressible 32 material. It shows a highly nonlinear and, due to the muscle and collagen fibers, an 33 anisotropic behavior.

To capture the specifics of this fiber-reinforced composite, Holzapfel and Ogden proposed a strain-energy function  $\Psi$  that is decomposed into a volumetric, an 36 isotropic and an anisotropic part, which consists of a transversely isotropic and an 37 orthotropic response, see [7, 11],

$$\Psi(\mathsf{C}) = \Psi_{\text{vol}}(J) + \Psi_{\text{iso}}(\mathsf{C}) + \Psi_{\text{trans}}(\mathsf{C}, \mathbf{f}_0) + \Psi_{\text{trans}}(\mathsf{C}, \mathbf{s}_0) + \Psi_{\text{ortho}}(\mathsf{C}, \mathbf{f}_0, \mathbf{s}_0). \tag{2}$$

Following [11], we describe the volume changing part by

$$\Psi_{\text{vol}}(J) = \frac{\kappa}{2} (\log J)^2. \tag{3}$$

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The bulk modulus  $\kappa > 0$  serves as a penalty parameter to enforce the (almost) incompressibility constraint. To model the isotropic ground substance we use a classical 41 exponential model, see [2],

$$\Psi_{\rm iso}(\mathsf{C}) = \frac{a}{2b} \left\{ \exp[b(J^{-2/3}I_1 - 3)] - 1 \right\},\tag{4}$$

where a > 0 is a stress-like and b is a dimensionless material parameter.  $I_1 = tr(C)$  43 is the first principal invariant of the right Cauchy-Green tensor C. In (2),  $\Psi_{\text{trans}}$  is 44 associated with the deformations in direction of the fiber directions. Following [7] 45 we describe the transversely isotropic response by using

$$\Psi_{\text{trans}}(\mathsf{C}, \mathbf{f}_0) = \frac{a_f}{2b_f} \left\{ \exp[b_f (J^{-2/3} I_{4f} - 1)^2] - 1 \right\}$$

$$\Psi_{\text{trans}}(\mathsf{C}, \mathbf{s}_0) = \frac{a_s}{2b_s} \left\{ \exp[b_s (J^{-2/3} I_{4s} - 1)^2] - 1 \right\},$$
(5)

with the invariants  $I_{4f} := \mathbf{f}_0 \cdot (\mathsf{C}\mathbf{f}_0)$  and  $I_{4s} := \mathbf{s}_0 \cdot (\mathsf{C}\mathbf{s}_0)$  and the material parameters 47  $a_f$ ,  $b_f$ ,  $a_s$  and  $a_f$  which are all assumed to be positive. It is worth to mention, that 48 in this model the transversely isotropic responses  $\Psi_{\text{trans}}$  only contribute in the cases 49  $I_{4f} > 1$ ,  $I_{4s} > 1$ , respectively. This corresponds to a stretch in a fiber direction, and 50 this is explained by the wavy structure of the muscle and collagen fibers. In par- 51 ticular, the fibers are not able to support compressive stress. Moreover, the fibers 52 are not active at low pressure, and the material behaves isotropically in this case. 53 In contrast, at high pressure the collagen and muscle fibers straighten and then they 54 govern the resistance to stretch of the material. This behavior of biological tissues 55 was observed in experiments and this is fully covered by the myocardium model as 56 described above. The stiffening effect at higher pressure also motivates the use of the 57 exponential function in the anisotropic responses of the strain energy  $\Psi$ .

Finally a distinctive shear behavior motivates the inclusion of an orthotropic part 59 in the strain energy function in terms of the invariant  $I_{8fs} = \mathbf{f}_0 \cdot (C\mathbf{s}_0)$ 60

$$\Psi_{\text{ortho}}(\mathsf{C}) = \frac{a_{fs}}{2b_{fs}} \left\{ \exp(b_{fs}J^{-2/3}I_{8fs}^2) - 1 \right\},\tag{6}$$

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Here  $a_{fs} > 0$  is a stress-like and  $b_{fs} > 0$  a dimensionless material constant.

Note that the material parameters can be fitted to an experimentally observed 62 response of the biological tissue. In the case of the myocardium, experimental data 63 and, consequently, parameter sets are very rare. Following [7] and [11], we use the 64 slightly adapted material parameters to be found in Table 1.

$\kappa = 3333.33 \text{ kPa},$	a = 33.445  kPa,	b = 9.242 (-),
$a_f = 18.535 \text{ kPa},$	$b_s = 10.446$ (-),	$b_f = 15.972$ (-),
$a_{fs} = 0.417 \text{ kPa},$	$a_s = 2.564 \text{ kPa},$	$b_{fs} = 11.602$ (-).

**Table 1.** Material parameters used in the numerical experiments [7, 11].

Note that similar models can also be used for the description of other biological 66 materials, e.g., arteries, cf. [6, 8].

#### 2 Finite Element Approximation

In this section we consider the variational formulation of the equilibrium equations 69 (1) with Dirichlet boundary conditions  $u=g_D$  on  $\Gamma_D$ , Neumann boundary conditions 70  $t:=\sigma(u)n=g_N$  on  $\Gamma_N$ ,  $\Gamma=\overline{\Gamma}_D\cup\overline{\Gamma}_N$ ,  $\Gamma_D\cap\Gamma_N=\emptyset$ , and n is the exterior normal 71 vector of  $\Gamma=\partial\Omega$ . In particular we have to find  $u\in[H^1(\Omega)]^3$ ,  $u=g_D$  on  $\Gamma_D$ , such 72 that

$$a(u,v) := \int_{\Omega} \sigma(u) : e(v) dx = \int_{\Omega} f \cdot v dx + \int_{\Gamma_{N}} g_{N} \cdot v ds_{x} =: F(v)$$
 (7)

is satisfied for all  $v \in [H^1(\Omega)]^3$ , v = 0 on  $\Gamma_D$ .

By introducing an admissible decomposition of the computational domain  $\Omega$  into 75 tetrahedra and by using piecewise quadratic basis functions  $\varphi_{\ell}$ , the Galerkin finite el- 76 ement discretization of the variational formulation (7) results in a nonlinear system 77 of algebraic equations, to find  $u_h$  satisfying an approximate Dirichlet boundary con- 78 dition  $u_h = Q_h g_D$  on  $\Gamma_D$ , and

$$K_{\ell}(u_h) = \int_{\Omega} \sigma(u_h) : e(\varphi_{\ell}) dx = \int_{\Omega} f \cdot \varphi_{\ell} dx + \int_{\Gamma_N} g_N \cdot \varphi_{\ell} ds_x = F_{\ell}.$$
 (8)

For the solution of the nonlinear system (8), i.e. of  $G(u_h) := K(u_h) - F = 0$ , we apply 80 Newton's method to obtain the recursion 81

$$u_h^{k+1} = u_h^k + \Delta u_h^k$$
,  $\mathsf{G}_h'(u_h^k) \Delta u_h^k = -G(u_h^k)$ ,

or, by using the definition of  $G(\cdot)$ ,

$$u_h^{k+1} = u_h^k + \Delta u_h^k, \quad \mathsf{K}_h'(u_h^k) \Delta u_h^k = -K(u_h^k).$$
 (9)

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For the computation of the linearized stiffness matrix  $K'_h(u_h^k)$  we need to evaluate the 83 derivative of the nonlinear material model as described in the previous section. For a 84 detailed presentation how to compute  $K'_h(u_h^k)$  in this particular case, see [5].

#### 3 Finite Element Tearing and Interconnecting

For the parallel solution of (9) we will use a finite element tearing and interconnect- 87 ing approach [4], see also [8, 14] and references given therein. For a bounded domain 88  $\Omega \subset \mathbb{R}^3$  we introduce a non-overlapping domain decomposition

$$\overline{\Omega} = \bigcup_{i=1}^{p} \overline{\Omega}_{i} \quad \text{with } \Omega_{i} \cap \Omega_{j} = \emptyset \quad \text{for } i \neq j, \quad \Gamma_{i} = \partial \Omega_{i}.$$
 (10)

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The local interfaces are given by  $\Gamma_{ij} := \Gamma_i \cap \Gamma_j$  for all i < j. The skeleton of the domain 90 decomposition (10) is denoted as

$$\Gamma_C := \bigcup_{i=1}^p \Gamma_i = \Gamma \cup \bigcup_{i < j} \overline{\Gamma}_{ij}.$$

Instead of the global problem (1) we now consider local subproblems to find the local 92 restrictions  $u_i = u_{|\Omega_i}$  satisfying partial differential equations

$$\operatorname{div}(\sigma(u_i)) + f(x) = 0 \quad \text{for } x \in \Omega_i,$$

the Dirichlet and Neumann boundary conditions  $u_i = g_D$  on  $\Gamma_i \cap \Gamma_D$ ,  $\sigma(u_i) n_i = g_N$  on 95  $\Gamma_i \cap \Gamma_N$ , and the transmission conditions  $u_i = u_i$ ,  $t_i + t_j = 0$  on  $\Gamma_{ij}$ , where  $t_i = \sigma(u_i)n_i$  is 96 the local boundary stress, and  $n_i$  is the exterior normal vector of the local subdomain 97 boundary  $\Gamma_i = \partial \Omega_i$ . Note that the local stress tensors  $\sigma(u_i)$  are defined locally by 98 using the stress-strain function  $\Psi$  as introduced in Sect. 1, and by using localized 99 parameters  $\kappa, k_1, k_2, c$  and fiber directions  $\beta_1, \beta_2$ . Hence, by reordering the degrees 100 of freedom, the linearized system (9) can be written as

$$\begin{pmatrix} \mathsf{K}'_{11}(u^k_{1,h}) & \mathsf{K}'_{1C}(u^k_{1,h}) \mathsf{A}_1 \\ & \cdot & & \cdot \\ & \mathsf{K}'_{pp}(u^k_{p,h}) & \mathsf{K}'_{pC}(u^k_{p,h}) \mathsf{A}_p \\ \mathsf{A}_1^\top \mathsf{K}'_{C1}(u^k_{1,h}) \cdot \mathsf{A}_p^\top \mathsf{K}'_{Cp}(u^k_{p,h}) \sum_{i=1}^p \mathsf{A}_i^\top \mathsf{K}'_{CC}(u^k_{i,h}) \mathsf{A}_i \end{pmatrix} \begin{pmatrix} \Delta \mathbf{u}^k_{1,I} \\ & \cdot \\ \Delta \mathbf{u}^k_{p,I} \\ \Delta \mathbf{u}^k_C \end{pmatrix} = - \begin{pmatrix} \mathsf{K}_1(u^k_{1,h}) \\ & \cdot \\ & \mathsf{K}_p(u^k_{p,h}) \\ & \sum_{i=1}^p \mathsf{A}_i^\top \mathsf{K}_C(u^k_{i,h}) \end{pmatrix}, \quad \text{102}$$

where the increments  $\Delta \mathbf{u}_{i,l}^k$  correspond to the local degrees of freedom within the 103 subdomain  $\Omega_i$ , and  $\Delta \mathbf{u}_C^k$  is related to all global degrees of freedom on the coupling boundary  $\Gamma_C$ . By introducing the tearing 105

$$\mathbf{w}_{i} = \begin{pmatrix} \Delta \mathbf{u}_{i,I}^{k} \\ \mathsf{A}_{i} \Delta \mathbf{u}_{C}^{k} \end{pmatrix}, \; \mathsf{K}_{i}' = \begin{pmatrix} \mathsf{K}_{ii}'(u_{i,h}^{k}) & \mathsf{K}_{iC}'(u_{i,h}^{k}) \\ \mathsf{K}_{Ci}'(u_{i,h}^{k}) & \mathsf{K}_{CC}'(u_{i,h}^{k}) \end{pmatrix}, \; \mathbf{f}_{i} = -\begin{pmatrix} \mathsf{K}_{i}(u_{i,h}^{k}) \\ \mathsf{K}_{C}(u_{i,h}^{k}) \end{pmatrix},$$

by applying the interconnecting  $\sum_{i=1}^{p} B_i \mathbf{w}_i = \mathbf{0}$ , and by using discrete Lagrange multipliers, we finally have to solve the system

$$\begin{pmatrix} \mathsf{K}_1' & \mathsf{B}_1^\top \\ & \ddots & \vdots \\ & \mathsf{K}_p' \; \mathsf{B}_p^\top \\ \mathsf{B}_1 \dots \mathsf{B}_p \end{pmatrix} \begin{pmatrix} \mathbf{w}_1 \\ \vdots \\ \mathbf{w}_p \\ \lambda \end{pmatrix} = \begin{pmatrix} \mathbf{f}_1 \\ \vdots \\ \mathbf{f}_p \\ \mathbf{0} \end{pmatrix}. \tag{11}$$

For the solution of the linear system (11) we follow the standard approach of tearing and interconnecting methods. In the case of a floating subdomain  $\Omega_i$ , i.e.  $\Gamma_i \cap \Gamma_D = 110$   $\emptyset$ , the local matrices  $K_i'$  are not invertible. Hence we introduce the Moore-Penrose 111 pseudo inverse  $K_i^{\dagger}$  to represent the local solutions as

$$\mathbf{w}_i = \mathsf{K}_i^{\dagger} (\mathbf{f}_i - \mathsf{B}_i^{\top} \lambda) + \sum_{k=1}^{6} \gamma_{k,i} \mathbf{v}_{k,i}, \tag{12}$$

where  $\mathbf{v}_{k,i} \in \ker \mathsf{K}_i'$  correspond to the rigid body motions of elasticity. Note that in this case we also require the solvability conditions

$$(\mathbf{f}_i - \mathsf{B}_i^{\top} \lambda, \mathbf{v}_{k,i}) = 0$$
 for  $i = 1, \dots, 6$ .

In the case of a non-floating subdomain, i.e. ker  $K_i = \emptyset$ , we may set  $K_i^{\dagger} = K_i^{-1}$ . As in [10] we may also consider an all-floating approach where also Dirichlet boundary conditions are incorporated by using discrete Lagrange multipliers.

In general, we consider the Schur complement system of (11) to obtain

$$\sum_{i=1}^{p} \mathsf{B}_{i} \mathsf{K}_{i}^{\dagger} \mathsf{B}_{i}^{\top} \lambda - \sum_{i=1}^{p} \sum_{k=1}^{6} \gamma_{k,i} \mathsf{B}_{i} \mathbf{v}_{k,i} = \sum_{i=1}^{p} \mathsf{B}_{i} \mathsf{K}_{i}^{\dagger} \mathbf{f}_{i}, \quad (\mathbf{f}_{i} - \mathsf{B}_{i}^{\top} \lambda, \mathbf{v}_{k,i}) = 0,$$

which can be written as

$$\begin{pmatrix} \mathsf{F} & -\mathsf{G} \\ \mathsf{G}^{\top} \end{pmatrix} \begin{pmatrix} \lambda \\ \gamma \end{pmatrix} = \begin{pmatrix} \mathbf{d} \\ \mathbf{e} \end{pmatrix} \tag{13}$$

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with 122

$$\mathsf{F} = \sum_{i=1}^{p} \mathsf{B}_{i} \mathsf{K}_{i}^{\dagger} \mathsf{B}_{i}^{\top}, \ \mathsf{G} = \sum_{i=1}^{p} \sum_{k=1}^{6} \mathsf{B}_{i} \mathbf{v}_{k,i}, \ \mathbf{d} = \sum_{i=1}^{p} \mathsf{B}_{i} \mathsf{K}_{i}^{\dagger} \mathbf{f}_{i}, \ e_{k,i} = (\mathbf{f}_{i}, \mathbf{v}_{k,i}).$$
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For the solution of the linear system (13) we use the projection  $P^{\top} := I - G(G^{\top}G)^{-1}G^{\top}_{124}$  and it remains to consider the projected system

$$\mathsf{P}^{\top}\mathsf{F}\lambda = \mathsf{P}^{\top}\mathbf{d} \tag{14}$$

which can be solved by using a parallel GMRES method with suitable preconditioning. Note that the initial approximate solution  $\lambda^0$  satisfies the compatibility condition  $\mathsf{G}^\top \lambda^0 = \mathbf{e}$ . In a post processing we finally recover  $\gamma = (\mathsf{G}^\top \mathsf{G})^{-1} \mathsf{G}^\top (\mathsf{F} \lambda - \mathbf{d})$ , and subsequently the desired solution (12).

Following [3] we are going to apply either the lumped preconditioner

$$PM^{-1} := \sum_{i=1}^{p} B_i K_i' B_i^{\top}, \tag{15}$$

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or the Dirichlet preconditioner

$$\mathsf{P}\mathsf{M}^{-1} := \sum_{i=1}^{p} \mathsf{B}_{i} \begin{pmatrix} \mathsf{0} & \mathsf{0} \\ \mathsf{0} & \mathsf{S}_{i} \end{pmatrix} \mathsf{B}_{i}^{\top}, \tag{16}$$

where

$$\mathsf{S}_{i} = \mathsf{K}'_{CC}(u^{k}_{i,h}) - \mathsf{K}'_{Ci}(u^{k}_{i,h})\mathsf{K}'^{-1}_{ii}(u^{k}_{i,h})\mathsf{K}'_{iC}(u^{k}_{i,h})$$
132

is the Schur complement of the local finite element matrix  $K'_i$ . Alternatively, one may also used scaled hypersingular boundary integral operator preconditioner as proposed in [9].

#### **4 Numerical Results**

In this section we present some examples to show the applicability of the FETI ap- 138 proach for the simulation of the myocardium. We consider a mesh of the left and the 139 right ventricle of a rabbit heart with given fiber and sheet directions, see Fig. 1, which is decomposed in 480 subdomains, see Fig. 2. To describe the anisotropic and nonlinear cardiac tissue, we use the material model (2) with the parameters given in Table 1. 142 Dirichlet boundary conditions are imposed on the top of the myocardium mesh. The 143 interior wall of the right ventricle is exposed to the pressure of 1 mmHg which is 144 modeled with Neumann boundary conditions. Although this pressure is rather low, 145 the material model as used is orthotropic. To simulate a higher pressure, an appropri- 146 ate time stepping scheme has to be used. However, this does not affect the number of 147 local iterations significantly. The local Moore Penrose pseudo inverse matrices are 148 realized with a sparsity preserving regularization and the direct solver package Par- 149 diso [12, 13]. The global nonlinear finite element system with 12.188.296 degrees 150 of freedom is solved by a Newton scheme, where the FETI approach is used in each 151 Newton step. For this specific example the Newton scheme needed six iterations. 152 Due to the non-uniformity of the subdomains the efficiency of a global preconditioner becomes more important. We consider both the classical FETI approach, as 154 well as the all-floating formulation. Besides no preconditioning we use the simple 155 lumped preconditioner (15) and the Dirichlet preconditioner (16). It turns out that the number of iterations for the all-floating formulation is approximately half the number of iterations for the standard approach. Moreover, the Dirichlet preconditioner 158 within the all-floating formulation requires only 108 iterations, with a computing 159 time of approximately 5 min. All computations were done at the Vienna Scientific 160 Cluster (VSC2) (Fig. 3).

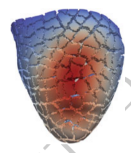
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#### FETI Methods for the Simulation of Biological Tissues

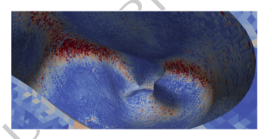


Fig. 1. Left and right ventricle of the rabbit heart. Mesh consists of 3.073.529 tetrahedrons and 547.680 vertices. Black lines indicate fiber directions  $\mathbf{f}_0$ . Point of view is from above showing the interior of the left and right ventricle

preconditioner iterations								
classical FETI								
none 941								
lumped, (15)	916							
Dirichlet, (16)	215							
all-floating FETI								
none	535							
lumped, (15)	401							
Dirichlet, (16)	108							



**Fig. 2.** The picture shows the displacement field of the rabbit heart with pressure applied in the *right* ventriculum. Point of view is from below showing the apex of the heart at the *bottom*. In the table the iteration numbers of the global GMRES method for different preconditioners are given



**Fig. 3.** Von Mises stress in the *right* ventricle. Point of view is from above looking inside the *right* ventricle

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# **Fast Summation Techniques for Sparse Shape** Functions in Tetrahedral hp-FEM

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**Summary.** This paper considers the hp-finite element discretization of an elliptic boundary 11 value problem using tetrahedral elements. The discretization uses a polynomial basis in which 12 the number of nonzero entries per row is bounded independently of the polynomial degree. 13 The authors present an algorithm which computes the nonzero entries of the stiffness matrix 14 in optimal complexity. The algorithm is based on sum factorization and makes use of the 15 nonzero pattern of the stiffness matrix.

1 Introduction 17

hp-finite element methods (hp-FEM), see e.g. [6, 9], have become very popular for 18 the approximation of solutions of boundary value problems with more regularity. In 19 order to obtain the approximate finite element solution numerically stable and fast, 20 the functions have to be chosen properly in hp-FEM. For quadrilateral and hexahe- 21 dral elements, tensor products of integrated Legendre polynomials are the prefered 22 basis functions, see [2]. For triangular and tetrahedral elements, the element can be 23 considered as collapsed quadrilateral or hexahedron. This allows us to use tensor 24 product functions. In order to obtain sparsity in the element matrices and a moder- 25 ate increase of the condition number, integrated Jacobi polynomials can be used, see 26 [3, 5, 7]. Then, it has been shown that the element stiffness and mass matrix have a 27 bounded number of nonzero entries per row, see [3–5] which results in a total number 28 of  $\mathcal{O}(p^d)$ , d=2,3, nonzero entries in two and three space dimensions, respectively. 29 However, the explicit computation of the nonzero entries is very involved.

This paper presents an algorithm which computes the element stiffness and mass 31 matrices in  $\mathcal{O}(p^3)$  operations in two and three space dimensions. The algorithm combines ideas based on sum factorization, [8], with the sparsity pattern of the matrices. 33 One other important ingredient is the fast evaluation of the Jacobi polynomials.

The outline of this paper is as follows. Section 2 defines  $H^1$ -conforming, i.e. 35 globally continuous piecewise polynomials, basis functions on the tetrahedron. The 36

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sum factorization algorithm is presented in Sect. 3. Section 4 is devoted to the evaluation of the Jacobi polynomials. The complexity of the algorithm is estimated in 38 Sect. 5.

#### 2 Definition of the Basis Functions

For the definition of our basis functions Jacobi polynomials are required. Let

$$p_n^{\alpha}(x) = \frac{1}{2^n n! (1-x)^{\alpha}} \frac{d^n}{dx^n} \left( (1-x)^{\alpha} (x^2 - 1)^n \right) \quad n \in \mathbb{N}_0, \ \alpha, \beta > -1$$
 (1)

be the *n*th Jacobi polynomial with respect to the weight function  $(1-x)^{\alpha}$ . The function  $p_n^{\alpha}$  is a polynomial of degree n, i.e.,  $p_n^{\alpha} \in \mathbb{P}_n((-1,1))$ , where  $\mathbb{P}_n(I)$  is the space 43 of all polynomials of degree n on the interval I. In the special case  $\alpha = 0$ , the functions  $p_n^0(x)$  are called Legendre polynomials. Moreover, let

$$\hat{p}_n^{\alpha}(x) = \int_{-1}^x p_{n-1}^{\alpha}(y) \, dy \quad n \ge 1, \quad \hat{p}_0^{\alpha}(x) = 1$$
 (2)

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be the *n*th integrated Jacobi polynomial. Several relations are known between the 46 different families of Jacobi polynomials, see e.g. [1]. In this paper, the relations

$$\begin{split} p_{n}^{\alpha-1}(x) &= \frac{1}{\alpha+2n} \left[ (\alpha+n) p_{n}^{\alpha}(x) - n p_{n-1}^{\alpha}(x) \right], \\ \hat{p}_{n+1}^{\alpha}(x) &= \frac{2n+\alpha-1}{(2n+2)(n+\alpha)(2n+\alpha-2)} \\ &\times \left( (2n+\alpha-2)(2n+\alpha)x + \alpha(\alpha-2) \right) \hat{p}_{n}^{\alpha}(x) \\ &- \frac{(n-1)(n+\alpha-2)(2n+\alpha)}{(n+1)(n+\alpha)(2n+\alpha-2)} \hat{p}_{n-1}^{\alpha}(x), \quad n \geq 1. \end{split} \tag{4}$$

are required.

Let  $\hat{\triangle}$  be the reference tetrahedron with the four vertices at (-1,-1,-1), 49 (1,-1,-1), (0,1,-1) and (0,0,1). On this element, the interior bubble functions 50

$$\phi_{ijk}(x, y, z) = u_i(x, y, z)v_{ij}(y, z)w_{ijk}(z), \quad i \ge 2, \ j, k \ge 1, i + j + k \le p$$
 (5)

are proposed for  $H^1$  elliptic problems in [3, (29)], where the auxiliary functions are

$$u_{i}(x,y,z) = \hat{p}_{i}^{0} \left(\frac{4x}{1-2y-z}\right) \left(\frac{1-2y-z}{4}\right)^{i},$$

$$v_{ij}(y,z) = \hat{p}_{j}^{2i-1} \left(\frac{2y}{1-z}\right) \left(\frac{1-z}{2}\right)^{j},$$

$$w_{ijk}(z) = \hat{p}_{k}^{2i+2j-2}(z).$$

In addition, there are vertex, face and edge based basis functions which can be 52 regarded as special cases of the above functions (5) for limiting cases of the indices 53 i, j and k, see [3] for more details. 54

Then, the element stiffness matrix for the Laplacian on the reference element  $\hat{\triangle}$  55 with respect to the interior bubbles reads as

$$\mathcal{K} = \left[ \int_{\hat{\triangle}} \nabla \phi_{ijk}(x, y, z) \cdot \nabla \phi_{i'j'k'}(x, y, z) \, \mathrm{d}(x, y, z) \right]_{i,j,k \le p,i'+j'+k' \le p}. \tag{6}$$

The transformation to the unit cube  $(-1,1)^3$  (Duffy trick) and the evaluation of the 57 nabla operation results in the integration of 21 different summands. More precisely, 58

$$\mathcal{K} = \sum_{m=1}^{21} \kappa_m \hat{\mathscr{J}}^{(m)}$$
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with known numbers  $\kappa_m \in \mathbb{R}$  and

$$\hat{\mathscr{J}}^{(m)} = \left[ \int_{-1}^{1} p_{x,1}(x) p_{x,2}(x) \, dx \right]$$

$$\times \int_{-1}^{1} \left( \frac{1-y}{2} \right)^{\gamma_{y}} p_{y,1}(y) p_{y,2}(y) \, dy$$

$$\times \int_{-1}^{1} \left( \frac{1-z}{2} \right)^{\gamma_{z}} p_{z,1}(z) p_{z,2}(z) \, dz \right]_{i+i+k < p:i'+i'+k' < p}.$$

The structure of the functions and coefficients is displayed in Table 1.

One summand is the term 62

$$\hat{\mathscr{J}}^{(6)} = \left( m_{ijk,i'j'k'} \right)_{i+j+k \le p,i'+j'+k' \le p} \tag{7}$$

which corresponds (before the Duffy trick) to

$$\begin{split} m_{ijk,i'j'k'} &= \int_{\hat{\triangle}} \hat{p}_i^0 \left( \frac{4x}{1 - 2y - z} \right) \hat{p}_{i'}^0 \left( \frac{4x}{1 - 2y - z} \right) \left( \frac{1 - 2y - z}{4} \right)^{i+i'} \\ &\times \hat{p}_j^{2i-1} \left( \frac{2y}{1 - z} \right) \hat{p}_{j'}^{2i'-1} \left( \frac{2y}{1 - z} \right) \left( \frac{1 - z}{2} \right)^{j+j'} \\ &\times p_{k-1}^{2i+2j-2}(z) p_{k'-1}^{2i'+2j'-2}(z) \, \mathrm{d}(x, y, z). \end{split}$$

The Duffy transformation applied to (7) gives

$$m_{ijk,i'j'k'} = \int_{-1}^{1} \hat{p}_{i}^{0}(x) \hat{p}_{i'}^{0}(x) dx \int_{-1}^{1} \left(\frac{1-y}{2}\right)^{i+i'+1} \hat{p}_{j'}^{2i'-1}(y) \hat{p}_{j}^{2i-1}(y) dy$$

$$\times \int_{-1}^{1} \left(\frac{1-z}{2}\right)^{i+j+i'+j'+2} p_{k-1}^{2i+2j-2}(z) p_{k'-1}^{2i'+2j'-2}(z) dz. \tag{8}$$

It has been shown in [3], this matrix has the sparsity pattern

$$m_{ijk,i'j'k'} = 0$$
 if  $(i, j, k, i', j', k') \in \mathfrak{S}_{ref}^{p}(ijk, i'j'k')$  (9)

	$p_{x,1}$	$p_{x,2}$	γ <sub>y</sub>	$p_{y,1}$	$p_{y,2}$	γz	$p_{z,1}$	$p_{z,2}$	t1.1
$\hat{\mathscr{J}}^{(1)}$	$p_{i-1}^0$	$p_{i'-1}^0$	i+i'-1	$\hat{p}_{i}^{2i-1}$	$\hat{p}_{i'}^{2i'-1}$	$\beta + \beta'$	$\hat{\mathbf{n}}^{-2+2\beta}$	$\hat{\beta}^{-2+2\beta'}$	t1.2
$\hat{\mathscr{J}}^{(2)}$	$\hat{p}_i^0$	$\hat{p}_{i'}^0$	i + i' + 1	$p_{j-1}^{2i-1}$	$p_{j'-1}^{2i'-1}$	$\beta + \beta'$	$\hat{p}_{L}^{-2+2\beta}$	$\hat{p}_{1}^{-2+2\beta'}$	t1.3
$\hat{\mathscr{J}}^{(3)}$	$p_{i-2}^{0}$	$\hat{p}^0_{i'}$	i+i'	$\hat{p}_{j}^{2i-1}$	$p_{j'-1}^{2i'-1}$	eta+eta'	$\hat{\mathbf{n}}^{-2+2\beta}$	$\hat{p}_{ij}^{-2+2p'}$	t1.4
$\hat{\mathscr{J}}^{(4)}$	$\hat{p}_i^0$	$p_{i'-2}^0$	i+i'	$p_{j-1}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta'$	$\hat{\boldsymbol{n}}^{-2+2\beta}$	$\hat{p}_{k'}^{-2+2\beta'}$	t1.5
$\hat{\mathscr{J}}^{(5)}$	$p_{i-2}^{0}$	$p_{i'-2}^0$	i + i' - 1	$\hat{p}_j^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$oldsymbol{eta}+oldsymbol{eta}'$	$\hat{p}_k^{-2+2\beta}$	$\hat{p}_{k'}^{-2+2\beta'}$	t1.6
Â(6)	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$\hat{p}_j^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta' + 2$	$p_{k-1}^{-2+2\beta}$	$p_{k'-1}^{-2+2\beta'}$	t1.7
$\hat{\mathscr{J}}^{(7)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-2}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta' + 1$	$\hat{p}_k^{-2+2\beta}$	$p_{k'-1}^{-2+2\beta'}$	t1.8
$\hat{\mathscr{J}}(8)$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-1}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta' + 1$	$\hat{p}_k^{-2+2\beta}$	$p_{k'-1}^{-2+2\beta'}$	t1.9
$\hat{\mathscr{J}}^{(9)}$	$p_{i-2}^{0}$	$\hat{p}^0_{i'}$	i+i'	$\hat{p}_j^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta' + 1$	$\hat{p}_k^{-2+2\beta}$	$p_{k'-1}^{-2+2\beta'}$	t1.10
$\hat{\mathscr{J}}^{(10)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$\hat{p}_j^{2i-1}$	$p_{j'-2}^{2i'-1}$	$\beta + \beta' + 1$	$p_{k-1}^{-2+2\beta}$	$\hat{p}_{k'}^{-2+2\beta'}$	t1.11
$\hat{\mathscr{J}}^{(11)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$\hat{p}_j^{2i-1}$	$p_{j'-1}^{2i'-1}$	$\beta + \beta' + 1$	$p_{k-1}^{-2+2\beta}$	$\hat{p}_{1}^{-2+2\beta'}$	t1.12
$\hat{\mathscr{J}}^{(12)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-2}^{2i-1}$	$p_{j'-2}^{2i'-1}$	$\beta + \beta'$	$\hat{p}_k^{-2+2\beta}$	$\hat{p}_{i,i}^{-2+2\beta'}$	t1.13
$\hat{\mathscr{J}}^{(13)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-1}^{2i-1}$	$p_{j'-2}^{2i'-1}$	$oldsymbol{eta}+oldsymbol{eta}'$	$\hat{p}_k^{-2+2\beta}$	$\hat{p}_{k'}^{-2+2\beta'}$	t1.14
$\hat{\mathscr{J}}^{(14)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-2}^{2i-1}$	$p_{j'-1}^{2i'-1}$	eta+eta'	$\hat{p}_k^{-2+2\beta}$	$\hat{\mathbf{n}}^{-2+2\beta'}$	t1.15
$\hat{\mathscr{J}}^{(15)}$	$\hat{p}_i^0$	$\hat{p}^0_{i'}$	i + i' + 1	$p_{j-1}^{2i-1}$	$p_{j'-1}^{2i'-1}$	eta+eta'	$\hat{p}_k^{-2+2\beta}$	$\hat{p}_{ij}^{-2+2\beta'}$	t1.16
$\hat{\mathscr{J}}^{(16)}$	$p_{i-2}^{0}$	$\hat{p}^0_{i'}$	i+i'	$\hat{p}_j^{2i-1}$	$p_{j'-2}^{2i'-1}$	eta+eta'	$\hat{p}_k^{-2+2\beta}$	$\hat{\boldsymbol{n}}^{-2+2\beta'}$	t1.17
$\hat{\mathscr{J}}^{(17)}$	$p_{i-2}^{0}$	$\hat{p}^0_{i'}$	i+i'	$\hat{p}_j^{2i-1}$	$p_{j'-1}^{2i'-1}$	eta+eta'	$\hat{p}_k^{-2+2\beta}$	$\hat{\mathbf{n}}^{-2+2\beta'}$	t1.18
$\hat{\mathscr{J}}^{(18)}$	$\hat{p}_i^0$	$p_{i'-2}^0$	i+i'	$\hat{p}_j^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta' + 1$	$p_{k-1}^{-2+2\beta}$	$\hat{p}_{ij}^{-2+2\beta'}$	t1.19
$\hat{\mathscr{J}}^{(19)}$	$\hat{p}_i^0$	$p_{i'-2}^0$	i+i'	$p_{j-2}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	eta+eta'	$\hat{p}_k^{-2+2\beta}$	$\hat{\mathbf{n}}^{-2+2\beta'}$	t1.20
$\hat{\mathscr{J}}^{(20)}$	$\hat{p}_i^0$	$p_{i'-2}^0$	i+i'	$p_{j-1}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	eta+eta'	$\hat{n}^{-2+2\beta}$	$\hat{p}_{ij}^{-2+2\beta'}$	t1.21
$\hat{\mathscr{J}}^{(21)}$	$p_{i-2}^{0}$	$p_{i'-2}^0$	i+i'-1	$\hat{p}_{j}^{2i-1}$	$\hat{p}_{j'}^{2i'-1}$	$\beta + \beta'$	$\hat{p}_k^{-2+2\beta}$	$\hat{p}_{k'}^{-2+2\beta'}$	t1.22

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**Table 1.** Integrands for  $\mathcal{K}$ , where  $\beta = i + j$ ,  $\beta' = i' + j'$ 

where 66

$$\mathfrak{S}^p_{ref}(ijk,i'j'k') = \{i+j+k \le p, i'+j'+k' \le p, |i-i'| \not\in \{0,2\} \\ \lor \quad |i-i'+j-j'| > 4 \qquad \lor \quad |i-i'+j-j'+k-k'| > 4\}$$

cf. [3, Theorem 3.3]. In the following the more general case

$$\mathfrak{S}^{p}(ijk,i'j'k') = \{i+j+k \le p,i'+j'+k' \le p,|i-i'| > 2$$

$$\vee |i-i'+j-j'| > 4 \qquad \vee |i-i'+j-j'+k-k'| > 4\}$$
(10)

is considered, e.g. the orthogonalities for |i - i'| = 1 are not assumed.

All 21 integrals give rise to a similar band structure as detailed above for  $\hat{\mathcal{J}}^{(6)}$ and can thus be treated in the same way as explained below for the particular case 70

m	$\kappa_m$
1,6,9,19	1
5,21	$\frac{5}{4}$
4,8,20	$c_1(i,j)$
7,19	$c_2(i,j)$
3,11,17	$c_1(i',j')$
2,15	$c_1(i,j)c_1(i',j')$
13	$c_1(i,j)c_2(i',j')$
10,16	$c_2(i',j')$
14	$c_1(i',j')c_2(i,j)$
21	$c_2(i,j)c_2(i',j')$

**Table 2.** Coefficients  $\kappa_m$  for  $\mathcal{H}$ , where  $c_1(i,j) = -\frac{1}{2} \frac{2i-1}{2i+2j-3}$  and  $c_2(i,j) = \frac{j-1}{2i+2j-3}$ .

of  $\hat{\mathscr{J}}^{(6)}$ . The only difference are shifts in the weights  $\alpha$  of the Jacobi polynomials or 71 changes of the weight functions (Table 2).

## 3 Sum Factorization

In this section, we present an algorithm for the fast numerical generation of the local 74 element matrices (6) for tetrahedra. The methods are based on fast summation tech-75 niques presented in [7, 8] and are carried out in detail for the example of the matrix 76  $\hat{\mathscr{S}}^{(6)}$  (8).

All one dimensional integrals in (8) are computed numerically by a Gaussian 78 quadrature rule with points  $x_k$ ,  $k=1,\ldots,p+1$  and corresponding weights  $\omega_k$ . The 79 points and weights are chosen such that

$$\int_{-1}^{1} f(x) \, dx = \sum_{l=1}^{p+1} \omega_l f(x_l) \quad \forall f \in \mathcal{P}_{2p}.$$
 (11)

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Since only polynomials of maximal degree 2p are integrated in (8), these integrals are evaluated exactly. Therefore, we have to compute

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$$m_{ijk,i'j'k'} = \sum_{l=1}^{p+1} \omega_l \hat{p}_i^0(x_l) \hat{p}_{i'}^0(x_l)$$

$$\times \sum_{m=1}^{p+1} \omega_m \left(\frac{1-x_m}{2}\right)^{i+i'+1} \hat{p}_{j'}^{2i'-1}(x_m) \hat{p}_j^{2i-1}(x_m)$$

$$\times \sum_{n=1}^{p+1} \omega_n \left(\frac{1-x_n}{2}\right)^{i+j+i'+j'+2} p_k^{2i+2j-2}(x_n) p_{k'}^{2i'+2j'-2}(x_n),$$

i.e., for all  $(i, j, k, i', j', k') \notin \mathbb{S}^p(ijk, i'j'k')$ , cf. (10), (9). This is done by the following 83 algorithm.

#### **Algorithm 3.1** 1. Compute

$$h_{i;i'}^{(1)} = \sum_{l=1}^{p+1} \omega_l \hat{p}_i^0(x_l) \hat{p}_{i'}^0(x_l)$$
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for all  $i, i' \in \mathbb{N}$  satisfying  $|i - i'| \le 2$  and  $i, i' \le p$ . 87

2. Compute

$$h_{i,j;i',j'}^{(2)} = \sum_{m=1}^{p+1} \omega_m \left(\frac{1-x_m}{2}\right)^{i+i'+1} \hat{p}_j^{2i-1}(x_m) \hat{p}_{j'}^{2i'-1}(x_m)$$
 89

for all  $i, j, i', j' \in \mathbb{N}$  satisfying  $|i - i'| \le 2$ ,  $|i + j - i' - j'| \le 4$ ,  $i + j \le p$  and 90  $i' + j' \le p$ .

3. Compute

$$h_{\beta,k;\beta,'k'}^{(3)} = \sum_{n=1}^{p+1} \omega_n \left(\frac{1-x_n}{2}\right)^{\beta+\beta'+2} p_k^{2\beta-2}(x_n) p_{k'}^{2\beta'-2}(x_n)$$
 93

for all  $k,k',\beta,\beta' \in \mathbb{N}$  satisfying  $|\beta-\beta'| \leq 4$ ,  $|\beta+k-\beta'-k'| \leq 4$ ,  $|\beta+k| \leq p$ and  $\beta' + k' < p$ . 96

4. For all 
$$(i, j, k, i', j', k') \notin \mathfrak{S}^p(ijk, i'j'k')$$
, set

$$m_{ijk,i'j'k'} = h_{i,i'}^{(1)} h_{i,j;i',j'}^{(2)} h_{i+j,k;i'+j',k'}^{(3)}.$$
 97

The algorithm requires the numerical evaluation of Jacobi and integrated Jacobi 98 polynomials at the Gaussian points  $x_l$ ,  $l=1,\ldots,p+1$ . In the next subsection, we present an algorithm which computes the required values  $\hat{p}_k^{\alpha}(x_l)$ ,  $m=1,\ldots,p+1$ , 100  $k=1,\ldots,p, \alpha=1,\ldots,2p$  in  $\mathcal{O}(p^3)$  operations. 101

## 4 Fast Evaluation of Integrated Jacobi Polynomials

The integrated Jacobi polynomials needed in the computation of  $m_{ijk,i'j'k'}$  (7) are 103  $\hat{p}_i^0(x), \hat{p}_i^{2i-1}(x)$  (progressing in odd steps with respect to the parameter  $\alpha$ ) and 104  $\hat{p}_k^{2i+2j-2}(x)$  (progressing in even steps with respect to the parameter  $\alpha$ ). For i+j+1 $k \le p$  with  $i \ge 2$  and  $j, k \ge 1$  this means that 106

$$\left[ \hat{p}_{i}^{0}(x) \right]_{2 \leq i \leq p}, \left[ \hat{p}_{j}^{3}(x) \right]_{1 \leq j \leq p}, \dots, \left[ \hat{p}_{j}^{2p-3}(x) \right]_{1 \leq j \leq p},$$

$$\left[ \hat{p}_{k}^{4}(x) \right]_{1 \leq k \leq p}, \dots, \left[ \hat{p}_{k}^{2p-4}(x) \right]_{1 \leq k \leq p}$$

are needed. Since one group proceeds in even, the other one in odd steps, the total of 108 integrated Jacobi polynomials that are needed is 109

$$\hat{p}_n^a(x), \qquad 1 \le n \le p-3, \quad 3 \le a \le 2p-3,$$

if we consider the integrated Legendre polynomials separately. However, integrating 111 both sides of (3) yields 112

$$\hat{p}_{n+1}^{\alpha-1}(x) = \frac{1}{2n+\alpha} \left( (n+\alpha) \hat{p}_{n+1}^{\alpha}(x) - n \hat{p}_{n}^{\alpha}(x) \right), \tag{113}$$

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valid for all  $n \ge 0$ . Using this relation starting from the integrated Jacobi polynomials 114 of highest degree, i.e.,  $\alpha = 2i - 1 = 2p - 3$ , the remaining Jacobi polynomials can 115 be computed using only two elements of the previous row. Note that for the initial 116 values n=1 we have  $\hat{p}_1^{\alpha}(x)=1+x$  for all  $\alpha$ . For assembling the polynomials of highest degree the three term recurrence (4) is used. Summarizing, the evaluation of 118 the functions at the Gaussian points can be done in  $\mathcal{O}(p^3)$  operations. This is optimal 119 in the three-dimensional case, but not in the two-dimensional case.

### 5 Complexity of the Algorithm

The cost of the last three steps is  $\mathcal{O}(p^3)$ , the first step requires  $\mathcal{O}(p^2)$  operations. 122 Together with the evaluation of the Jacobi polynomials, the algorithm requires in 123 total  $\mathcal{O}(p^3)$  flops.

This algorithm uses only the sparsity structure (10). Since all matrices  $\hat{\mathscr{I}}^{(m)}$ , 125  $m=1,\ldots,21$ , have a similar sparsity structure of the form (10), this algorithm can be 126 extended to all ingredients which are required for assembling/computing the element 127 stiffness matrix (6) for the Laplacian, see [3]. The algorithm can also be extended 128 to mass matrices or matrices arising from the discretization of elliptic problems in 129 H(curl) and H(div), see [4]. For two-dimensional problems, the third step of the 130 algorithm is not necessary. However, the values  $h_{i,j;i',j'}^{(2)}$  have to be computed. Since this requires  $\mathcal{O}(p^3)$  floating point operations, the total cost in 2D is also  $\mathcal{O}(p^3)$ . 132

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## **AUTHOR QUERY**

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# A Non-overlapping Quasi-optimal Optimized Schwarz Domain Decomposition Algorithm for the Helmholtz **Equation**

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1 Introduction 13

In this paper, we present a new non-overlapping domain decomposition algorithm 14 for the Helmholtz equation. We are particularly interested in the method introduced 15 by P.-L. Lions [6] for the Laplace equation and extended to the Helmholtz equa- 16 tion by B. Després [3]. However, this latest approach provides slow convergence 17 of the iterative method due to the choice of the transmission conditions. Thus, in 18 order to improve the convergence, several methods were developed [4, 5, 9, 10]. 19 The main idea in [5, 9] consists in computing a more accurate approximation of the 20 Dirichlet-to-Neuman (DtN) operator than the one proposed in [3] by using partic- 21 ular local transmission conditions. We propose in this work a different approach to 22 approximate the DtN map. We mainly use Padé approximants to suitably localize the 23 nonlocal representation of the DtN operator [8, 11]. This results in an algorithm with 24 quasi-optimal convergence properties.

#### 2 Model Problem and Non-overlapping Domain Decomposition Method

For the sake of simplicity, we limit ourselves to the evaluation of the two-dimensional 28 time-harmonic scattering wave by an obstacle denoted by K. The three-dimensional 29 case is treated similarly without adding any difficulty. We consider the model prob- 30 lem given by the system

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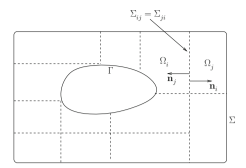


Fig. 1. Example of 2D non-overlapping domain decomposition method

$$\begin{cases} \Delta u + k^2 u = 0 & \text{in } \mathbb{R}^2 \backslash K, \\ u = f & \text{on } \Gamma = \partial K, \\ \lim_{|x| \to \infty} |x|^{1/2} (\partial_{|x|} u - iku) = 0, \end{cases}$$
 (1)

composed of the Helmholtz equation, the Dirichlet condition on  $\Gamma$  (TE polarization in electromagnetics) where  $f=-e^{ik\alpha\cdot x}$  describes the incident plane wave with 33  $|\alpha|=1$  and k is the wavenumber, and the Sommerfeld radiation condition. To 34 solve (1), we combine the absorbing boundary condition method [1, 2] with non-overlapping domain decomposition methods. The absorbing boundary conditions 36 method consists of truncating the computational domain using an artificial interface 57, and reducing the system (1) to the following one

$$\begin{cases} \Delta u + k^2 u = 0 & \text{in } \Omega, \\ u = f & \text{on } \Gamma, \\ \partial_{\mathbf{n}} u + \mathcal{B} u = 0 & \text{on } \Sigma, \end{cases}$$
 (2)

where  $\Omega$  is the bounded domain enclosed by  $\Sigma$  and  $\Gamma$ ,  $\mathscr{B}$  indicates the approximation of the Dirichlet-to-Neuman (DtN) operator, and  $\mathbf{n}$  is the outward normal to  $\Sigma$ . We are interested in the domain decomposition method introduced in [3, 6]. The first step of this approach consists in splitting  $\Omega$  into several subdomains  $\Omega_i$ ,  $i = 1, \ldots, N$ , such 42 that

- $\overline{\Omega} = \bigcup_{i=1}^{N} \overline{\Omega}_i \ (i=1,\ldots,N),$
- $\Omega_i \cap \Omega_j = \emptyset$ , if  $i \neq j$ ,  $(i, j = 1, \dots, N)$ ,
- $\partial \Omega_i \cap \partial \Omega_j = \overline{\Sigma}_{ij} = \overline{\Sigma}_{ji} \ (i, j = 1, ..., N)$  is the artificial interface (see Fig. 1) separating  $\Omega_i$  from  $\Omega_j$  as long as its interior  $\Sigma_{ij}$  is not empty.

Then, applying the Lions-Després algorithm, the solution of the initial problem (1) 48 is reduced to an iterative procedure, where each iteration is performed by solving the 49 local problems 50

DDM with Generalized Impedance Boundary Conditions

$$\begin{cases} \Delta u_{i}^{(n+1)} + k^{2} u_{i}^{(n+1)} = 0 & \text{in } \Omega_{i}, \\ u_{i}^{(n+1)} = f_{i} & \text{on } \Gamma_{i}, \\ \partial_{\mathbf{n}_{i}} u_{i}^{(n+1)} + \mathcal{B} u_{i}^{(n+1)} = 0 & \text{on } \Sigma_{i} \end{cases}$$
(3a)

$$\partial_{\mathbf{n}_i} u_i^{(n+1)} + \mathcal{S} u_i^{(n+1)} = g_{ij}^{(n)} \quad \text{on } \Sigma_{ij}, \tag{3b}$$

and forming the quantities to be transmitted through the interfaces

$$g_{ij}^{(n+1)} = -\partial_{\mathbf{n}_j} u_j^{(n+1)} + \mathcal{S} u_j^{(n+1)} = -g_{ij}^{(n)} + 2\mathcal{S} u_j^{(n+1)} \quad \text{on } \Sigma_{ij}, \tag{4}$$

where  $u_i = u|_{\Omega_i}$ ,  $\mathbf{n}_i$  (resp.  $\mathbf{n}_j$ ) is the outward unit normal of the boundary of  $\Omega_i$  52 (resp.  $\Omega_i$ ), i = 1, ..., N, j = 1, ..., N,  $\Gamma_i = \partial \Omega_i \cap \Gamma$  and  $\Sigma_i = \partial \Omega_i \cap \Sigma$ . Note that 53 the boundary condition on  $\Gamma_i$  (resp.  $\Sigma_i$ ) does not take place if the interior of  $\partial \Omega_i \cap \Gamma$  54 (resp.  $\partial \Omega_i \cap \Sigma$ ) is the empty set.

#### 3 New Transmission Conditions

It is well established that the convergence of the domain decomposition algorithms 57 depends on the choice of the transmission operator  $\mathcal{S}$ . In the original method pro- 58 posed by B. Després [3], the usual approximation of the DtN operator  $\mathcal{S}u = -iku$  59 is used. The resulting algorithm does not treat efficiently the evanescent modes of 60 the iteration operator which impairs the iterative method [9]. In order to improve the 61 convergence, two techniques, based on the modification of the operator  $\mathcal{S}$ , were 62 proposed. First, the optimized Schwarz method introduced by Gander et al. [5]. 63 It consists of using local second-order approximations of the DtN operator  $\mathcal{S}u = 64$  $\delta u + \gamma \partial_s^2 u$ , where  $\partial_s$  is the tangential derivative operator, and the coefficients  $\delta$  65 and  $\gamma$  are optimized using the rate of convergence obtained in the case of the 66 half-plane. The second method, called the "evanescent modes damping algorithm" 67 (EMDA), was introduced by Boubendir et al. [9, 10]. In this case, S is chosen as 68  $\mathcal{S}u = -\iota ku + \mathcal{X}u$  where  $\mathcal{X}$  is a self-adjoint positive operator. We only consider 69 here the usual case where  $\mathscr{X}$  is a real-valued positive coefficient. In this paper we 70 propose a new "square-root" transmission operator [7, 8, 11] that takes the following 71 form:

$$\mathscr{S}u = -\iota k \operatorname{Op}\left(\sqrt{1 - \frac{\xi^2}{k_{\varepsilon}^2}}\right) u,\tag{5}$$

where 73

$$k_{\varepsilon} = k + \iota \varepsilon \tag{6}$$

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is a complexified wavenumber, and the notation  $\sqrt{z}$  designates the principal deter- 74 mination of the square-root of a complex number z with branch-cut along the nega- 75 tive real axis. This choice of the square-root operator is motivated by developments 76 of absorbing boundary conditions (ABC) for scattering problems [1, 2]. Generally 77 speaking, the usual techniques to develop absorbing boundary conditions consists 78 mainly in using Taylor expansions to approximate the symbol of the DtN operator. 79 However, these approximations prevent the modelling of the three parts describing 80 the wave (propagating, evanescent and transition) at the same time, which affects, in 81 return, the final accuracy of the solution. This problem can be solved by high-order 82 local ABC introduced in [7, 8], which uses (5) to model all the scattering modes: 83 propagating, evanescent as well as (in an approximate way) grazing. The localization is performed with complex Padé approximants, and the coefficient  $\varepsilon$  in (6) can 85 then be chosen to minimize spurious reflections at the boundary. In the context of 86 domain decomposition methods, this optimization of  $\varepsilon$  improves the spectrum of the 87 iteration operator on these grazing modes. As it is shown in [8], the optimal value of 88 this parameter is given by  $\varepsilon = 0.4k^{1/3}\mathcal{H}^{2/3}$ , where  $\mathcal{H}$  is the mean curvature on the 89 interface.

## 4 Localization of the Square-Root Operator Using Padé Approximants

Because the square-root operator (5) is nonlocal, its use in the context of finite 93 element method is ineffective since it would lead to consider full matrices for the 94 transmission boundaries. A localization process of this operator can be efficiently 95 done by using partial differential (local) operators and obtain sparse matrices. This 96 is performed [7, 8, 11] in rotating branch-cut approximation of the square-root and 97 then applying complex Padé approximants of order  $N_p$ , 98

$$\sqrt{1 - \frac{\xi^{2}}{k_{\varepsilon}^{2}}} u \approx R_{N_{p}}^{\alpha} \left(-\frac{\xi^{2}}{k_{\varepsilon}^{2}}\right) u$$

$$= C_{0}u + \sum_{\ell=1}^{N_{p}} A_{\ell} \left(\frac{-\xi^{2}}{k_{\varepsilon}^{2}}\right) \left(1 + B_{\ell} \left(\frac{-\xi^{2}}{k_{\varepsilon}^{2}}\right)\right)^{-1} u, \tag{7}$$

which correspond to the complex Padé approximation

$$\sqrt{1+z} \approx R_{N_p}^{\alpha}(z) = C_0 + \sum_{\ell=1}^{N_p} \frac{A_{\ell}z}{1+B_{\ell}z},$$
 (8)

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and where the complex coefficients  $C_0$ ,  $A_\ell$  and  $B_\ell$  are given by

$$C_0 = e^{i\frac{\alpha}{2}} R_{N_p}(e^{-i\alpha} - 1), A_{\ell} = \frac{e^{-\frac{i\alpha}{2}} a_{\ell}}{(1 + b_{\ell}(e^{-i\alpha} - 1))^2}, B_{\ell} = \frac{e^{-i\alpha} b_{\ell}}{1 + b_{\ell}(e^{-i\alpha} - 1)}.$$

Here,  $\alpha$  is the angle of rotation,  $(a_{\ell},b_{\ell})$ ,  $\ell=1,\ldots,N_p$ , are the standard real Padé 101 coefficients

$$a_{\ell} = \frac{2}{2N_p + 1} \sin^2(\frac{\ell \pi}{2N_p + 1}), b_{\ell} = \cos^2(\frac{\ell \pi}{2N_p + 1}),$$
 (9)

and  $R_{N_p}$  is the real Padé approximant of order  $N_p$ 

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$$\sqrt{1+z} \approx R_{N_p}(z) = 1 + \sum_{\ell=1}^{N_p} \frac{a_{\ell}z}{1+b_{\ell}z}.$$
 (10)

For a variational representation, the approximation of the Padé-localized squareroot transmission operators is realized by using auxiliary coupled functions [7, 11]

$$\mathscr{S}u = -\iota k \left(C_0 u + \sum_{\ell=1}^{N_p} A_\ell \operatorname{div}_{\Sigma_d} \left(\frac{1}{k_{\mathcal{E}}^2} \nabla_{\Sigma_d} \varphi_\ell\right)\right) \quad \text{on } \Sigma_d, \tag{11}$$

where the functions  $\varphi_{\ell}$ ,  $\ell=1,...,N_p$ , are defined on any artificial interface  $\Sigma_d$  as the 106 solutions of the surface PDEs

$$(1 + B_{\ell} \operatorname{div}_{\Sigma_d}(\frac{1}{k_{\varepsilon}^2} \nabla_{\Sigma_d})) \varphi_{\ell} = u.$$
(12)

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The resulting transmitting condition is a Generalized Impedance Boundary Condition, and is denoted by  $\mathrm{GIBC}(N_p,\alpha,\varepsilon)$  for the Padé approximation with  $N_p$  auxiliary functions, for an angle of rotation  $\alpha$  and a damping parameter  $\varepsilon$ . The lowest-order approximation  $\mathscr{S} = -ikI$  (resp.  $\mathscr{S} = -iku + \mathscr{X}u$ ) is denoted by IBC(0) (resp. 111 IBC( $\mathscr{X}$ )).

#### **5 Numerical Results**

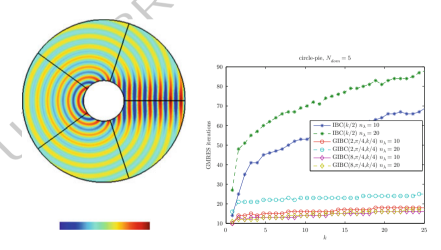
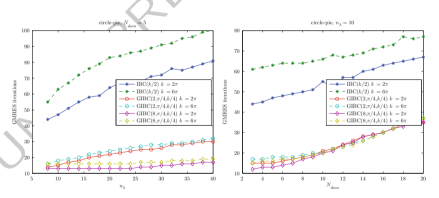


Fig. 2. Left: decomposition of the computational domain. Right: iteration number with respect to the wavenumber k for two densities of discretization  $n_{\lambda}$ 

The numerical tests presented here concern the scattering of a plane wave by 114 a unit sound-soft circular cylinder. We truncate the computational domain using a 115 circle of radius equal to 4, on which the second-order Bayliss-Turkel absorbing condition [1] is set (see problem (2)). We perform these numerical tests on partitions 117 of the type displayed in Fig. 2, and we refer to them as "circle-pie". We use a finite 118 element method with linear (P1) basis functions to approximate the solution in each 119 subdomain. The implementation of this method with Padé approximants is described 120 in [11]. The iterative problem is solved using GMRES and the iterations are stopped 121 when the initial residual has decreased by a factor of  $10^{-6}$ .

We begin by testing the iterative method with respect to the wavenumber k. Let 123 us consider the number of subdomains  $N_{\rm dom} = 5$ . Because the interfaces are straight, 124 as depicted on the left picture of Fig. 2,  $\varepsilon$  cannot be optimized as described in Sect. 3. 125 However, numerical simulations show that  $\varepsilon = k/4$  is an appropriate choice for this 126 kind of interfaces. On the right picture of Fig. 2, we represent the behavior of the 127 number of iterations. We choose two densities of discretization points per wave- 128 length  $n_{\lambda}$ . We compare the new algorithm noted GIBC $(N_p, \pi/4, \varepsilon)$ , where  $N_p$  is the Padé number and  $\pi/4$  the angle of rotation, with the EMDA algorithm designated by 130 IBC(k/2). In this latest case, the number of iterations clearly increases with respect 131 to k and  $n_{\lambda}$ . However, for GIBC( $N_p$ ,  $\pi/4$ ,  $\varepsilon$ ), the convergence rate is almost independent of both the wavenumber and density of discretization points per wavelength. 133 In particular, the convergence for  $N_p = 2$  and  $N_p = 8$  is similar. This means that the cost of the solution when solving local problems is comparable to the other methods 135 with usual local transmission conditions (see [11] for more details).

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**Fig. 3.** Number of iterations with respect to the density of discretization  $n_{\lambda}$  and the number of subdomains N<sub>dom</sub>

In Fig. 3, we show the number of iterations with respect to: (i) the density of discretization points per wavelength  $n_{\lambda}$  for two wavenumbers k, and (ii) the number of 138 subdomains  $N_{dom}$ . We can see that for a small Padé number ( $N_p = 2$ ), the convergence is almost independent of the mesh size. A larger choice of  $N_p$  will provide an 140 optimal result. We also see that the number of iterations with respect to the number 141

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of	subdomains	does 1	not (	deteriorate	with	increasing	values	of $N_p$	or $k$ ,	contrary to	0 1	42
ΙB	C(k/2).										1	43

6 Conclusion 144

We designed in this paper a new non-overlapping domain decomposition algorithm 145 for the Helmholtz equation with quasi-optimal convergence properties. It is based on 146 a suitable approach which consists in using Padé approximants to approximate the 147 DtN operator. The analysis of this new approach can be found in [11], as well as 148 several numerical tests including the three-dimensional case.

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# A Continuous Approach to FETI-DP Mortar Methods: Application to Dirichlet and Stokes Problem

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**Summary.** In this contribution we extend the FETI-DP mortar method for elliptic problems 9 introduced by Bernardi et al. [2] and Chacón Vera [3] to the case of the incompressible Stokes 10 equations showing that the same results hold in the two dimensional setting. These ideas 11 extend easily to three dimensional problems. Finally some numerical tests are shown as a 12 conclusion. This contribution is a condensed version of a more detailed forthcoming paper. 13 We use standard notation, see for instance [1].

### 1 Incompressible Stokes Equations

Let  $\Omega\subset\mathbb{R}^2$  be a polygonal domain. We look for  $u\in\mathbf{H}^1_0(\Omega)=(H^1_0(\Omega))^2$  and  $p\in L^2(\Omega)$  such that  $\int_\Omega p=0$  and

$$(\nabla u, \nabla v)_{\Omega} - (p, div(v))_{\Omega} = (f, v)_{\Omega}, \quad \forall v \in \mathbf{H}_0^1(\Omega)$$
$$- (q, div(u))_{\Omega} = 0, \qquad \forall q \in L^2(\Omega).$$

We better accommodate the restriction on the pressure by adding a new scalar 18 unknown: we look for a pair of values  $(u,\tau)\in \mathbf{H}^1_0(\Omega)\times\mathbb{R}$  and  $p\in L^2(\Omega)$  such 19 that

$$\begin{split} (\nabla u, \nabla v)_{\Omega} - (p, div(v))_{\Omega} + t \, (\tau - \int_{\Omega} p) &= (f, v)_{\Omega}, \quad \forall (v, t) \in \mathbf{H}^{1}_{0}(\Omega) \times \mathbb{R} \\ &- (q, div(u))_{\Omega} - \tau \int_{\Omega} q &= 0, \qquad \quad \forall q \in L^{2}(\Omega). \end{split}$$

Set  $W = \mathbf{H}_0^1(\Omega) \times \mathbb{R}$  normed by  $\|\underline{v}\|_W^2 = \|(v,t)\|_W^2 = \|\nabla v\|_{0,\Omega}^2 + t^2$  for any  $\underline{v} = (v,t) \in \mathbb{R}$  given by 22 W, let  $(\cdot, \cdot)_W$  be the scalar product on W and  $b: W \times L^2(\Omega) \mapsto \mathbb{R}$  given by 22

$$b(q,(v,t)) = -(q,div(v))_{\Omega} - t \int_{\Omega} q.$$
 23

Then, we look for  $\underline{u} = (u, \tau) \in W$  and  $p \in L^2(\Omega)$  such that

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$$(u,v)_W + b(p,v) = (f,v)_Q, \quad \forall v \in W$$
 (1)

$$b(q, u) = 0, \quad \forall q \in L^2(\Omega).$$
 (2)

25

It is quite straightforward to see that:

**Lemma 1.** There exists a positive constant  $\beta > 0$  such that for all  $p \in L^2(\Omega)$ 

$$\sup_{(v,t)\in W} \frac{b(p,(v,t))}{\|(v,t)\|_{W}} \ge \sup_{v\in \mathbf{H}_{0}^{1}(\Omega),t\in\mathbb{R}} \frac{b(p,(v,t))}{(\|\nabla v\|_{0,\Omega}^{2} + t^{2})^{1/2}} \ge \beta \|p\|_{0,\Omega}. \tag{3}$$

As a consequence, problem (1)–(2) is well posed and its unique solution is the one of 27 the original Stokes problem with Dirichlet homogeneous boundary conditions. 28

Next, we split  $\Omega = \bigcup_{s=1}^{S} \Omega^{s}$  with nonoverlaping polygonal subdomains, suppose that 29

$$\Gamma_{s,t} = \partial \Omega^s \cap \partial \Omega^t$$
 30

is either an edge (i.e., a segment), a crosspoint or empty and, finally, consider  $\mathscr{E}_0=31$   $\{\varGamma_e\}_{e=1,\dots,E}$  the sorted set of all edges inside  $\Omega$ . We suppose that each  $\Omega^s$  is of area 32  $\mathscr{O}(H^2)$  and shape regular while each  $\varGamma_e$  is of length  $\mathscr{O}(H)$  for some fixed H>0. 33 The set of all vertices of the polygonal subdomains  $\Omega^s$  that are not on  $\partial\Omega$  will be 34 called **cross points** and denoted by  $\mathscr{C}$ . Finally, we denote by  $[v]_{\varGamma_e}$  the jump across 35 any interface  $\varGamma_e$ .

We take

$$\begin{split} X_{\delta} &= \{v \in L^2(\Omega); v^s = v_{|_{\Omega^s}} \in H^1(\Omega^s) \cap H^1_0(\Omega), \ 1 \leq s \leq S\}, \\ X &= \{v \in X_{\delta}, \ [v]_{\Gamma_e} \in H^{1/2}_{00}(\Gamma_e), \ \forall \, \Gamma_e \in \mathcal{E}_0\}. \end{split}$$

With  $\mathbf{X} = X \times X$  we construct  $\mathbf{V} = \mathbf{X} \times \mathbb{R}$  and represent by  $\underline{v} = (v, t)$  any element 38 of  $\mathbf{V}$  where  $v \in \mathbf{X}$  and  $t \in \mathbb{R}$ .  $\mathbf{V}$  is Hilbert space with norm  $\|\underline{v}\|_{\mathbf{V}}^2 = |v|_{\mathbf{X}}^2 + t^2$  where, 39 thanks to Poincaré's inequality, the norm of v is

$$|v|_{\mathbf{X}} = \{ \sum_{s=1}^{S} \|\nabla v^{s}\|_{0,\Omega^{s}}^{2} + \sum_{e=1}^{E} \|[v]_{\Gamma_{e}}\|_{1/2,00,\Gamma_{e}}^{2} \}^{1/2}.$$

Here,  $\|\cdot\|_{1/2,00,\Gamma_e}$  is the norm induced by the scalar product  $(\cdot,\cdot)_{1/2,00,\Gamma_e}$  on  $H_{00}^{1/2}(\Gamma_e)$ , 41 see [5]. To simplify, let  $\{\cdot,\cdot\}_{\Gamma_e}=(\cdot,\cdot)_{1/2,00,\Gamma_e}$ . For the pressure space we consider 42  $\mathbf{M}=\prod_{s=1}^S L^2(\Omega^s)(\approx L^2(\Omega))$  and define the continuous bilinear form  $b:\mathbf{M}\times\mathbf{V}\mapsto\mathbb{R}$  43 given by

$$b(q,\underline{v}) = -\sum_{s=1}^S (q^s,div(v^s))_{\Omega^s} - t\sum_{s=1}^S \int_{\Omega^s} q^s, \quad orall q^s \in L^2(\Omega^s).$$

Next, for each  $\Gamma_e \in \mathscr{E}_0$  we take  $\mathbf{H}_{00}^{1/2}(\Gamma_e) = (H_{00}^{1/2}(\Gamma_e))^2$ , and handle the Lagrange 45 multipliers for the jumps with the space  $\mathbf{N} = \prod_{e=1}^E \mathbf{H}_{00}^{1/2}(\Gamma_e)$ .

We propose to look for  $\underline{u} = (u, \tau) \in \mathbf{V}$ ,  $p = \{p^s\}_s \in \mathbf{M}$  and  $\lambda = \{\lambda_e\}_e \in \mathbf{N}$  such 47 that

$$\begin{split} \sum_{s=1}^{S} (\nabla u^{s}, \nabla v^{s})_{\Omega^{s}} + \sum_{e=1}^{E} \{[u]_{\Gamma_{e}}, [v]_{\Gamma_{e}}\}_{\Gamma_{e}} + \tau t \\ - \sum_{s=1}^{S} (p^{s}, div(v^{s}))_{\Omega^{s}} - t \sum_{s=1}^{S} \int_{\Omega^{s}} p^{s} + \sum_{e=1}^{E} \{\lambda_{e}, [v]_{\Gamma_{e}}\}_{\Gamma_{e}} = \sum_{s=1}^{S} (f, v^{s})_{\Omega^{s}}, \\ - \sum_{s=1}^{S} (q^{s}, div(u^{s}))_{\Omega^{s}} - \tau \sum_{s=1}^{S} \int_{\Omega^{s}} q^{s} = 0, \\ \sum_{e=1}^{E} \{\mu_{e}, [u]_{\Gamma_{e}}\}_{\Gamma_{e}} = 0 \\ = (v, t) \in \mathbf{V}, \ q = \{q^{s}\}_{s} \in \mathbf{M} \ \text{and} \ \mu = \{\mu_{e}\}_{e} \in \mathbf{N}. \end{split}$$

for all  $\underline{v} = (v,t) \in \mathbf{V}$ ,  $q = \{q^s\}_s \in \mathbf{M}$  and  $\mu = \{\mu_e\}_e \in \mathbf{N}$ .

We see that we added the jumps to the elliptic terms and replaced the pair- 50 ings  $H_{00}^{-1/2}(\Gamma)-H_{00}^{1/2}(\Gamma)$  for the normal fluxes on the edges by the scalar product 51 in  $H_{00}^{1/2}(\Gamma)$ . As a consequence, we have made a regularization of order 1 for the 52 Lagrange multipliers and now all terms are suitable to compute in a Galerkin ap- 53 proach. Moreover, the solution to this problem is that of the incompressible Stokes 54 equations on  $\Omega$ .

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Next, we elliminate via a standard Schur process the primal variables u and p 56 in terms of the dual variable  $\lambda$ , and obtain a dual problem that once solved will 57 give the correct boundary data for the primal variables. Thanks to the fact that the 58 elliptic part is the scalar product on V, that the inf-sup condition for the bilinear form 59 b is achieved with velocities without jumps and that the inf-sup condition for c is 60 achieved with velocities with jumps, our dual problem is a well posed symmetric 61 positive definite problem.

## 2 Finite Dimensional Approach

We consider a conforming triangulation  $\mathcal{T}_h$ , h is the mesh size, of  $\overline{\Omega}$  that contains 64 the skeleton  $\mathscr{E}_0$  as union of edges of triangles and such that on each edge only one 65 partition is inherited from both sides. As  $\mathcal{T}_h$  is also compatible with the subdivision 66 of  $\Omega$ , its restriction to each  $\overline{\Omega}_s$  gives a mesh  $\mathscr{T}_h^s$  on  $\overline{\Omega}^s$ . We use the Taylor-Hood finite 67 element for the velocity and pressure pair on each subdomain. Define the family of 68 subspaces  $\{Y_h\}_h \subset H_0^1(\Omega)$  and  $\{Q_h\}_h \subset H^1(\Omega)$  given by

$$\begin{split} Y_h &= \{ v \in H^1_0(\Omega); \ v_{|_{\kappa}} \in \mathbb{P}_2(\kappa), \ \forall \kappa \in \mathscr{T}_h \}, \\ Q_h &= \{ p \in H^1(\Omega); \ p_{|_{\kappa}} \in \mathbb{P}_1(\kappa), \ \forall \kappa \in \mathscr{T}_h \} \end{split}$$

where  $\mathbb{P}_r(\kappa)$  is the space of polynomials of degree less or equal to r in the two 70 variables x and y. On each subdomain, we take also 71

$$Y_h(\Omega^s) = Y_h \cap H^1(\Omega^s), \quad Q_h(\Omega^s) = Q_h \cap H^1(\Omega^s), \ s \leq S.$$

Consider now  $\mathbf{X}_h = X_h \times X_h$ , where  $X_h$  is the broken version of  $Y_h$  given by

$$X_h = \{ v \in L^2(\Omega); \ v^s \in Y_h^s, \ \forall \ s = 1, 2, \dots, S,$$
  
and  $v$  is continuous at every cross point in  $\mathscr{C} \} \subset X$ ,

define  $\mathbf{V}_h = \mathbf{X}_h \times \mathbb{R}$ ,  $\mathbf{M}_h = \prod_{s=1}^S Q_h(\Omega^s)$  and finally  $\mathbf{N}_h \subset \mathbf{N}$  is given by the restriction of functions in  $\mathbf{X}_h$  to the skeleton  $\mathscr{E}_0$ .

The discrete uniform inf-sup condition for c on the pair  $\mathbf{V}_h$  and  $\mathbf{N}_h$  is by now a 75 well known result and the discrete uniform inf-sup condition for b is a consequence 76 of Theorem 1.12 pp. 130 in [4]. The idea is to use locally on each subdomain  $\Omega^s$  the 77 stability of the pair  $\mathbb{P}_2 - \mathbb{P}_1$  and that of the pair  $\mathbb{P}_2 - \mathbb{P}_0$  globally on the substructures 78  $\Omega^s$  of  $\Omega$ . This inf-sup condition is achieved with a discrete continuous function 79 in the wohle of  $\Omega$  and, as a consequence, the continuous setting is replicated and 80 the equation for the multiplier can be solved via Conjugate Gradient Method (CG) 81 without preconditioner. Then, we have

- 1. An external computational cicle, the CG for the Lagrange multiplier with a fixed 83 number of iterations independent of the discretization parameter *h* and 84
- At each iteration of this external cicle, the resolution of a primal problem of the 85 form:

Find 
$$(\underline{w}_h, q_h) \in \mathbf{V}_h \times \mathbf{M}_h$$
 such that

$$(\underline{w}_h, \underline{v}_h)_{\mathbf{V}} + b(q_h, \underline{v}_h) = (\xi, \underline{v}_h) \quad \forall \underline{v}_h \in \mathbf{V}_h,$$
$$b(p, \underline{w}_h) = 0 \quad \forall p \in \mathbf{M}_h$$

where for the initial residuous  $r_0$  we have  $(\xi,\underline{\nu}_h)=\sum_{s=1}^S(f,\nu_h^s)_{\Omega^s}$  and for the 88 iteration  $m\geq 0$  we have  $(\xi,\underline{\nu}_h)=\sum_{e=1}^E\{\{d_m\}_e,[\nu_h]_{\Gamma_e}\}_{\Gamma_e}=0$  89

A closer inspection to the general form of this saddle point problem for the primal 90 variables shows that the solution can be obtained by means of independent solves 91 per subdomain. Ordering the unknows per subdomains,  $x^s = (u^s, p^s)$  and  $x^C = u^C$ , 92 the linear system for the primal variables is

$$\begin{pmatrix} M_{11} & M_{1,2} & \dots & \dots & M_{1,S} & M_{1,C} & D_1 \\ M_{21} & M_{2,2} & M_{2,3} & \dots & \dots & M_{2,C} & D_2 \\ M_{31} & M_{3,2} & M_{3,3} & M_{3,4} & \dots & \dots & M_{3,C} & D_3 \\ \vdots & \ddots & \ddots & \ddots & \ddots & \vdots & \vdots & \vdots \\ \vdots & \ddots & \ddots & \ddots & \ddots & \vdots & \vdots & \vdots \\ \vdots & \dots & \dots & M_{S,S-1} & M_{S,S} & M_{S,C} & D_S \\ M_{1,C}^t & M_{2,C}^t & \dots & \dots & M_{S-1,C}^t & M_{S,C}^t & M_{C,C} & 0 \\ D_1^t & D_2^t & \dots & \dots & D_{S-1}^t & D_S^t & 0^t & 1 \end{pmatrix} \begin{pmatrix} x^1 \\ x^2 \\ x^3 \\ \vdots \\ \vdots \\ x^S \\ x^C \\ \tau \end{pmatrix} = \begin{pmatrix} b^1 \\ b^2 \\ b^3 \\ \vdots \\ \vdots \\ b^S \\ b^C \\ 0 \end{pmatrix}$$

where the different blocks are of the form

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$$M_{s,s} = \begin{pmatrix} A_{s,s} & B_{s,s} \\ B_{s,s}^t & 0 \end{pmatrix}, M_{s,s'} = \begin{pmatrix} A_{s,s'} & 0 \\ 0 & 0 \end{pmatrix}, M_{s,C} = \begin{pmatrix} A_{s,C} \\ B_{s,C}^t \end{pmatrix}, M_{C,C} = A_{C,C}$$
 95

here each block  $M_{s,s}$  is similar to a standard Stokes matrix on the subdomain  $\Omega^s$ , 96 but with our interface contributions, each block  $M_{s,s'}$  is sparse and contains the 97 interaction through interfaces of the domain  $\Omega^s$  with  $\Omega^{s'}$ , the rectangular blocks  $M_{s,C}$  98 contains the interaction with the crosspoints and  $M_{C,C}$  contains the interaction of the 99 crosspoints with themselves. Although this linear system couples all the subdomains 100 it can be solved by means of the Preconditioned Conjugate Gradient Method using 101 as a preconditioner the matrix P formed by the main blocks

$$P = \begin{pmatrix} M_{11} & 0 & \cdots & \cdots & 0 & M_{1,C} & D_1 \\ 0 & M_{2,2} & 0 & \cdots & 0 & M_{2,C} & D_2 \\ 0 & 0 & M_{3,3} & 0 & \ddots & M_{3,C} & D_3 \\ \vdots & \ddots & \ddots & \ddots & \ddots & \vdots & \vdots \\ \cdots & \cdots & \cdots & 0 & M_{S,S} & M_{S,C} & D_S \\ M_{1,C}^t & M_{2,C}^t & \cdots & M_{S-1,C}^t & M_{S,C}^t & M_{C,C} & 0 \\ D_1^t & D_2^t & \cdots & D_{S-1}^t & D_S^t & 0^t & 1 \end{pmatrix}.$$

Therefore, the main task here is the resolution of a linear system of the form Px = b 103 which is done using a Schur complement process for the variables  $x^C$  and  $\tau$ . The 104 equations are 105

$$(M_{C,C} - \sum_{s=1}^{S} M_{s,C}^{t} M_{s,s}^{-1} M_{s,C}) x^{C} - \sum_{s=1}^{S} M_{s,C}^{t} M_{s,s}^{-1} D_{s} \tau = b^{C} - \sum_{s=1}^{S} M_{s,C}^{t} M_{s,s}^{-1} b^{s},$$

$$\sum_{s=1}^{S} D_{s}^{t} M_{s,s}^{-1} M_{s,C} x^{C} + (\sum_{s=1}^{S} D_{s}^{t} M_{s,s}^{-1} D_{s} - 1) \tau = \sum_{s=1}^{S} D_{s}^{t} M_{s,s}^{-1} b^{s}.$$

We finally write  $x^C$  in terms of  $\tau$  and solve first for  $\tau$ , next  $x^C$  and finally compute all 106 the  $x^s$ . As a consequence, the main job is performed with independent solves of the matrices  $M_{s,s}$  that can be performed independently, i.e., computations of the form 108

$$M_{s,s}^{-1}b^s$$
,  $M_{s,s}^{-1}M_{s,C}$ ,  $M_{s,s}^{-1}D_s$ .

#### **3 Some Numerical Tests**

For L = 1, 2, 3, ... integer we consider on  $\Omega_L = [0, L] \times [0, 1]$  the exact solution 111

$$u(x,y) = \begin{pmatrix} -\sin^3(\pi x L^{-1})\sin^2(\pi y)\cos(\pi y) \\ -L^{-1}\sin^2(\pi x L^{-1})\sin^3(\pi y)\cos(\pi x L^{-1}) \end{pmatrix}, \quad p(x,y) = \frac{x^2}{L^2} - y^2$$
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and partition  $\Omega_L$  into  $\Omega_L^s = (s-1,s) \times (0,1)$  for  $s=1,2,\ldots,L$ . For the dual problem we start our iteration process with  $\lambda_{0,e} = 0$  on each  $\Gamma_e$  and stop all iterations according to a relative residual less than  $10^{-6}$ . In this example the gradients control the jumps and there is no need to introduce them in the elliptic part; then the blocks  $M_{s,t}$  are null for  $s \neq t$ . Then, there is no need for a PCG in the internal cycle. The following 117 Table 1 shows that the iteration count for the dual problem is mesh independent on 118 different configurations Table 2 shows relative errors with respect to the true solution

	h = 1/24	h = 1/48	h = 1/96
L=4	17	17	17
$\parallel L = 8$	23	24	24
L=16	37	39	39

**Table 1.** Mesh independent iteration count for the dual problem on different configurations and for different values of h on  $\Omega_L = [0,L] \times [0,1]$ . The number of subdomains is L given by  $\Omega^s = [s-1,s] \times [0,1]$  for  $s=1,2,3,\ldots,L$ 

u and p on  $\Omega_L$  Finally, we take on  $\Omega=(0,1)^2$  the exact solution

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**Table 2.** Relative errors in velocity field and pressure for different values of h on  $\Omega_L = [0, L] \times [0, 1]$  and with the same configuration as in Table 1

$$u(x,y) = \begin{pmatrix} -\sin^3(\pi x)\sin^2(\pi y)\cos(\pi y) \\ -\sin^2(\pi x)\sin^3(\pi y)\cos(\pi x) \end{pmatrix}, \quad p(x,y) = (x - 0.25)^2(y - 0.25)^2$$
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and partition  $\Omega$  into 4 equal subdomains with a cross point at (0.5,0.5). Table 3 122 shows the results and we see that the number of iterations is independent of the mesh 123 size again (Fig. 1).

	Dual	Initial PCG	Final PCG		
h	# Iters	# Iters	# Iters	eu(h)	ep(h)
1/12	7	22	20	6.9e - 4	4.2e-03
1/24	7	21	20	8.8e - 5	1.0e - 03
1/48	7	23	21	1.2e-5	2.5e-04
1/96	7	23	23	1.4e-6	8.3e-05

**Table 3.** Results obtained when subdividing the domain  $\Omega = (0,1)^2$  into four subdomains with a cross point at (0.5,0.5)

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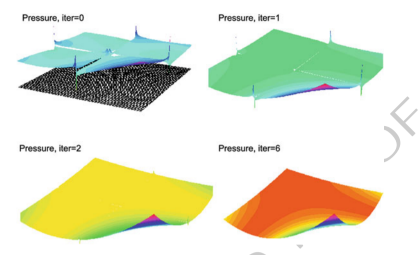


Fig. 1. Inital iteration with the underlying mesh and some contiguous iterations for the computed pressure

4 Conclusions 125

We presented a FETI-DP Mortar method applied to incompressible Stokes equations. 126 Continuity at crosspoints is retained and the jumps across interfaces are included in 127 the continuous formulation. The Lagrange multipliers are represented by their Riesz- 128 canonical isometry, which improves their regularity from  $H_{00}^{-1/2}(\Gamma)$  to  $H_{00}^{1/2}(\Gamma)$ , and 129 the mortaring is performed using the  $H_{00}^{1/2}(\Gamma)$  scalar product for each interface  $\Gamma$ . As a consequence, continuous bounds are replicated at the discrete level and no stabi- 131 lization is required. In this setting we solve a dual problem by a CG that has a mesh 132 independent condition number. The primal problems involved include the effect of 133 the coupling between neighboring subdomains at interfaces and are solved by PCG. 134 Still independent solves per subdomains are possible.

The advantage of the continuous framework introduced is the clear sight of the 136 effect of condensing all information on subdomains and interfaces before the discrete 137 work starts and the use of, to our belief, the most appropriated norms on subdomains 138 and interfaces that make no necessary the use of mesh dependent norms for obtaining 139 stability.

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# One-Shot Domain Decomposition Methods for Shape Optimization Problems

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1 Introduction 9

Shape optimization aims to optimize an objective function by changing the shape of the computational domain. In recent years, shape optimization has received considerable attentions. On the theoretical side there are several publications dealing with the existence of solution and the sensitivity analysis of the problem; see e.g., [6] and references therein. On the practical side, optimal shape design has played an important role in many industrial applications, for example, aerodynamic shape design [7], artery bypass design [1, 10], and so on. In this paper, we propose a general framework for the parallel solution of shape optimization problems, and study it in detail for the optimization of an artery bypass problem.

For PDE constrained optimization problems, there are two basic approaches: 19 nested analysis and design and simultaneous analysis and design (one-shot meth- 20 ods). As computers become more powerful in processing speed and memory capac- 21 ity, one-shot methods become more attractive due to their higher degree of paral- 22 lelism, better scalability, and robustness in convergence. The main challenges in the 23 one-shot approaches are that the nonlinear system is two to three times larger, and 24 the corresponding indefinite Jacobian system is a lot more ill-conditioned and also 25 much larger. So design a preconditioner that can substantially reduce the condition 26 number of the large fully coupled system and, at the same time, provides the scalabil- 27 ity for parallel computing becomes a very important stage in the one-shot methods. 28 There are several recent publications on one-shot methods for PDE constrained op- 29 timization problems. In [5], a reduced Hessian sequential quadratic programming 30 method was introduced for an aerodynamic design problem. In [4], a parallel full 31 space method was introduced for the boundary control problem where a Newton- 32 Krylov method is used together with Schur complement type preconditioners. In [9] 33 and [8], an overlapping Schwarz based Lagrange-Newton-Krylov approach (LNKSz) 34 was investigated for some boundary control problems. As far as we know no one has 35 studied shape optimization problems using LNKSz, which has the potential to solve 36 very large problems on machines with a large number of processors (np). The previ-

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ous work on LNKSz doesn't consider the change of the computational domain which 38 makes the study much more difficult and interesting.

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## 2 Shape Optimization on a Moving Mesh

We consider a class of shape optimization problems governed by the stationary incompressible Navier-Stokes equations defined in a two dimensional domain  $\Omega_{\alpha}$ . Our 42 goal is to computationally find the optimal shape for part of the boundary  $\partial\Omega_{lpha}$  such 43 that a given objective function  $J_o$  is optimized. We represent the part of the boundary 44 by a smooth function  $\alpha(x)$  determined by a set of parameters  $\mathbf{a} = (a_1, a_2, \dots, a_n)$ . By 45 changing the shape defined by  $\alpha(x)$ , one can optimize certain properties of the flow. 46 In this paper, we focus on the minimization of the energy dissipation in the whole 47 flow field and use the integral of the squared energy deformation as the objective 48 function [6]

$$\min_{\mathbf{u},\alpha} J_o(\mathbf{u},\alpha) = 2\mu \int_{\Omega_{\alpha}} \varepsilon(\mathbf{u}) \cdot \varepsilon(\mathbf{u}) dx dy + \frac{\beta}{2} \int_{I} (\alpha'')^2 dx$$
subject to
$$\begin{cases}
-\mu \Delta \mathbf{u} + \mathbf{u} \cdot \nabla \mathbf{u} + \nabla p = \mathbf{f} & \text{in } \Omega_{\alpha}, \\
\nabla \cdot \mathbf{u} = 0 & \text{in } \Omega_{\alpha}, \\
\mathbf{u} = \mathbf{g} & \text{on } \Gamma_{inlet}, \\
\mathbf{u} = \mathbf{0} & \text{on } \Gamma_{wall},
\end{cases}$$

$$\mu \frac{\partial \mathbf{u}}{\partial \mathbf{n}} - p \cdot \mathbf{n} = \mathbf{0} & \text{on } \Gamma_{outlet}, \\
\alpha(a) = z_1, \quad \alpha(b) = z_2,$$
(1)

where  $\mathbf{u} = (u, v)$  and p represent the velocity and pressure,  $\mathbf{n}$  is the outward unit 50 normal vector on  $\partial \Omega_{\alpha}$  and  $\mu$  is the kinematic viscosity.  $\Gamma_{inlet}$ ,  $\Gamma_{outlet}$  and  $\Gamma_{wall}$  rep- 51 resent the inlet, outlet and wall boundaries, respectively; see Fig. 1. f is the given 52 body force and **g** is the given velocity at the inlet  $\Gamma_{inlet}$ .  $\varepsilon(\mathbf{u}) = \frac{1}{2}(\nabla \mathbf{u} + (\nabla \mathbf{u})^{\mathbf{T}})$  is the 53 deformation tensor for the flow velocity **u** and  $\beta$  is a nonnegative constant. I = [a, b] 54 is an interval in which the shape function  $\alpha(x)$  is defined. In the constraints, the first 55 five equations are the Navier-Stokes equations and boundary conditions and the last 56 two equations indicate that the optimized boundary should be connected to the rest 57 of the boundary and  $z_1$  and  $z_2$  are two given constants. The last term in the objective 58 function is a regularization term providing the regularity of  $\partial \Omega_{\alpha}$ .

The optimization problem (1) is discretized with a LBB-stable (Ladyzhenskaya- 60 Babuška-Brezzi)  $Q_2 - Q_1$  finite element method. Since the computational domain of 61 the problem changes during the optimization process, the mesh needs to be modified 62 following the computational domain. Generally speaking, there are two strategies to 63 modify the mesh. One is mesh reconstruction which often guarantees a good new 64 mesh but is computationally expensive. The other strategy is moving mesh which 65 is cheaper but the deformed mesh may become ill-conditioned when the boundary 66 variation is large. In our test case the boundary variations are not very large, so we 67

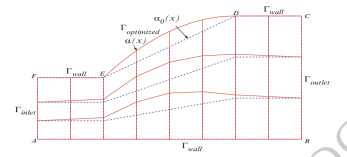


Fig. 1. The initial domain  $\Omega_{\alpha_0}$  (dashed line) and deformed domain  $\Omega_{\alpha}$  (solid line) over a simple mesh. The boundary  $\Gamma_{optimized}$  (ED) denotes the part of the boundary whose shape is computed by the optimization process

use the latter strategy. The moving of the mesh is simply described by Laplace's 68 equations.

$$\begin{cases}
-\Delta \delta_{\mathbf{x}} = \mathbf{0} & \text{in } \Omega_{\alpha_0}, \\
\delta_{\mathbf{x}} = \mathbf{g}_{\alpha} & \text{on } \partial \Omega_{\alpha_0},
\end{cases}$$
(2)

where  $\delta_{\bf x}$  is the mesh displacement and  ${\bf g}_{\alpha}=(g_{\alpha}^x,g_{\alpha}^y)$  is the displacement on the 70 boundary determined by  $\alpha(x)$ . Note that  ${\bf g}_{\alpha}$  is obtained automatically during the 71 iterative solution process. For example, in Fig. 1,  $g_{\alpha}^x=0$  and  $g_{\alpha}^y=\alpha(x)-\alpha_0(x)$ . 72 The Eqs. (2) are discretized with a  $Q_2$  finite element method. The discretized shape 73 optimization problem is given as follows

$$\min_{\mathbf{u}, \mathbf{a}, \delta_{\mathbf{x}}} J_{o}(\mathbf{u}, \mathbf{a}, \delta_{\mathbf{x}}) = \mu \mathbf{u}^{\mathbf{T}} \mathbf{J} \mathbf{u} + \frac{\beta}{2} \mathbf{J}_{\alpha}$$
subject to
$$\begin{cases}
\mathbf{K} \mathbf{u} + \mathbf{B}(\mathbf{u}) \mathbf{u} - \mathbf{Q} \mathbf{p} = \mathbf{F}_{\mathbf{f}} + \mathbf{F}_{\mathbf{u}}, \\
\mathbf{Q}^{\mathbf{T}} \mathbf{u} = \mathbf{0}, \\
\mathbf{D} \delta_{\mathbf{x}} = \mathbf{F}_{\mathbf{x}}, \\
\mathbf{A}_{\mathbf{a}} = \mathbf{F}_{\mathbf{a}}.
\end{cases} \tag{3}$$

Here  $\mathbf{F_f}$  refers to the discretized body force,  $\mathbf{F_u}$  and  $\mathbf{F_x}$  refer to the Dirichlet boundary 75 condition for  $\mathbf{u}$  and  $\delta_{\mathbf{x}}$ , respectively, and  $\mathbf{A_a}$  and  $\mathbf{F_a}$  are the geometric constrains. Note 76 that  $\mathbf{K}$ ,  $\mathbf{B}(\mathbf{u})$ ,  $\mathbf{Q}$  and  $\mathbf{J}$  depend on the grid displacement  $\delta_{\mathbf{x}}$ , while  $\mathbf{D}$  is independent of 77  $\delta_{\mathbf{x}}$ . Here  $\delta_{\mathbf{x}}$  is treated as an optimization variable and the moving mesh equations are 78 viewed as constraints of the optimization problem which are solved simultaneously 79 with the other equations.

# 3 One-Shot Lagrange-Newton-Krylov-Schwarz Methods

We use a Lagrange multiplier method to transform the optimization problem (3) 82 to a nonlinear system G(X) = 0 which is solved by an inexact Newton method. 83

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Given an initial guess  $X^0$ , at each iteration,  $k = 0, 1, \dots$ , we use a GMRES method 84 to approximately solve the preconditioned system 85

$$\mathbf{H}^{k}(\mathbf{M}^{k})^{-1}(\mathbf{M}^{k}\mathbf{d}^{k}) = -\mathbf{G}^{k},\tag{4}$$

to find a search direction  $\mathbf{d}^k$ , where  $\mathbf{H}^k = \nabla_X \mathbf{G}(\mathbf{X}^k)$  is the Jacobian matrix of the 86 nonlinear function,  $\mathbf{G}^k = \mathbf{G}(\mathbf{X}^k)$  and  $(\mathbf{M}^k)^{-1}$  is an additive Schwarz preconditioner 87 [11] defined as

$$(\mathbf{M}^k)^{-1} = \sum_{l=1}^{N_p} (R_l^{\delta})^{\mathbf{T}} (\mathbf{H}_l^k)^{-1} R_l^{\delta},$$
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where  $\mathbf{H}_l^k = R_l^{\delta} \mathbf{H}^k (R_l^{\delta})^{\mathsf{T}}$ ,  $R_l^{\delta}$  is a restriction operator from  $\Omega_{\alpha}$  to the overlapping 90 subdomain,  $\delta$  is the size of the overlap which is understood in terms of the number 91 of elements; i.e.,  $\delta = 8$  means the overlapping size is 8 layers of elements, and  $N_p$  92 is the number of subdomains which is equal to np in this paper. After approximately 93 solving (4), the new approximate solution is defined as  $\mathbf{X}^{k+1} = \mathbf{X}^k + \tau^k \mathbf{d}^k$ , and the 94 step length  $\tau^k$  is selected by a cubic line search.

# **4 Numerical Experiments**

The algorithm introduced in the previous sections is applicable to general shape optimization problems governed by incompressible Navier-Stokes equations. Here we study an application of the algorithm for the incoming part of a simplified artery bypass problem [2] as shown in Fig. 2. Our solver is implemented using PETSc [3]. All computations are performed on an IBM BlueGene/L supercomputer at the National Center for Atmospheric Research. Unstructured meshes, which are generated with CUBIT and partitioned with ParMETIS, are used in this paper.

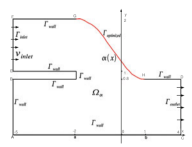
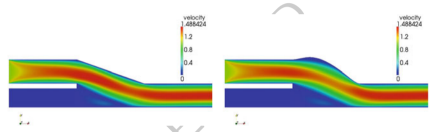


Fig. 2. The incoming part of a simplified bypass model; The *red* boundary  $\Gamma_{optimized}$  denotes the part of the boundary whose shape is to be determined by the optimization process

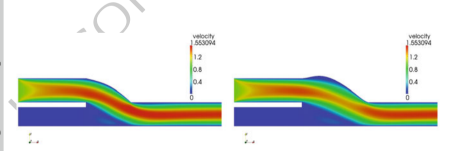
This is the incoming part of a bypass: www.reshealth.org/images/greystone/em\delimiter"026E30F\_2405.gif

Without the blockage, the flow is supposed to go from AB to CD, but now we assume that AB is blocked and the flow has to go through EF. For simplicity, we let the thickness EF be fixed and the body force  $\mathbf{f} = \mathbf{0}$  in the Navier-Stokes equations. 106 The shape of the bypass is determined by the curve GH as in Fig. 2. The boundary 107 conditions on the inlet  $\Gamma_{intlet}$  are chosen as a constant  $v_{in}$ , no-slip boundary conditions are used on the walls  $\Gamma_{wall}$ . On the outlet section  $\Gamma_{outlet}$ , the free-stress boundary 109 conditions are imposed; see (1). We use a polynomial  $\alpha(x) = \sum_{i=1}^p a_i x^i$  with p = 7 to 110 represent the part of the boundary that needs to be optimized. Other shape functions 111 can be used, but here we simply follow [1]. The goal is to compute the coefficients 112  $\mathbf{a} = (a_1, \dots, a_p)$ , such that the energy loss is minimized.

In all experiments, we use a hand-coded Jacobian matrix. The Jacobian system 114 in each Newton step is solved by a right-preconditioned restarted GMRES with an 115 absolute tolerance of  $10^{-10}$ , a relative tolerance of  $10^{-3}$ , and a restart at 100. We stop 116 the Newton iteration when the nonlinear residual is decreased by a factor of  $10^{-6}$ .



**Fig. 3.** Velocity distribution of the initial (*left*) and optimal shapes (*right*). The initial shape is given by a *straight line*.  $\beta = 0.01$  and Re = 100

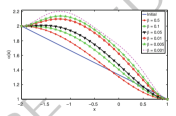


**Fig. 4.** Velocity distribution of the initial (*left*) and optimal shapes (*right*). The initial shape is given as  $\alpha(x) = 0.4 + 0.45x^2 + 0.15x^3$ .  $\beta = 0.01$  and Re = 100

In the first test case, we set the Reynolds number  $Re = \frac{Lv_{in}}{\mu}$  to 100, where  $L = \frac{117}{118}$  1.0 cm is the artery diameter,  $v_{in} = 1.0$  cm/s is the inlet velocity and  $\mu = 0.01$  cm<sup>2</sup>/s. 119

We solve the problem on a mesh with about 18,000 elements.  $\beta=0.01$  and the degrees of freedom (DOF) is 589,652. The initial shape is given by a straight line, and 121 Fig. 3 shows the velocity distribution of the initial (left) and optimal shapes (right). 122 The energy dissipation of the optimized shape is reduced by about 5.13 % compared 123 to the initial shape. Figure 4 is the velocity distribution of another initial shape (left) 124 which is given as  $\alpha(x)=0.4+0.45x^2+0.15x^3$  and the corresponding optimal shape 125 (right). The reduction of the energy dissipation of this case is about 11.96 %. Figures 3 and 4 show that we can obtain nearly the same optimal shape from different 127 initial shapes.

In the test case showed in Fig. 3, if we add a small inlet velocity at the boundary 129 AB, which is equal to that the blood flow is not totally blocked, the computed optimal 130 shape would be different from what is shown in Fig. 3. If we move the boundary 131 AB towards CD (A from (-5,0) to (-3,0) and B from (-5,0.8) to (-3,0.8)), the 132 optimal shape is nearly the same as Fig. 3 since the flow in the "dead area" doesn't 133 impact much of the optimal solution.



**Fig. 5.** The initial shape and optimal shapes with different values of parameter  $\beta$ . DOF = 589,652 and Re = 100

The regularization parameter  $\beta$  in the objective function is very important for shape optimization problems. From Table 1 we see that reducing  $\beta$  can increase the reduction of the energy dissipation ("Init.", "Opt." and "Reduction" are the initial, 137 optimized and reduction of the energy dissipation in the table), but the number of 138 Newton (Newton) and the average number of GMRES iterations per Newton (GM-139 RES) and the total compute time in seconds (Time) increase, which means that the 140 nonlinear algebraic system is harder to solve when  $\beta$  is small. This is because the 141 boundary of  $\Omega_{\alpha}$  is more flexible and may become irregular when  $\beta$  is too small. Figure 5 shows the initial shape and the optimized shapes obtained with different values of  $\beta$ . From this figure we see that  $\beta$  controls the boundary deformation.

To show the parallel scalability of the algorithm, two meshes with DOF = 145589,652 and DOF = 928,572 are considered. The strong scalability of our algorithm 146 is good; see Fig. 6 and Table 2, which show that the speedup is almost linear when 147 np is small. As expected in one-level Schwarz methods, the preconditioner becomes 148 worse as the number of subdomains increases. 149

Table 3 shows some results for different *Re*. Judging from the increase of the 150 number of linear and nonlinear iterations, it is clear that the problem becomes harder 151

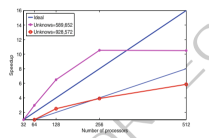
One-Shot Domain Decomposition Methods for Shape Optimization Problems

Table 1. Effect of the	parameter $\beta$ . DOF	= 589,652, Re = 100.
------------------------	-------------------------	----------------------

В	Newton	CMDES	Time	Energy Dissipation		
P	i New toll	GMRES Time		Init.	Opt.	Reduction
0.05	4					4.27%
0.01	5	441.40				5.13%
0.005	5	439.00	599.77	1.17	1.10	5.98%
0.001	6	510.67	747.78	1.17	1.10	5.98%

**Table 2.** Parallel scalability for two different size grids.  $\beta = 0.1$ , overlap = 6 and Re = 100.

		F=589,		DOF = 928,572			
np	Newton	<b>GMRES</b>	Time	Newton	GMRES	Time	
32	4	124.50	2959.73	_	-	)—	
64	4	179.25	980.48	4	146.50	2121.52	
128	4	346.75	455.69	4	330.00	844.62	
256	4	533.25	280.96	4	520.75	541.97	
512	4	917.50	282.07	4	861.00	361.08	



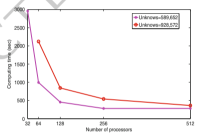


Fig. 6. The speedup and the total compute time for two different mesh sizes. Re = 100

as we increase the Re. On the other hand, we achieve higher percentage of reduction  $_{152}$  of energy dissipation in the harder to solve situations.

**Table 3.** The impact of *Re*.  $\beta = 0.1$ , *overlap* = 8, *DOF* = 589,652, np = 128.

D	NT /	CMDEC	Tr'			ssipation
Re	Newton	GMRES	11me	Init.	Opt.	Reduction
100	4	346.75	456.83	1.17	1.13	3.42%
200	4	372.00	470.16	0.65	0.62	4.62%
300	6					6.05%
600	7	721.71	1035.84	7.43	6.97	6.19%

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#### 5 Conclusions and Future Work

We developed a parallel one-shot LNKSz for two-dimensional shape optimization 155 problems governed by incompressible Navier-Stokes equations. We tested the algorithms for an artery bypass design problem with more than 900,000 DOF and up to 157 512 processors. The numerical results show that our method is quite robust with re- 158 spect to the Re and the regularization parameter. The strong scalability is almost ideal 159 when np is not too large. In the future, we plan to study some multilevel Schwarz 160 methods which may improve the scalability when np is large.

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# A Schur Complement Method for Compressible **Navier-Stokes Equations**

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Summary. Domain decomposition methods were first developed for elliptic problems, taking 9 advantage of the strong regularity of their solutions. In the last two decades, many investiga- 10 tions have been devoted to improve the performance of these methods for elliptic and parabolic 11 problems. The situation is less clear for hyperbolic problems with possible singular solutions. 12 In this paper, we will discuss a nonoverlapping domain decomposition method for nonlinear 13 hyperbolic problems. We use the finite volume method and an implicit version of the Roe 14 approximate Riemann solver, and propose a new interface variable inspired by Dolean and 15 Lanteri [1]. The new variable makes the Schur complement approach simpler and allows the treatment of diffusion terms. Numerical results for the compressible Navier-Stokes equations 17 in various 2D and 3D configurations such as the Sod shock tube problem or the lid driven 18 cavity problem show that our method is robust and efficient. Comparisons of performances on 19 parallel computers with up to 512 processors are also reported.

1 Introduction 21

When solving a nonlinear partial differential equation by an implicit scheme, one 22 classically ends by solving a nonlinear algebraic system using a Newton method. 23 At each step of this method we have to solve a linear system  $\mathcal{A}(U^k)U^{k+1} = b(U^k)$ . 24 This task is computationally expensive in particular since the matrix  $\mathscr{A}$  is usually 25 non-symmetric and very ill-conditioned. It is therefore necessary to find an efficient 26 preconditioner.

When the size of the system is large (as in the case of 3D computations), the par- 28 allel solution on multiple processors is essential to obtain reasonable computation 29 times. Currently in the thermal hydraulic code, FLICA-OVAP (see [2]), the matrix 30  $\mathcal{A}$  and the right hand side b are stored on multiple processors and the system is 31 solved in parallel with a Krylov solver (classical incomplete factorization). Unfor- 32 tunately, the parallel preconditioners of FLICA-OVAP only perform well on a few 33 processors. In contrast, if we want to increase the number of processors these par- 34 allel preconditioners perform poorly. Tests were run on different test cases and led 35

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us to conclude that it is often better not to use these parallel preconditioners, espe- 36 cially for 3D problems. This strategy does not make an optimal use of the available 37 computational power. Hence we seek for more efficient methods to distribute the 38 computations. We study and use a domain decomposition method as an alternative 39 to the classical distribution.

The paper is organized as follows. In Sects. 2 and 3, we present the mathematical 41 model and its numerical schemes. In Sect. 4, we first review the domain decomposition method proposed by Dolean and Lanteri [1] based on a Schwarz algorithm. We 43 then introduce a new interface variable which makes the Schur complement approach 44 simpler and allows for the treatment of diffusion terms. Section 5 presents a set of 45 numerical experiments to validate our method, compares it with that of [1] concern- 46 ing the robustness and efficiency and presents the scalability and the performance of 47 different preconditioners.

#### 2 Mathematical Model

The simplest model of FLICA-OVAP consists of the following three balance laws 50 for the mass, the momentum and the energy:

$$\begin{cases}
\frac{\partial \rho}{\partial t} + \nabla \cdot \mathbf{q} &= 0 \\
\frac{\partial \mathbf{q}}{\partial t} + \nabla \cdot \left( \mathbf{q} \otimes \frac{\mathbf{q}}{\rho} + p \mathbb{I}_d \right) - \nu \Delta \left( \frac{\mathbf{q}}{\rho} \right) = 0 \\
\frac{\partial (\rho E)}{\partial t} + \nabla \cdot \left[ (\rho E + p) \frac{\mathbf{q}}{\rho} \right] - \lambda \Delta T &= 0
\end{cases} \tag{1}$$

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where  $\rho$  is the density,  ${\bf v}$  the velocity,  ${\bf q}=\rho{\bf v}$  the momentum, p the pressure,  $\rho e$  the 52 internal energy,  $\rho E = \rho e + \frac{||\mathbf{q}||^2}{2\rho}$  the total energy, T the absolute temperature, v the 53 viscosity and  $\lambda$  the thermal conductivity. We close the system (1) by the ideal gas 54 law  $p = (\gamma - 1)\rho e$ . For the sake of simplicity, we consider constant viscosity and 55 conductivity, and neglect the contribution of viscous forces in the energy equation. By denoting  $U = (\rho, \mathbf{q}, \rho E)^t$  the vector of conserved variables, the Navier–Stokes 57 system (1) can be written as a nonlinear system of conservation laws:

$$\frac{\partial U}{\partial t} + \nabla \cdot (\mathscr{F}^{conv}(U)) + \nabla \cdot \left(\mathscr{F}^{diff}(U)\right) = 0, \tag{2}$$

where 
$$\mathscr{F}^{conv}(U) = \begin{pmatrix} \mathbf{q} \\ \mathbf{q} \otimes \frac{\mathbf{q}}{\rho} + p \mathbb{I}_d \\ (\rho E + p) \frac{\mathbf{q}}{\rho} \end{pmatrix}, \, \mathscr{F}^{diff}(U) = \begin{pmatrix} 0 \\ -\nu \nabla (\frac{\mathbf{q}}{\rho}) \\ -\lambda \nabla T \end{pmatrix}.$$
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#### 3 Numerical Method

The conservation form (2) allows for the definition of weak solutions, which can 61 be discontinuous ones. Discontinuous solutions such as shock waves are of great 62 importance in transient calculations. In order to correctly capture shock waves, one 63 needs a robust, low diffusive conservative scheme. The finite volume framework is 64 the most appropriate setup to write discrete equations that express the conservation 65 laws at each cell (see [3]).

We decompose the computational domain into N disjoint cells  $C_i$  with volume 67  $v_i$ . Two neighboring cells  $C_i$  and  $C_j$  have a common boundary  $\partial C_{ij}$  with area  $s_{ij}$ . We 68 denote N(i) the set of neighbors of a given cell  $C_i$  and  $\mathbf{n}_{ij}$  the exterior unit normal 69 vector of  $\partial C_{ij}$ . Integrating the system (2) over  $C_i$  and setting  $U_i(t) = \frac{1}{v_i} \int_{C_i} U(x,t) dx$  70 and  $U_i^n = U_i(n\Delta t)$ , the discretized equations can be written:

$$\frac{U_i^{n+1} - U_i^n}{\Delta t} + \sum_{j \in N(i)} \frac{s_{ij}}{v_i} \left( \overrightarrow{\Phi}_{ij}^{conv} + \overrightarrow{\Phi}_{ij}^{diff} \right) = 0.$$
 (3)

with: 
$$\overrightarrow{\Phi}_{ij}^{conv} = \frac{1}{s_{ij}} \int_{\partial C_{ij}} \mathscr{F}^{conv}(U^{n+1}) \cdot \mathbf{n}_{ij} ds$$
,  $\overrightarrow{\Phi}_{ij}^{diff} = \frac{1}{s_{ij}} \int_{\partial C_{ij}} \mathscr{F}^{diff}(U^{n+1}) \cdot \mathbf{n}_{ij} ds$ .

To approximate the convection numerical flux  $\overrightarrow{\Phi}_{ij}^{conv}$  we solve an approximate 73 Riemann problem at the interface  $\partial C_{ij}$ . Using the Roe local linearisation of the fluxes 74 [4], we obtain the following formula:

$$\overrightarrow{\Phi}_{ij}^{conv} = \frac{\mathscr{F}^{conv}(U_i^{n+1}) + \mathscr{F}^{conv}(U_j^{n+1})}{2} \cdot \mathbf{n}_{ij} - \mathscr{D}(U_i^{n+1}, U_j^{n+1}) \frac{U_j^{n+1} - U_i^{n+1}}{2} \quad (4)$$

$$= \mathscr{F}^{conv}(U_i^{n+1}) \mathbf{n}_{ij} + A^{-}(U_i^{n+1}, U_j^{n+1}) (U_j^{n+1} - U_i^{n+1}), \quad (5)$$

where  $\mathscr{D}$  is an upwinding matrix,  $A(U_i^{n+1}, U_j^{n+1})$  the Roe matrix and  $A^{\pm} = \frac{A \pm \mathscr{D}}{2}$ . 76 The choice  $\mathscr{D} = 0$  gives the centered scheme, whereas  $\mathscr{D} = |A|$  gives the upwind 77 scheme. For the Euler equations, we can build  $A(U_i^{n+1}, U_j^{n+1})$  explicitly using the 78 Roe averaged state (see [3]).

The diffusion numerical flux  $\overrightarrow{\Phi}_{ij}^{diff}$  is approximated on structured meshes using 80 the formula:

$$\overrightarrow{\Phi}_{ij}^{diff} = D(\frac{U_i^{n+1} + U_j^{n+1}}{2})(U_j^{n+1} - U_i^{n+1})$$
(6)

with the matrix  $D(U) = \begin{pmatrix} 0 & \mathbf{0} & 0 \\ \frac{v\mathbf{q}}{\rho^2} & \frac{-v}{\rho} \mathbb{I}_d & 0 \\ \frac{\lambda}{c_v} \left( \frac{c_v T}{\rho} - \frac{||\mathbf{q}||^2}{2\rho^3} \right) & \frac{\mathbf{q}^t \lambda}{\rho^2 c_v} & -\frac{\lambda}{c_v \rho} \end{pmatrix}$ , where  $c_v$  is the heat 82 capacity at constant volume.

3.1 Newton Scheme

Finally, since  $\sum_{j\in N(i)} \mathscr{F}^{conv}(U_i^{n+1}).\mathbf{n}_{ij} = 0$ , using (5) and (6) the Eq. (3) of the nu-

$$\frac{U_i^{n+1} - U_i^n}{\Delta t} + \sum_{j \in N(i)} \frac{s_{ij}}{v_i} \{ (A^- + D)(U_i^{n+1}, U_j^{n+1}) \} (U_j^{n+1} - U_i^{n+1}) = 0.$$
 (7)

The system (7) is nonlinear, hence we use the following Newton iterative method to 87 obtain the required solutions: 88

$$\frac{\delta U_i^{k+1}}{\Delta t} + \sum_{j \in N(i)} \frac{s_{ij}}{v_i} \left[ (A^- + D)(U_i^k, U_j^k) \right] \left( \delta U_j^{k+1} - \delta U_i^{k+1} \right) 
= -\frac{U_i^k - U_i^n}{\Delta t} - \sum_{j \in N(i)} \frac{s_{ij}}{v_i} \left[ (A^- + D)(U_i^k, U_j^k) \right] (U_j^k - U_i^k),$$
(8)

where  $\delta U_i^{k+1} = U_i^{k+1} - U_i^k$  is the variation of the *k*-th iterate that approximates the solution at time n+1.

## 4 Domain Decomposition Method

The principle of the domain decomposition method by Schur complement is to decompose the global problem into independent subproblems solved on each processor.

More precisely, if we want to solve the problem:

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$$\begin{cases} \frac{\partial U}{\partial t} + \nabla \cdot \mathscr{F}(U) = 0 \text{ in } \Omega \\ BU = g & \text{on } \partial \Omega \end{cases}$$
 (9)

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on a partition of the original domain  $\Omega = \bigcup_{I=1}^K \Omega_I$ , defining  $U_I$  as the restriction of the 95 solution U in the subdomain  $\Omega_I$ , the algorithm of the domain decomposition method 96 is then written as: 97

$$\begin{cases} \frac{\partial U_I}{\partial t} + \nabla \cdot \mathscr{F}(U_I) = 0 \text{ in } \Omega \\ BU_I = g & \text{on } \partial \Omega \cap \partial \Omega_I \\ C_I U_I = C_I U_J & \text{on } \partial \Omega_I \cap \partial \Omega_j \end{cases}$$
(10)

where  $C_I$  is an interface operator which we will clarify later.

#### 4.1 Dolean and Lanteri Interface Variable

In the article [1], in order to make the subsystem (10) solution independent, Dolean to et al introduced a redundant variable  $\Phi^{DL}_{ij}$  at the domain interface between two cells to and  $j:\Phi^{DL}_{ij}=A^+_{Roe,\mathbf{n}_{i,j}}U_i-A^-_{Roe,\mathbf{n}_{i,j}}U_j$  and then defined the orthogonal projectors to the eigenvectors subspaces such that

$$P^{-}(U_{i},U_{j})\delta\phi_{ij}^{Do} = A_{Roe,\mathbf{n}_{i,j}}^{-}\delta U_{j}^{k+1}, P^{+}(U_{i},U_{j})\delta\phi_{ij}^{Do} = -A_{Roe,\mathbf{n}_{i,j}}^{+}\delta U_{i}^{k+1}$$
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This strategy can only be applied to the Euler equations (Eq. (2) with no viscosity and heat conductivity terms) using the upwind scheme. In order to include diffusion terms in the model and to use various schemes, we introduce a new interface variable  $\Phi_{ij}$  at the domain interface between two cells i and j:

$$\Phi_{ij} = U_j - U_i \tag{11}$$

#### 4.2 A New Interface Variable

In the case where the cell i of the subdomain I is at the boundary and has to communicate with the neighboring subdomains, we can rewrite the system (8) as:

$$\begin{split} \frac{\delta U_i^{k+1}}{\Delta t} & + \sum_{j \in I, j \in N(i)} \frac{s_{ij}}{v_i} \left[ (A^- + D)(U_i^k, U_j^k) \right] \left( \delta U_j^{k+1} - \delta U_i^{k+1} \right) \\ & = -\frac{U_i^k - U_i^n}{\Delta t} - \sum_{j \in N(i)} \frac{s_{ij}}{v_i} \left[ (A^- + D)(U_i^k, U_j^k) \right] (U_j^k - U_i^k) \\ & - \sum_{j \notin I, j \in N(i)} \left[ (A^- + D)(U_i^k, U_j^k) \right] \delta \phi_{ij} \end{split}$$

By defining  $\mathcal{U}_I = (U_1, \dots, U_m)^t$  the unknown vector of the subdomain I and

$$\delta\phi_{IJ} = (\delta\phi_{ij})_{i \in I, j \in J, j \in N(i)}$$
(12)

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and by denoting  $P = A^- + D$ , we can write the linear system as:

$$\mathscr{A}(\mathscr{U}_I^k)\delta\mathscr{U}_I^{k+1} = b_I(\mathscr{U}^n, \mathscr{U}^k) - \sum_{I \in N(I)} P(\mathscr{U}_I^k, \mathscr{U}_J^k)\delta\phi_{IJ}$$
(13)

By taking into account Eqs. (11)–(13), we can build an extended system that distinguishes the internal unknowns from the interface ones:

$$\begin{pmatrix}
\mathscr{A}_{1} & 0 & \dots & | P_{1} \\
0 & \mathscr{A}_{2} & 0 & \dots & | P_{2} \\
\dots & \dots & \dots & \dots \\
\frac{0}{M_{1}} & \dots & \dots & M_{N} & \mathbb{I}
\end{pmatrix}
\begin{pmatrix}
\mathscr{S}\mathscr{U}_{1} \\
\mathscr{S}\mathscr{U}_{2} \\
\dots \\
\mathscr{S}\mathscr{U}_{N} \\
\mathscr{S}\Phi
\end{pmatrix} = \begin{pmatrix}
b_{1} \\
b_{2} \\
\dots \\
b_{N} \\
b_{\phi}
\end{pmatrix}$$
(14)

where  $\mathscr{A}_I$  is the matrix that couples the unknowns associated with internal cells of  $\Omega_I$  whereas  $M_I$  enables us to build  $\delta \Phi$ , the interface unknown on all coupling subdomain interfaces, from the  $\delta U_I$ . The internal unknowns can be eliminated in favor of the interface ones to yield the following interface system:

$$S\delta\phi = b_{\phi} \tag{15}$$

with 120

$$(S\delta\phi)_{IJ} = \delta\phi_{IJ} + M_{IJ}\mathscr{A}_I^{-1} \sum_{K \in N(I)} P_{IK}\delta\phi_{IK} + M_{JI}\mathscr{A}_J^{-1} \sum_{K \in N(J)} P_{JK}\delta\phi_{JK}$$
$$(b_\phi)_{IJ} = M_{IJ}A_I^{-1}b_I + M_{JI}A_J^{-1}b_J$$

The Eq. (15) can be solved by, e.g., GMRES, BICGStab, or the Richardson methods. 121

**Page 577** 

### 5 Numerical Results

#### 5.1 Validation 123

Figures 1 and 2 present the profile of the pressure after 10 time steps using the upwind 124 scheme with CFL = 10 for the Euler equations. Our initial state is a pressurized ball 125 at the center of a closed box and for t > 0 there are waves which propagate and reflect 126 all over the box. The gas expands in the box and we can see the shock waves and the 127 rarefaction waves. The solution is solved on a cartesian mesh of  $200 \times 200$  cells.

Figures 3 and 4 show the streamlines of the steady state obtained using centered 129 scheme to solve a lid driven cavity flow at Reynolds number 400 on a cartesian 130  $50 \times 50$  mesh. The lid speed is 1 m/s, the maximum Mach number of the flow is 131 0.008. According to these results, we obtain the same solutions by using single or

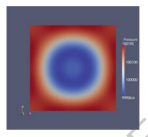


Fig. 1. Profile of the pressure at time step 10 on one processor

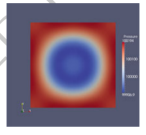
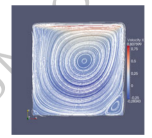
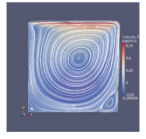


Fig. 2. Profile of the pressure at time step 10 on four processors



**Fig. 3.** Streamlines of  $V_x$  on one processor



**Fig. 4.** Streamlines of  $V_x$  on four processors

multiple domains.

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#### 5.2 Scalability

We now study the robustness and the scalability of our numerical method using the 135 same test as presented in Sect. 5.1. In Figs. 5 and 6, we compare the parallel efficiency 136 of different preconditioners on 2D and 3D computations and with two and four processors. We see that without the preconditioner the solver is scalable. However, when

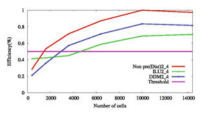


Fig. 5. Parallel efficiency for 2D Lid driven cavity

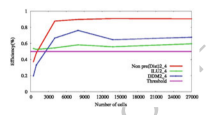


Fig. 6. Parallel efficiency for 3D Lid driven cavity

we use the Incomplete LU preconditioner, the scalability is not optimal especially for 139 3D problems. Our method proves better than ILU when we increase the number of 140 cells in each subdomain. In Fig. 7, we compare the robustness of different methods

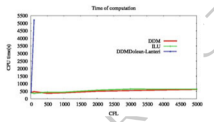


Fig. 7. Comparisons of parallelism in 3D Detonation, global mesh =  $50 \times 50 \times 50$ 

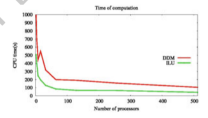


Fig. 8. Time of computation, 1 time step, global mesh =  $96 \times 96 \times 96$ 

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using the detonation problem. This problem is solved on a catersian  $50 \times 50 \times 50$ cell mesh on two processors. The computation time of Dolean and Lanteri method 143 increases rapidly because it needs many Newton iterations for convergence at each 144 time step. In Fig. 8, we compare the scalability of the ILU preconditioner and of our 145 method using the lid driven cavity problem solved on a global catersian  $96 \times 96 \times 96$ cell mesh. The computation time of the domain decomposition method is higher than 147 that of the ILU preconditioner due to the large number of Schur complement itera- 148 tions.

6 Conclusion 150

We have presented a new interface variable which allows for the treatment of diffusion terms and the use of various numerical schemes. We also compared the efficiency and the scalability of our method with the classical distributed computations 153

and the method of Dolean and al. Our approach seems promising but we still need to find an efficient preconditioner for the Schur complement in order to reduce its computational time.	
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# Numerical Study of the Almost Nested Case in a **Multilevel Method Based on Non-nested Meshes**

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**Summary.** Partial differential equations in complex domains are very flexibly discretized by 7 finite elements with unstructured meshes. For such problems, the challenging task to construct 8 coarse level spaces for efficient multilevel preconditioners can in many cases be solved by a 9 semi-geometric approach, which is based on a hierarchy of non-nested meshes. In this paper, 10 we investigate the connection between the resulting semi-geometric multigrid methods and the truly geometric variant more closely. This is done by considering a sufficiently simple computational domain and treating the geometric multigrid method as a special case in a family of 13 almost nested settings. We study perturbations of the meshes and analyze how efficiency and 14 robustness depend on a truncation of the interlevel transfer. This gives a precise idea of which 15 results can be achieved in the general unstructured case.

1 Introduction

This paper is about multilevel methods for an efficient solution of partial differential 18 equations in complicated domains. Our particular purpose is to provide additional 19 insight into the design of coarse spaces in case of unstructured finite element meshes. 20 We study an approach of semi-geometric preconditioning based on non-nested mesh 21 hierarchies motivated by Cai [2], Chan et al. [3, 4], Griebel and Schweitzer [6], 22 Toselli and Widlund [8], and Xu [9]. This is a concept with rather weak requirements 23 (yet still in a variational setting) compared with other geometry-based methods. The 24 main contribution of the present paper is a numerical study of the almost nested 25 case, which establishes a connection between the multilevel methods based on non- 26 nested meshes and the standard variant. Combined with our investigations of mesh 27 perturbations, this allows for the determination of a suitable truncation parameter for 28 the interlevel transfer. As a result, the efficiency of the completely nested case is in 29 large part retained.

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This section aims at a semi-geometric preconditioning framework. We introduce a 32 multiplicative multilevel preconditioner based on a hierarchy of non-nested meshes. 33 This is done in a way which allows for a powerful convergence analysis as well as 34 an efficient implementation. 35

Let  $\Omega \subset \mathbb{R}^d$  be a Lipschitz domain of dimension  $d \in \{2,3\}$ . For a right hand 36 side  $\mathscr{F} \in H^{-1}(\Omega)$  and a positive function  $\alpha \in L^{\infty}(\Omega)$  bounded away from zero, we 37 consider the variational model problem

$$u \in H_0^1(\Omega): \quad a(u,v) := (\alpha \nabla u, \nabla v)_{L^2(\Omega)} = \mathscr{F}(v), \quad \forall v \in H_0^1(\Omega).$$
 (1)

For a Galerkin discretization of problem (1), let  $(\mathscr{T}_\ell)_{\ell\in\mathbb{N}}$  be a family of *non-nested* 39 shape regular meshes of domains  $(\Omega_\ell)_{\ell\in\mathbb{N}}$ . We denote the set of nodes of  $\mathscr{T}_\ell$  by  $\mathscr{N}_\ell$  40 and abbreviate  $n_\ell:=|\mathscr{N}_\ell|$ . At each level  $\ell$ , we consider the space  $X_\ell$  of Lagrange 41 conforming finite elements of first order and denote its nodal basis as  $\Lambda_\ell=(\lambda_p^\ell)_{p\in\mathscr{N}_\ell}$  42 with  $\lambda_p^\ell(q)=\delta_{pq},\,p,q\in\mathscr{N}_\ell$ . For simplicity, we assume that  $\Omega_L=\Omega$  and  $X_L\subset H_0^1(\Omega)$  43 for a fixed finest level  $L\geq 2$ . In addition, let  $\Omega_\ell\supset\Omega$  for all  $\ell\in\{0,\ldots,L-1\}$ . The 44 basic idea how the setting can be chosen is exemplarily illustrated in Fig. 1 (left) for 45 an unstructured fine mesh with structured coarse meshes.

In the following, we consider an iterative method to efficiently solve the discrete 47 problem, namely the ill-conditioned equation 48

$$\mathbf{A}_L \mathbf{u}_L = \mathbf{F}_L \quad \text{in } \mathbb{R}^{n_L}.$$

Here,  $\mathbf{A}_L \in \mathbb{R}^{n_L \times n_L}$  is the stiffness matrix associated with  $X_L$ , i.e.,  $(\mathbf{A}_L)_{pq} := a(\lambda_p^L, \lambda_q^L)$  49 for  $p, q \in \mathcal{N}_L$ , and the right hand side  $\mathbf{F}_L \in \mathbb{R}^{n_L}$  is given by  $(\mathbf{F}_L)_p := \mathscr{F}(\lambda_p^L)$  for 50  $p \in \mathcal{N}_L$ .

For the construction of an appropriate coarse space hierarchy, let the spaces 52  $(X_\ell)_{\ell=0,\dots,L}$  be connected by the prolongation operators  $(\Pi_{\ell-1}^\ell)_{\ell=1,\dots,L}$ , namely 53

$$\Pi_{\ell-1}^{\ell}: X_{\ell-1} \to X_{\ell}, \quad \forall \ \ell \in \{1, \dots, L\}.$$

The choice of a concrete transfer concept generating a set of suitable linear operators 54  $(\Pi_{\ell-1}^{\ell})_{\ell=1,\dots,L}$  in practice is discussed in full detail in [5]. An example is nodal interpolation. Now, let  $V_L := X_L$ ; we emphasize that the fine space will not be touched in 56 the present framework. We construct a nested sequence of spaces  $(V_{\ell})_{\ell=0,\dots,L}$  via 57

$$V_{\ell} := \Pi_{L-1}^{L} \cdots \Pi_{\ell}^{\ell+1} X_{\ell}, \quad \forall \ \ell \in \{0, \dots, L-1\}.$$

The images of the compositions of the given operators determine the coarse spaces. 58 With the nodal bases  $(\Lambda_\ell)_{\ell=0,\dots,L}$ , matrix representations  $\Pi^\ell_{\ell-1} \in \mathbb{R}^{n_\ell \times n_{\ell-1}}$  of 59  $\Pi^\ell_{\ell-1}$  can be computed for  $\ell \in \{1,\dots,L\}$  via  $\Pi^\ell_{\ell-1} \mathbf{v} := \Phi^{-1}_\ell(\Pi^\ell_{\ell-1}\Phi_{\ell-1}(\mathbf{v}))$  for all 60  $\mathbf{v} \in \mathbb{R}^{n_{\ell-1}}$  with the coordinate isomorphisms  $\Phi_\ell : \mathbb{R}^{n_\ell} \to X_\ell$ . Assume that these matrices have full rank. Then, bases of  $(V_\ell)_{\ell=0,\dots,L-1}$  can recursively be defined by

The Almost Nested Case in a Multilevel Method Based on Non-nested Meshes

$$\widetilde{\lambda}_q^{\ell} := \sum_{p \in \mathcal{N}_{\ell+1}} (\boldsymbol{\Pi}_{\ell}^{\ell+1})_{pq} \widetilde{\lambda}_p^{\ell+1}, \quad \forall \ q \in \mathcal{N}_{\ell},$$

starting with  $\widetilde{\lambda}_q^L := \lambda_q^L$  for  $q \in \mathscr{N}_L$ . The new coordinate isomorphisms with respect 63 to the bases  $\widetilde{\Lambda}_\ell := (\widetilde{\lambda}_p^\ell)_{p \in \mathscr{N}_\ell}, \ell \in \{0,\dots,L\}$ , will be denoted by  $\widetilde{\Phi}_\ell : \mathbb{R}^{n_\ell} \to V_\ell$ . Moreover,  $\mathbf{M}_{\ell} \in \mathbb{R}^{n_{\ell} \times n_{\ell}}$  is the mass matrix with respect to  $\widetilde{\Lambda}_{\ell}$ , i.e.,  $(\mathbf{M}_{\ell})_{pq} := (\widetilde{\lambda}_{p}^{\ell}, \widetilde{\lambda}_{q}^{\ell})_{L^{2}(\Omega)}$ for  $p, q \in \mathcal{N}_{\ell}$ ,  $\ell \in \{0, \dots, L\}$ .

Note that the mapping  $\Pi_{\ell-1}^{\ell}$  between the given spaces  $X_{\ell-1}$  and  $X_{\ell}$  usually does 67 not act on  $V_{\ell-1}$  directly. Still, the matrix  $\Pi_{\ell-1}^{\ell}$  determines a linear transfer operator 68  $\widetilde{\Pi}_{\ell-1}^{\ell}: V_{\ell-1} \to V_{\ell}$  by

$$v \mapsto \widetilde{\Pi}_{\ell-1}^{\ell} v := \widetilde{\Phi}_{\ell}(\boldsymbol{\Pi}_{\ell-1}^{\ell} \widetilde{\Phi}_{\ell-1}^{-1}(v)), \quad \forall v \in V_{\ell-1}, \quad \forall \ell \in \{1, \dots, L\}.$$

One can easily see that  $\widetilde{\Pi}_{\ell-1}^{\ell}$  is the natural embedding because it interpolates the 70 respective basis exactly. Thus, we can regard the matrix  $\Pi_{\ell-1}^{\ell}$  as an algebraic representation of the natural embedding of  $V_{\ell-1}$  into  $V_{\ell}$ . Consequently, the  $L^2$ -projection 72 from  $V_{\ell}$  to  $V_{\ell-1}$  is represented by the matrix  $\mathbf{M}_{\ell-1}^{-1}(\mathbf{\Pi}_{\ell-1}^{\ell})^T\mathbf{M}_{\ell} \in \mathbb{R}^{n_{\ell-1}\times n_{\ell}}$ . This 73 holds true for any imaginable set of operators between the original non-nested spaces 74  $(X_{\ell})_{\ell=0,\dots,L}$ ; no special structure is required.

With this information we can summarize our efforts as follows. From the completely unrelated finite element spaces  $(X_{\ell})_{\ell=0,...,L}$  we have constructed a sequence of 77 nested spaces  $(V_\ell)_{\ell=0,...,L}$  such that the given prolongation operators  $(\Pi_{\ell-1}^\ell)_{\ell=1,...,L}$  78 induce the natural embeddings  $(V_{\ell-1} \hookrightarrow V_{\ell})_{\ell=1,\dots,L}$  by their matrix representations 79  $(\Pi_{\ell-1}^{\ell})_{\ell=1,...,L}$  with respect to the original bases  $(\Lambda_{\ell})_{\ell=0,...,L}$ . In particular, the coarse 80 level matrices for the nested spaces with the respective bases  $\tilde{\Lambda}_{\ell}$ , as customary in a 81 variational approach, can be written as 82

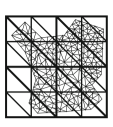
$$\boldsymbol{A}_{\ell-1} = (\boldsymbol{\Pi}_{\ell-1}^{\ell})^T \boldsymbol{A}_{\ell} \, \boldsymbol{\Pi}_{\ell-1}^{\ell} \in \mathbb{R}^{n_{\ell-1} \times n_{\ell-1}}, \quad \forall \ \ell \in \{1, \dots, L\}.$$

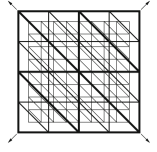
If  $A_L$  is symmetric positive definite and if  $\Pi_{\ell-1}^{\ell}$  has full rank for all  $\ell \in \{1, ..., L\}$ , 83 the respective coarse level matrices  $(\mathbf{A}_{\ell})_{\ell=0,\dots,L-1}$  are symmetric positive definite, 84 too. Note that the bandwidth of the coarse matrices depends on the transfer concept 85 employed to obtain the prolongation operators.

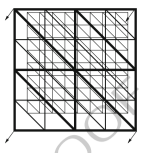
The multiplicative Schwarz method studied in this paper is the symmetric multi- 87 grid  $\mathscr{V}$ -cycle in the novel space hierarchy  $(V_{\ell})_{\ell=0,\dots,L}$ , which combines (Gauß–88 Seidel) smoothing and coarse level correction in the standard way. Naturally, only 89 multiplications with the matrices  $(\Pi_{\ell-1}^{\ell})_{\ell=1,\dots,L}$  and their transposes appear in the 90 interlevel transfer of the algorithm; no mass matrices need to be inverted. Given the 91 meshes  $(\mathscr{T}_{\ell})_{\ell=0,...L}$  and a suitable transfer concept, we can compute all auxiliary 92 matrices in a setup phase.

For a complete convergence analysis of this class of algorithms, which puts the 94 semi-geometric approach into the well-known context of [1], we refer to [5]. There, 95 we carefully distinguish between the generally different domains  $(\Omega_\ell)_{\ell=0,...,L}$  and 96 elaborate requirements for the meshes and the interlevel transfer to obtain a quasi- 97 optimal result.

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**Fig. 1.** Simplified sketch in d=2. Basic idea of the coarse space construction based on nonnested meshes (left). Concerning the experiments: scaling (center) and translation (right) of the coarse meshes keeping the respective fine mesh fixed. We emphasize that all computations are in d=3

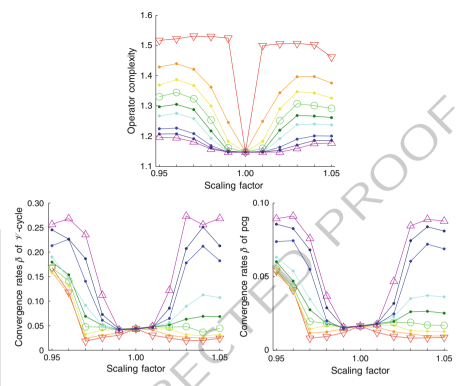
The geometric nature of the construction usually requires some modifications of 99 the meshes and operators, e.g., to ensure full rank. Moreover, a prevalent technique 100 to keep the operator complexity  $\mathscr{C}_{\text{op}} := \sum_{\ell=0}^L n_\ell^A/n_L^A$  small, where  $n_\ell^A$  is the number 101 of non-zero entries of  $A_{\ell}$ , is truncation of the prolongation operators by deleting the 102 entries of  $(\Pi_{\ell-1}^{\ell})_{\ell=1,...,L}$  which are less than a truncation parameter  $\varepsilon_{\rm tr} > 0$  times the maximal entry in the respective row. Afterwards, the modified rows are rescaled such 104 that the row totals remain unchanged; see [7]. All this is done in the setup before the computation of the respective Galerkin products (2). In this paper, we choose  $\Pi_{\ell-1}^{\ell}$  as standard nodal interpolation in  $X_\ell$  for  $\ell \in \{1, ..., L\}$ , namely  $\Pi_{\ell-1}^\ell v := \sum_{p \in \mathscr{N}_\ell} v(p) \lambda_p^\ell$ for all  $v \in X_{\ell-1}$ , and refer to [5] for a detailed discussion.

# 3 Numerical Studies

#### 3.1 The Almost Nested Limiting Case

We consider a hierarchy of four nested meshes  $(\mathcal{I}_{\ell})_{\ell=0,\dots,3}$  of the unit cube in  $\mathbb{R}^3$  111 where the coarsest mesh consists of 768 elements with 189 nodes. Throughout the 112 study, we keep the finest mesh  $\mathscr{T}_L = \mathscr{T}_3$  with 393,216 elements and 68,705 nodes 113 fixed. In contrast, the coarse domains  $(\Omega_\ell)_{\ell < 3}$  and the corresponding coarse meshes  $(\mathscr{T}_\ell)_{\ell < 3}$  are scaled around the center with a different factor between 0.95 and 1.05  $\,$  115 for each set of tests; see Fig. 1 (center).

In the semi-geometric framework, it is absolutely necessary to perform a truncation procedure to retain the optimality of the algorithms. Otherwise, one can in 118 general not prevent the appearance of very small and thus irrelevant entries in the 119 prolongation matrices. We study the complexity of the constructed space hierarchy 120 and the convergence of the semi-geometric multigrid method (stand-alone or in a 121 preconditioned conjugate gradient method) for a variety of values for the parame- 122 ter  $\varepsilon_{\rm tr}$  in [0.01, 0.49]. Note that, for linear finite elements associated with simplicial 123 meshes, it does generally not make sense to choose  $\varepsilon_{tr}$  greater than or equal to 0.5. 124



**Fig. 2.** The complexity measure  $\mathscr{C}_{op}$  (top) and the convergence rates  $\bar{\rho}_{\mathscr{V}(2,2)}$  (left) and  $\bar{\rho}_{\mathscr{V}(2,2)}^{pcg}$ (right) of a semi-geometric multigrid method, plotted versus the scale of the coarse meshes. Each line represents a different parameter  $\varepsilon_{tr} \in [0.01, 0.49]$ . The marked lines correspond to the values 0.01 ( $\nabla$ ), 0.20 ( $\Diamond$ ) and 0.49 ( $\triangle$ ), respectively

This is because such a choice would result in deleting entries even in case of perfectly 125 nested meshes, leaving nodes without direct coupling to the next coarser level.

The results of the experiments with scaled  $(\Omega_{\ell})_{\ell < 3}$  are illustrated in Fig. 2. Each 127 single line represents either the complexity  $\mathscr{C}_{op}$  or one of the asymptotic convergence 128 rates  $ar{
ho}_{\mathscr{V}(2,2)}$  and  $ar{
ho}_{\mathscr{V}(2,2)}^{
m pcg}$  for a fixed parameter  $arepsilon_{
m tr}$  plotted versus the scale of the 129 coarse meshes. The lines corresponding to the extreme  $\varepsilon_{tr}$ -values 0.01 and 0.49 are marked by downward and upward triangles, respectively; an intermediate value of 131 0.20 is marked by circles. Table 1 contains the numbers for these three values. We 132 stop with the scales 0.95 and 1.05, respectively. For smaller factors, the convergence 133 rates further increase quite fast as less and less of the computational domain  $\Omega=134$  $\Omega_L$  is covered by the coarse meshes; the complexity measures do not change much 135 in this case. For larger factors, the convergence rates slowly increase whereas the 136 complexity measures decrease. This is due to the fact that more and more elements 137 of the coarse meshes lie completely outside the computational domain.

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scale	$\mathscr{C}_{op}$	$\bar{\rho}_{\mathscr{V}(2,2)}$	$ar{ ho}_{\mathscr{V}(2,2)}^{ m pcg}$	$\mathscr{C}_{op}$	$\bar{\rho}_{\mathscr{V}(2,2)}$	$ar{ ho}_{\mathscr{V}(2,2)}^{ m pcg}$	$\mathscr{C}_{\operatorname{op}}$	$\bar{\rho}_{\mathscr{V}(2,2)}$	$ar ho_{\mathscr{V}(2,2)}^{ m pcg}$	
0.95	1.52	0.169	0.054	1.33	0.168	0.055	1.20	0.256	0.089	t1.1
0.96	1.52	0.118	0.041	1.34	0.142	0.043	1.19	0.268	0.091	t1.2
0.97	1.53	0.018	0.008	1.32	0.048	0.020	1.18	0.235	0.076	t1.3
0.98	1.53	0.026	0.009	1.25	0.047	0.018	1.16	0.112	0.037	t1.4
0.99	1.52	0.031	0.012	1.16	0.041	0.015	1.15	0.041	0.016	t1.5
1.00	1.15	0.044	0.016	1.15	0.044	0.016	1.15	0.044	0.016	t1.6
1.01	1.50	0.031	0.012	1.16	0.048	0.017	1.15	0.048	0.018	t1.7
1.02	1.51	0.025	0.009	1.25	0.047	0.019	1.15	0.122	0.047	t1.8
1.03	1.51	0.020	0.008	1.31	0.048	0.019	1.16	0.273	0.085	t1.9
1.04	1.50	0.020	0.008	1.30	0.037	0.017	1.18	0.256	0.089	t1.10
1.05	1.46	0.024	0.009	1.29	0.045	0.017	1.18	0.269	0.088	t1.11
										t1.12
	ε	$t_{\rm tr} = 0.01$		ε	$t_{\rm tr} = 0.20$	)	ε	$t_{\rm tr} = 0.49$	)	t1.13

**Table 1.** Studying the convergence behavior for a family of almost nested meshes associated with the unit cube. The middle row (scale 1.00) corresponds to the completely nested case in which the approach coincides with the standard geometric multigrid method.

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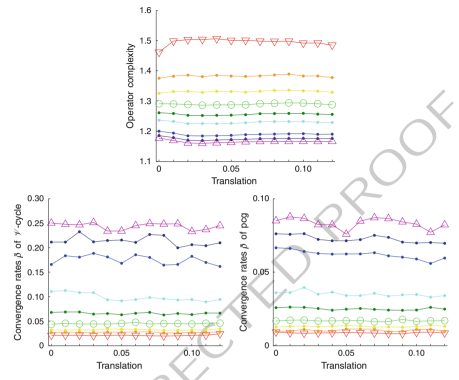
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## 3.2 Robustness of the Coarse Level Hierarchy

The second experiment is to further investigate the influence of perturbations of the 140 meshes on the coarse level hierarchy and the multigrid performance. Here, we consider different translations of the coarse meshes associated with the cube of scale 1.05 142 in direction of the unit vector  $(\frac{2}{3}, \frac{2}{3}, \frac{1}{3})^T \in \mathbb{R}^3$  by sizes up to 0.12. In this case, the computational domain  $\Omega = \Omega_L$  is covered by the domains  $(\Omega_\ell)_{\ell \leq L}$  for almost the entire range of translations; see Fig. 1 (right). Basic robustness of the semi-geometric 145 construction is demonstrated by the results in Fig. 3 where the parameter  $\varepsilon_{tr}$  again 146 varies in the interval [0.01, 0.49].

#### 4 Discussion of the Results

As expected and observed in the vast majority of experiments, the convergence rates 149 principally increase with increasing truncation parameter, which indicates that the 150 constructed coarse spaces have adequate approximation power. Note that the deterioration of the convergence behavior is usually rather slow, though. It is evident that 152 the semi-geometric methods, which leave the coarse meshes flexible, coincide with 153 the standard geometric variants in the special case of nested meshes. In addition, an 154 important observation from Sect. 3.1 is that both the complexities  $\mathscr{C}_{op}$  and the convergence rates of the geometric multigrid methods are retained in case the meshes 156 are almost nested if a suitable parameter  $\varepsilon_{\rm tr}$  is applied; see the discussion below. 157 This also indicates that our construction is robust in the sense that the coarse level 158 hierarchy (and with it the multigrid convergence) only varies slightly if the coarse 159 meshes themselves change slightly. Perturbations of the meshes are irrelevant for the 160



**Fig. 3.** The numbers  $\mathscr{C}_{op}$  (top),  $\bar{\rho}_{\mathscr{V}(2,2)}$  (left), and  $\bar{\rho}_{\mathscr{V}(2,2)}^{pcg}$  (right). Each line represents a different parameter  $\varepsilon_{tr} \in [0.01, 0.49]$  plotted versus the size of the coarse mesh translation

efficiency of the methods. This can also be seen clearly in the experiments described 161 in Sect. 3.2.

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As a general rule, we observe the following effects in Sect. 3.1. The larger the parameter  $\varepsilon_{\rm tr}$  the less sensitive is the complexity  $\mathscr{C}_{\rm op}$  to changes of the coarse meshes. 164 The smaller  $\varepsilon_{\rm tr}$  the less sensitive are the convergence rates to changes of the coarse meshes. In our examples, the convergence actually improves in case of small perturbations for sufficiently small  $\varepsilon_{\rm tr}$ . This is of course accompanied by a rapid increase of  $\mathscr{C}_{on}$ . The choice  $\varepsilon_{tr} = 0.20$  (which is, interestingly enough, a standard value in many 168 algebraic multigrid algorithms) is a reasonable attempt to achieve the two competing 169 goals. It manages to keep the convergence rates almost constant for a rather broad 170 range of different problem sizes while leading to an only moderate increase of  $\mathscr{C}_{on}$ .

Finally, let us compare to the general semi-geometric case. For an unstructured 172 mesh with similar size (64,833 nodes) approximating a ball, the measured rates, 173  $\bar{
ho}_{\mathscr{V}(2,2)}=0.060$  and  $\bar{
ho}_{\mathscr{V}(2,2)}^{\mathrm{pcg}}=0.024$ , are not much worse than the ones produced by the geometric method on the cube with completely nested meshes,  $\bar{\rho}_{\mathscr{V}(2,2)} = 0.044$  175 and  $\bar{\rho}_{\mathcal{V}(2,2)}^{\text{pcg}} = 0.016$ . However, for unstructured meshes without natural coarse level 176 hierarchy, it seems impossible to achieve this fast convergence with an operator complexity as small as 1.15 which is easily obtained in the structured case. 178 For comparison, we have  $\mathscr{C}_{op} = 1.38$  for the ball. A whole series of experiments 179 studying the asymptotics of the semi-geometric preconditioners can be found in [5]. 180

5 Conclusion 181

In this paper, we reported on numerical studies of a class of preconditioners based on 182 non-nested meshes. Considering the almost nested case, we determined a truncation 183 parameter  $\varepsilon_{tr} = 0.20$  of the interlevel transfer to be reasonable in order to ensure that the efficiency of the completely nested case is in large part retained. Moreover, per- 185 turbations of the meshes turned out to be irrelevant for the efficiency of the methods. 186

Our results also show that, in the variational coarse space construction, it is ap- 187 propriate to choose auxiliary meshes mimicking geometric coarsening, which leads 188 to particularly small hierarchical overhead (less than 40%). This is in contrast to the 189 non-variational variant of the auxiliary space method [9] where both analysis and ex- 190 periments indicate that the sizes of the original space and of the auxiliary space need 191 to be comparable in a quite restrictive sense such that  $\mathscr{C}_{op}$  is usually clearly larger 192 than two.

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# **BDDC** for Higher-Order Discontinuous Galerkin Discretizations

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**Summary.** The BDDC algorithm is extended to a large class of discontinuous Galerkin (DG) 7 discretizations of second order elliptic problems in two spatial dimensions. An estimate of 8  $C(1 + \log(p^2H/h))^2$  is obtained for the condition number of the preconditioned system where 9 C is a constant independent of p, h or H. Numerical simulations are presented which confirm 10 the theoretical results

1 Introduction 12

A Balancing Domain Decomposition by Constraints (BDDC) method is presented 13 for the solution of a discontinuous Galerkin (DG) discretization of a second-order 14 elliptic problem in two dimensions. BDDC was originally introduced in [8] for the 15 solution of continuous finite element discretizations. Mandel and Dohrmann [13] 16 later proved a condition number bound of  $\kappa \leq C(1+\log(H/h))^2$  for preconditioned system of a continuous finite element discretization of second order elliptic problems. Pavarino [15] and Klawonn et al. [11] extended the BDDC algorithm 19 to higher-order finite element methods and proved a condition number bound of 20  $\kappa \leq C(1+\log(p^2H/h))^2$ . Further analysis of BDDC methods and their connection 21 to FETI methods has been presented in [12, 14].

While domain decomposition methods have been widely studied for continuous finite element discretizations, relatively little work has been performed for discontinuous Galerkin discretizations. Previous work on domain decomposition methods for DG discretizations include [1, 10] and [9]. This work presents a BDDC 26 method applied to a large class of DG methods considered in the unified analysis 27 of [2]. A key component for the development and analysis of the BDDC algorithm 28 involves presenting the DG discretization as the sum of element-wise "local" bilinear 29 forms. The element-wise perspective leads naturally to the appropriate choice for the 30 subdomain-wise local bilinear forms. Additionally, this perspective enables a connection to be drawn between the DG discretization and a related continuous finite 32 element discretization. As a result of this connection, the condition number bound 33

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for the BDDC preconditioned system for a large class of conservative and consistent 34 DG methods is identical to that for continuous finite element methods.

#### 2 DG Discretization

Consider the second order elliptic equation in a domain  $\Omega \subset \mathbb{R}^2$ :

$$-\nabla \cdot (\rho \nabla u) = f \qquad \text{in } \Omega, \qquad u = 0 \qquad \text{on } \partial \Omega \tag{1}$$

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with positive  $\rho > 0 \in L^{\infty}(\Omega)$ ,  $f \in L^{2}(\Omega)$ . Let the triangulation  $\mathscr{T}$  be a partition of  $\Omega$  38 into triangles or quadrilaterals. In order to simplify the presentation we assume that 39  $\rho$  takes on a constant value,  $\rho_{\kappa}$  on each element  $\kappa$ . Define  $\mathscr{E}$  to be the union of edges 40 of elements  $\kappa$ . Additionally, define  $\mathscr{E}^i \subset \mathscr{E}$  and  $\mathscr{E}^{\partial} \subset \mathscr{E}$  to be the set of interior, 41 respectively boundary edges. Note that any edge  $e \in \mathcal{E}^l$  is shared by two adjacent 42 elements  $\kappa^+$  and  $\kappa^-$  with corresponding outward pointing normal vectors  $n^+$  and 43  $n^-$ . Let  $\mathscr{P}^p(\kappa)$  denote the space of polynomials of order at most p on  $\kappa$  and define 44 the following finite element space  $W_h^p := \{ w_h \in \mathbf{L}^2(\Omega) : w_h|_{\kappa} \in \mathscr{P}^p(\kappa) \quad \forall \kappa \in \Omega \}.$  45 Note that traces of functions  $u_h \in W_h^P$  are in general double valued on each edge, 46  $e \in \mathcal{E}^i$ , with values  $u_h^+$  and  $u_h^-$  corresponding to traces from elements  $\kappa^+$  and  $\kappa^-$  47 respectively. On  $e\in\mathscr{E}^\partial$  , associate  $u_h^+$  with the trace taken from the element,  $\kappa^+\in\mathscr{T}_h$ , 48 neighbouring e. The weak form of (1) on each element is given by:  $\forall w_h \in \mathscr{P}^p(\kappa)$ 

$$(\rho \nabla u_h, \nabla w_h)_{\kappa} - \langle \rho(u_h^+ - \hat{u}_h) \mathbf{n}^+, \nabla w_h^+ \rangle_{\partial \kappa} + \langle \hat{\mathbf{q}}_h, w_h^+ \mathbf{n}^+ \rangle_{\partial \kappa} = (f, w_h)_{\kappa}$$
(2)

where  $(\cdot,\cdot)_{\kappa} := \int_{\kappa}$  and  $\langle\cdot,\cdot\rangle_{\partial\kappa} := \int_{\partial\kappa}$ . Superscript  $^+$  is used to explicitly denote values on  $\partial \kappa$ , taken from  $\kappa$ . For all  $w_h \in W_h^p$ ,  $\hat{w}_h = \hat{w}_h(w_h^+, w_h^-)$  is a single valued 51 numerical trace on  $e \in \mathcal{E}^i$ , while  $\hat{w}_h = 0$  for  $e \in \mathcal{E}^{\partial}$ . Note that  $\hat{u}_h = 0$  on  $e \in \mathcal{E}^{\partial}$ , cor-52 responds to weakly enforced homogeneous boundary conditions on  $\partial\Omega$ . Similarly 53  $\hat{q} = \hat{q}(\rho^+, \rho^-, \nabla u_h^+, \nabla u_h^-, u_h^+, u_h^-)$  is a single valued numerical flux approximating 54  $\mathbf{q} = \rho \nabla u$  on  $e \in \mathcal{E}$ . Summing over all elements gives:

$$a(u_h, w_h) = (f, w_h)_{\Omega} \qquad \forall w_h \in W_h^p$$
(3)

A key component, required for the development and analysis of the algorithms pre- 56 sented, is to express the global bilinear form  $a(u_h, w_h)$  as the sum of element-wise 57 contributions  $a_{\kappa}(u_h, w_h)$  such that

$$a(u_h, w_h) = \sum_{\kappa \in \mathscr{T}} a_{\kappa}(u_h, w_h) \tag{4}$$

where  $a_{\kappa}(u_h, w_h)$  is a symmetric, positive semi-definite "local bilinear form". In particular, the local bilinear form should have a compact stencil, such that  $a_{\kappa}(u_h, w_h)$  is 60 a function of only  $u_h$ ,  $\nabla u_h$  in  $\kappa$ , and  $u_h^+$ ,  $\nabla u_h^+$  and  $\hat{u}_h$  on  $\partial \kappa$ . The local bilinear form 61 is written as:

$$a_{\kappa}(u_{h}, w_{h}) = (\rho \nabla u_{h}, \nabla w_{h})_{\kappa} - \langle \rho(u_{h}^{+} - \hat{u}_{h}) \boldsymbol{n}^{+}, \nabla w_{h}^{+} \rangle_{\partial \kappa} + \langle \hat{\boldsymbol{q}}_{h}^{+}, (w_{h}^{+} - \hat{w}_{h}) \boldsymbol{n}^{+} \rangle_{\partial \kappa}$$
$$= (\rho \nabla u_{h}, \nabla w_{h})_{\kappa} - \langle \rho [u]_{h}^{+}, \nabla w_{h}^{+} \rangle_{\partial \kappa} + \langle \hat{\boldsymbol{q}}_{h}^{+}, [w_{h}]^{+} \rangle_{\partial \kappa}$$
(5)

where  $\hat{\boldsymbol{q}}_h^+ = \hat{\boldsymbol{q}}_h^+(\rho^+, \nabla u_h^+, u_h^+, \hat{u}_h)$  is a "local numerical flux". The choice of the 63 numerical trace  $\hat{u}_h$  and flux  $\hat{\boldsymbol{q}}_h$  define the particular DG method considered. Table 1 64 lists the numerical traces and fluxes for the DG methods considered in this paper, 65 while Table 2 lists the corresponding local bilinear forms.

DC M 41 1	•	^	^+	
DG Method	$\hat{u}_h$	$oldsymbol{q}_h$	$q_h$	t1.1
ĪP	$\{u_h\}$	$-\left\{  ho  abla u_h  ight\} + rac{\eta_e}{h} \left\{  ho \left[\!\left[ u_h  ight]\!\right]^\pm  ight\}$	$-\rho^+\nabla u_h^+ + \frac{\eta_e}{h}\rho^+ \llbracket \rho u_h \rrbracket^+$	t1.2
BR2	$\{u_h\}$	$-\left\{  ho  abla u_h ight\} + \eta_e \left\{  ho r_e(\llbracket u_h  rbracket^\pm)  ight\}$	$-\rho^+\nabla u_h^+ + \eta_e\rho^+r_e(\llbracket u_h\rrbracket^+)$	t1.3
Brezzi	$\{u_h\}$	$\{oldsymbol{q}_h\} + \eta_e \left\{  ho r_e(\llbracket u_h  rbracket^{\pm})  ight\}$	$\boldsymbol{q}_h^+ + \eta_e \rho^+ r_e(\llbracket u_h \rrbracket^+)$	t1.4
LDG	$\{u_h\} - \beta \cdot \llbracket u_h \rrbracket$	$\{\boldsymbol{q}_h\} + \beta  [\![\boldsymbol{q}_h]\!] + \frac{2\eta_e}{h}  \{\rho  [\![u_h]\!]^{\pm}\}$	$oldsymbol{q}_h^+ + rac{\eta_e}{h} ho^+ \llbracket u_h  brace^+$	t1.5
CDG	$\{u_h\}-\beta\cdot \llbracket u_h \rrbracket$	$\left\{oldsymbol{q}_{h}^{e}\right\} + oldsymbol{eta}\left[oldsymbol{q}_{h}^{e} ight] + rac{2\eta_{e}}{h}\left\{ ho\left[\!\left[u_{h} ight]\!\right]^{\pm} ight\}$	$q_h^{e+} + \frac{\eta_e}{h} \rho^+ \llbracket u_h  brace^+$	t1.6

**Table 1.** Numerical fluxes for different DG methods. (IP: Interior Penalty, BR2: [3], Brezzi: [4], LDG: [5] CDG: [16])

 $\begin{array}{ll} & & & & \\ \hline \text{Method } a_{\kappa}(u_{h},w_{h}) & & & \\ \hline \text{IP} & & & & & \\ g + \sum_{e \in \partial \kappa} \frac{\eta_{e}}{h_{e}} \left\langle \rho \left[\!\left[u_{h}\right]\!\right]^{+}, \left[\!\left[w_{h}\right]\!\right]^{+}\right\rangle_{e} & & \\ \hline \text{BR2} & & & & \\ g + \sum_{e \in \partial \kappa} \eta_{e} \left(\rho r_{e}(\left[\!\left[u_{h}\right]\!\right]^{+}), r_{e}(\left[\!\left[w_{h}\right]\!\right]^{+})\right)_{\kappa} & & \\ \hline \text{Brezzi} & & & & \\ g + \left(\rho r_{\kappa}(\left[\!\left[u_{h}\right]\!\right]^{+}), r_{\kappa}(\left[\!\left[w_{h}\right]\!\right]^{+})\right)_{\kappa} + \sum_{e \in \partial \kappa} \eta_{e} \left(\rho r_{e}(\left[\!\left[u_{h}\right]\!\right]^{+}), r_{e}(\left[\!\left[w_{h}\right]\!\right]^{+})\right)_{\kappa} \\ \hline \text{LDG} & & & & \\ g + \left(\rho r_{\kappa}(\left[\!\left[u_{h}\right]\!\right]^{+}), r_{\kappa}(\left[\!\left[w_{h}\right]\!\right]^{+})\right)_{\kappa} + \sum_{e \in \partial \kappa} \frac{\eta_{e}}{h_{e}} \left\langle \rho \left[\!\left[u_{h}\right]\!\right]^{+}, \left[\!\left[w_{h}\right]\!\right]^{+}\right)_{e} \\ \hline \hline \text{CDG} & & & & \\ g + \sum_{e \in \partial \kappa} \left(\rho r_{e}(\left[\!\left[u_{h}\right]\!\right]^{+}), r_{e}(\left[\!\left[w_{h}\right]\!\right]^{+})\right)_{\kappa} + \sum_{e \in \partial \kappa} \frac{\eta_{e}}{h_{e}} \left\langle \rho \left[\!\left[u_{h}\right]\!\right]^{+}, \left[\!\left[w_{h}\right]\!\right]^{+}\right\rangle_{e} \\ \hline \\ \hline \text{Where } g = \left(\rho \nabla u_{h}, \nabla w_{h}\right)_{\kappa} - \left\langle \rho \left[\!\left[u_{h}\right]\!\right]^{+}, \nabla w_{h}^{+}\right\rangle_{\partial \kappa} - \left\langle \rho \nabla u_{h}, \left[\!\left[w_{h}\right]\!\right]^{+}\right\rangle_{\partial \kappa} \\ \end{array} \qquad 12.1 \\ \end{array}$ 

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**Table 2.** Elementwise bilinear form for different DG methods

In the definition of the different DG methods,  $\{u_h\} = \frac{1}{2}(u_h^+ + u_h^-)$  and  $[\![u_h]\!] = 67$   $u_h^+ n^+ + u_h^- n^-$  are average and jump operators on  $e \in \mathscr{E}^i$ . Additionally, a second set 68 of jump operators involving the numerical trace  $\hat{u}$  are given by  $[\![u_h]\!]^+ = u_h^+ n^+ + \hat{u}_h n^-$  69 and  $[\![u_h]\!]^- = \hat{u}_h n^+ + u_h^- n^-$ . Define  $\mathbf{q}_h = -\rho(\nabla u_h - r_\kappa([\![u_h]\!]^+))$  and  $\mathbf{q}_h^e = -\rho(\nabla u_h - 70)$   $[\![u_h]\!]^+$ ) where  $r_\kappa(\phi)$  and  $r_e(\phi) \in [\![\mathscr{P}^p(\kappa)]\!]^n$  are lifting operators defined such that: 71  $(r_\kappa(\phi), \mathbf{v}_h)_\kappa = \langle \phi, \mathbf{v}_h^+ \rangle_\kappa$  and  $(r_e(\phi), \mathbf{v}_h)_\kappa = \langle \phi, \mathbf{v}_h^+ \rangle_e$ ,  $\forall \mathbf{v}_h \in [\![\mathscr{P}^p(\kappa)]\!]^n$ . Additionally, 72 on each edge in  $\mathscr{E}$ ,  $\eta_e$  is a penalty parameter, while  $\beta = \frac{1}{2} S_{\kappa^+}^{\kappa^-} n^+ + S_{\kappa^-}^{\kappa^+} n^-$  is a vector 73 where  $S_{\kappa^+}^{\kappa^-} \in \{0,1\}$  is a switch defined, such that  $S_{\kappa^+}^{\kappa^+} + S_{\kappa^-}^{\kappa^+} = 1$ .

Consider using a nodal basis on each element  $\kappa$  to define  $W_h^p$ . Figure 1 shows 75 graphically the nodal degrees of freedom involved in defining the local bilinear form. 76 For the IP, BR2 and Brezzi schemes, the numerical trace  $\hat{u}_h$  on an edge/face depends 77 on both  $u_h^+$  and  $u_h^-$ . Hence the local bilinear form corresponds to all nodal degrees 78 of freedom defining  $u_h$  on  $\kappa$  as well as nodal values on all edge/faces of  $\partial \kappa \cap \mathscr{E}^i$  79

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corresponding to the trace of  $u_h$  from elements neighbouring  $\kappa$ . On the other hand, 80 for the LDG and CDG methods, the numerical trace  $\hat{u}_h$  takes on the value of  $u_h^+$  if 81  $S_{\kappa^+}^{\kappa^-} = 0$  or  $u_h^-$  if  $S_{\kappa^+}^{\kappa^-} = 1$ . Hence the local bilinear form corresponds only to degrees 82 of freedom defining  $u_h$  on  $\kappa$  and nodal values corresponding to the trace of  $u_h$  on 83 neighbouring elements across edge/faces of  $\partial \kappa \cap \mathcal{E}^i$  for which  $S_{\kappa^+}^{\kappa^-} = 1$ .

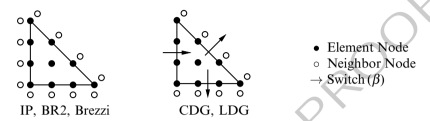


Fig. 1. Degrees of freedom involved in "local" bilinear form

The element-wise bilinear form  $a_{\kappa}(u_h, u_h)$  satisfies

$$a_{\kappa}(u_h, u_h) \ge 0 \tag{6}$$

85

95

with  $a_{\kappa}(u_h, u_h) = 0$  iff  $u_h = \hat{u}_h = K$  for some constant K. The proof of (6) closely 86 follows the proof of boundedness and stability of the different DG methods presented 87 in [2]. As a result it is possible to show that the bilinear form is equivalent to a 88 quadratic form based on the value of  $u_h$  at the nodes x: 89

$$ca_{\kappa}(u_h, u_h) \leq \rho_{\kappa} p^4 h^{n-2} \sum_{\boldsymbol{x}_i, \boldsymbol{x}_j \in \kappa \cup \kappa'} (u_h(\boldsymbol{x}_i) - u_h(\boldsymbol{x}_j))^2 \leq Ca_{\kappa}(u_h, u_h)$$
 (7)

where c and C are constants independent of h, p and  $\rho$ , while  $\mathbf{x}_i, \mathbf{x}_j$  are the nodes 90 on  $\kappa$  defining the basis for  $u_h$  and nodes on  $\partial \kappa'$  defining a basis for the trace  $u_h^-$  91 from neighbours  $\kappa'$  of  $\kappa$ . Using the quadratic form in (7) a connection may be drawn 92 between the DG discretization a continuous finite element discretization on a subtriangulation (See for example [6] Lemma 4.3). Further details are given in [7].

# 3 Domain Decomposition

Consider a partition of the domain  $\Omega$  into substructures  $\Omega_i$  such that  $\bar{\Omega} = \bigcup_{i=1}^N \bar{\Omega}_i$ . 96 The substructures  $\Omega_i$  are disjoint shape regular polygonal regions of diameter O(H), 97 consisting of a union of elements in  $\mathscr{T}$ . Assume that  $\rho(\mathbf{x})$  takes on a constant value, 98  $\rho_i$ , within each subdomain  $\Omega_i$ . Additionally, assume that each element  $\kappa$  in  $\Omega_i$  with 99 an edge e on  $\partial \Omega_i \cap \partial \Omega_j$  has neighbours only in  $\Omega_i \cup \Omega_j$ .

Define the local interface  $\Gamma_i = \partial \Omega_i \setminus \partial \Omega$  and global interface  $\Gamma$  by  $\Gamma = \bigcup_{i=1}^N \Gamma_i$ . 101 Denote by  $W_{\Gamma}^{(i)}$  the space of discrete nodal values on  $\Gamma_i$  which correspond to degrees 102

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of freedom shared between  $\Omega_i$  and neighbouring subdomains  $\Omega_j$ , while  $W_I^{(i)}$  denotes 103 the space of discrete unknowns local to a single substructure  $\Omega_i$ . In particular, note 104 that for the IP, BR2 and Brezzi et al. methods  $W^{(i)}_\Gamma$  includes for each edge  $e \in \Gamma_i$  105 degrees of freedom defining two sets of trace values  $u^+$  from  $\kappa^+ \in \Omega_i$  and  $u^-$  for 106  $\kappa^- \in \Omega_i$ . Thus,  $W_i^{(i)}$  corresponds to nodal values strictly interior to  $\Omega_i$  or on  $\partial \Omega_i \backslash \Gamma_i$ . 107 On the other hand, for the CDG and LDG methods  $W_{\Gamma}^{(i)}$  includes for each edge  $e \in \Gamma_i$  108 degrees of freedom defining a single trace value corresponding to either  $u^+$  from 109  $\kappa^+ \in \Omega_i$  if  $S_{\kappa^+}^{\kappa^-} = 0$  or  $u^-$  from  $\kappa^- \in \Omega_j$  if  $S_{\kappa^+}^{\kappa^-} = 1$ . Hence,  $W_I^{(i)}$  corresponds to 110 nodal values interior to  $\Omega_i$  and on  $\partial \Omega_i \backslash \Gamma_i$  as well as nodal values defining  $u^+$  on 111  $e \in \Gamma_i$  for which  $S_{\kappa^+}^{\kappa^-} = 1$ .

Similarly, define  $\hat{W}_{\Gamma}$  as the space of degrees of freedom shared among multiple 113 subdomains and  $W_I$  as the space of degrees of freedom which correspond only to a 114 single subdomain. Note that  $W_I$  is equal to the product space  $W_I := \Pi_{i=1}^N W_I^{(i)}$ , while in general  $\hat{W}_{\Gamma} \subset W_{\Gamma} := \Pi_{i=1}^N W_{\Gamma}^{(i)}$ . Define local operators  $R_{\Gamma}^{(i)} : \hat{W}_{\Gamma} \to W_{\Gamma}^{(i)}$  which 116 extract the local degrees of freedom on  $\Gamma_i$  from those on  $\Gamma$ . Additionally define a 117 global operator  $R_{\Gamma}: \hat{W}_{\Gamma} \to W_{\Gamma}$  which is formed by a direct assembly of  $R_{\Gamma}^{(i)}$ . The 118 discrete form of (3) is written as:

$$\begin{bmatrix} A_{II} & A_{\Gamma I}^T \\ A_{\Gamma I} & A_{\Gamma \Gamma} \end{bmatrix} \begin{bmatrix} u_I \\ u_{\Gamma} \end{bmatrix} = \begin{bmatrix} b_I \\ b_{\Gamma} \end{bmatrix}. \tag{8}$$

where  $u_I$  and  $u_{\Gamma}$  corresponds to degrees of freedom associated with  $W_I$  and  $\hat{W}_{\Gamma}$  120 respectively. Since the degrees of freedom associated with  $W_I$  are local to a particular 121 substructure they may be locally eliminated to obtain a system

$$\hat{S}_{\Gamma}u_{\Gamma} = g_{\Gamma} \tag{9}$$

122

where  $\hat{S}_{\Gamma} = A_{\Gamma\Gamma} - A_{\Gamma I}A_{II}^{-1}A_{\Gamma I}^{T}$  and  $g_{\Gamma} = b_{\Gamma\Gamma} - A_{\Gamma I}A_{II}^{-1}b_{\Gamma I}$ .  $\hat{S}_{\Gamma}$  and  $g_{\Gamma}$  may be 123 formed by a direct assembly: 124

$$\hat{S}_{\Gamma} = \sum_{i=1}^{N} R_{\Gamma}^{(i)^{T}} S_{\Gamma}^{(i)} R_{\Gamma}^{(i)} \qquad g_{\Gamma} = \sum_{i=1}^{N} R_{\Gamma}^{(i)^{T}} g_{\Gamma}^{(i)}$$
 (10)

where 
$$S_{\Gamma}^{(i)} = A_{\Gamma\Gamma}^{(i)} - A_{\Gamma I}^{(i)} A_{II}^{(i)^{-1}} A_{\Gamma I}^{(i)^{T}}$$
 and  $g_{\Gamma}^{(i)} = b_{\Gamma}^{(i)} - A_{\Gamma I}^{(i)} A_{II}^{(i)^{-1}} b_{I}^{(i)}$ .

4 BDDC 126

A BDDC preconditioner is used to solve the Schur complement problem (9). A full 127 description of the BDDC preconditioner is given by Li and Widlund [12]. In order to 128 define the BDDC preconditioner  $W_{\Gamma}^{(i)}$  is reparameterize into two orthogonal spaces 129  $W_{II}^{(i)}$  and  $W_{\Delta}^{(i)}$ . The primal space  $W_{II}^{(i)}$  is the space of discrete unknowns corresponding to functions with a constant value of  $\hat{u}$  on each edge of substructure  $\Omega_i$ . The

dual space,  $W_A^{(i)}$  is the space of discrete unknowns corresponding to functions which have zero mean value of  $\hat{u}$  on  $\Gamma_i$ . For continuous finite element discretizations, different primal degrees of freedom such as subdomain corners have also been used, 134 however these are not explored in this work. The BDDC algorithm is implemented 135 using a change of basis as described in [12]. The partially assembled space is defined 136 as  $\tilde{W}_{\Gamma}=\hat{W}_{\Pi}\oplus\left(\Pi_{i=1}^{N}W_{\Delta}^{(i)}\right)$ , where  $\hat{W}_{\Pi}$ , single valued on  $\Gamma$ , is formed by assembling the local primal spaces,  $W^{(i)}_{\Pi}$ . Define additional local operators  $\bar{R}^{(i)}_{\Gamma}: \tilde{W}_{\Gamma} \to W^{(i)}_{\Gamma}$ which extract the degrees of freedom in  $\tilde{W}_{\Gamma}$  corresponding to  $\Gamma_i$ . The global operator  $\bar{R}_{\Gamma}: \tilde{W}_{\Gamma} \to W_{\Gamma}$  is formed by a direct assembly of  $\bar{R}_{\Gamma}^{(i)}$ . Also define the global operator 140  $\tilde{R}_{\Gamma}: \hat{W}_{\Gamma} \to \tilde{W}_{\Gamma}$ . The partially assembled Schur complement matrix  $\tilde{S}$ , is given by: 141

$$\tilde{S}_{\Gamma} = \sum_{i=1}^{N} \bar{R}_{\Gamma}^{(i)^{T}} S_{\Gamma}^{(i)} \bar{R}_{\Gamma}^{(i)}$$
(11)

The scaled operator  $\tilde{R}_{D,\Gamma}:\hat{W}_{\Gamma}\to \tilde{W}_{\Gamma}$  is obtained by multiplying the entries of  $\tilde{R}_{\Gamma}$  142 corresponding to  $W_{\Lambda}^{(i)}$  by  $\delta_i^{\dagger}(x)$ , where  $\delta_i^{\dagger}(x)$  defined for each nodal degree of freedom in  $W_{\Gamma}^{(i)}$  on  $\partial\Omega_i$  and  $\partial\Omega_j$  as  $\delta_i^{\dagger}=\frac{\rho_i^{\hat{Y}}}{\rho_i^{\hat{Y}}+\rho_j^{\hat{Y}}}, \gamma\in[1/2,\infty)$ . The BDDC preconditioner 144  $M_{\mathrm{BDDC}}^{-1}:\hat{W}_{\Gamma}\to\hat{W}_{\Gamma}$  is given by:

$$M_{\rm BDDC}^{-1} = \tilde{R}_{D,\Gamma}^T \tilde{S}_{\Gamma}^{-1} \tilde{R}_{D,\Gamma} \tag{12}$$

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The condition number of the preconditioner operator  $M_{\rm BDDC}^{-1}\hat{S}$  is bounded by 146  $C(1 + \log(p^2H/h))^2$  where C is a constant independent of p, h, H or  $\rho$ . This is the same condition number bound as obtained by Klawonn et al. [11] for a continuous 148 finite element discretization. Proof of this condition number bound closely follows 149 that presented by Tu [17] for mixed finite element methods, which in turn builds upon 150 the work of [6]. The key idea is to connect the DG discretization to a related continuous finite element discretization on a subtriangulation of  $\mathcal{T}$ . The ability to connect the DG discretization to the continuous finite element discretization is a direct 153 result of (7) (see [6]). The existing theory for continuous finite elements developed 154 in [13, 15] and [11] is then leveraged to obtain the desired condition number bound. 155 Further details are provided in [7].

#### 5 Numerical Results

This section presents numerical results using the BDDC preconditioner introduced 158 in Sect. 4. For each numerical experiment the linear system resulting from the DG 159 discretization is solved iteratively using a Preconditioned Conjugate Gradient (PCG) 160 method, starting from zero initial condition until  $l_2$  norm of the residual is decreased by a factor of  $10^{10}$ . The domain  $\Omega = (0,1)^2$  is partitioned into  $N \times N$  square subdomains  $\Omega_i$  with side lengths H such that  $N = \frac{1}{H}$ . Each subdomain is the union of triangular elements obtained by bisecting squares of side length h. In the first numerical 164

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experiment (1) is solved on  $\Omega$  with  $\rho = 1$  and f chosen such that the exact solution is 165 given by  $u = \sin(\pi x)\sin(\pi y)$ . Table 3 shows the number of PCG iteration required to converge varying N,  $\frac{H}{h}$  and p for each of the DG discretization considered. Table 3 167 also gives the Lanczos estimate of the maximum eigenvalue of the preconditioned 168 system. The minimum eigenvalue is bounded below by unity as with continuous 169 finite element methods. As expected the number of iterations is independent of the 170 number of subdomains and only weakly dependent on the number of elements per 171 subdomain or the solution order.

$\frac{1}{H}$	$\frac{H}{h}$	p	IP	BR2	Brezzi	LDG	CDG
2			12 (12.1)	15 (12.0)	15 (7.7)	11 (6.1)	12 (5.9)
4			22 (14.3)	27 (14.0)	23 (9.2)	24 (7.4)	24 (7.1)
8	8	4	31 (15.2)	34 (14.8)	30 (9.8)	28 (7.7)	27 (7.5)
16			33 (15.3)	36 (14.9)	32 (9.9)	29 (8.0)	28 (7.8)
32			33 (15.3)	36 (14.9)	32 (9.9)	29 (7.9)	27 (7.7)
	2		25 (10.9)	29 (10.9)	26 (6.9)	23 (5.2)	23 (5.3)
	4		29 (13.0)	34 (12.8)	28 (8.3)	26 (6.4)	25 (6.2)
8	8	4	31 (15.2)	34 (14.8)	30 (9.8)	28 (7.8)	27 (7.5)
	16		33 (17.6)	36 (17.1)	33 (11.5)	29 (9.3)	29 (9.1)
	32		35 (20.2)	38 (19.4)	34 (13.4)	32 (11.0)	31(10.7)
		1	32 (11.1)	36 (13.8)	28 (8.1)	26 (5.9)	25 (5.6)
		2	31 (12.9)	34 (14.1)	29 (8.7)	26 (6.4)	26 (6.3)
8	8	4	31 (15.2)	34 (14.8)	30 (9.8)	28 (7.8)	27 (7.5)
		8	34 (18.4)	37 (16.2)	34 (11.7)	31 (9.9)	32 (9.6)
		16	36 (22.5)	38 (18.6)	38 (14.4)	34 (12.8)	36 (12.2)

**Table 3.** Iteration count ( $\lambda_{max}$ ) for BDDC preconditioner using different DG methods

In the second numerical experiment the behaviour of the preconditioner for large 173 jumps in the coefficient  $\rho$  is examined. For this numerical experiment only the CDG 174 discretization is used. The domain is partitioned in a checkerboard pattern with  $\rho=1$  175 on half of the subdomains and  $\rho = 1,000$  in the remaining subdomains. Initially set 176  $\delta_i^\dagger=rac{1}{2},$  which corresponds to setting  $\gamma=0,$  which does not satisfy the assumption 177  $\gamma \in [1/2, \infty)$ . Poor convergence of the BDDC algorithm is seen in Table 4a. Next  $\delta_i^{\dagger}$  178 is set to  $\delta_i^{\dagger} = \frac{\rho_i}{\rho_i + \rho_j}$  which corresponds to  $\gamma = 1$ . With this choice of  $\delta_i^{\dagger}$  the good 179 convergence properties of the BDDC algorithm is recovered as shown in Table 4b.

6 Conclusions 181

The BDDC preconditioner has been extended to a large class of DG discretizations 182 for second-order elliptic problems. The condition number of the BDDC precondi- 183 tioned system is bounded by  $C(1 + \log(p^2H/h))^2$ , with constant C independent of 184 p, h, H or the coefficient  $\rho$ . This is the same condition number bound previously 185 proven for continuous finite element methods. Numerical results confirm the theory. 186

(a) $\delta_i^{\dagger} = \frac{1}{2}, \frac{H}{h} = 8$									
			$\frac{1}{H}$						
p	2	4	8	16	32				
1	51	119	179	215	232				
3	55	133	207	267	232 316 361				
5	59	153	242	306	361				

(	(b)	$\delta_i^{\dagger}$	=	$\frac{\rho_i}{\rho_i + \mu}$	$\frac{1}{0_j}, \frac{F_j}{F_j}$	$\frac{d}{dt} = 8$				
		$\frac{1}{H}$								
	p	2	4	8	16	32				
	1	4	7	14	18	19				
	3	4	7	15	18 18 19	19				
	5	4	7	14	19	20				

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**Table 4.** Iteration count for BDDC preconditioner using the CDG method with  $\rho = 1$  or 1000.

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# **ARAS2 Preconditioning Technique for CFD Industrial** 2 Cases

3

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Schwarz (RAS) iterative process [2] writes:

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1 Introduction 9

The convergence rate of a Krylov method such as the Generalized Conjugate Resid- 10 ual (GCR) [6] method, to solve a linear system  $Au = f, A = (a_{ij}) \in \mathbb{R}^{m \times m}, u \in$  $\mathbb{R}^m, f \in \mathbb{R}^m$ , decreases with increasing condition number  $\kappa_2(A) = ||A||_2 ||A^{-1}||_2$ of the non singular matrix A. Left preconditioning techniques consist of solving 13  $M^{-1}Au = M^{-1}f$  such that  $\kappa_2(M^{-1}A) \ll \kappa_2(A)$ . The Additive Schwarz (AS) preconditioning is built from the adjacency graph G = (W, E) of A, where  $W = \{1, 2, ..., m\}$  15 and  $E = \{(i, j) : a_{ij} \neq 0\}$  are the edges and vertices of G. Starting with a nonoverlapping partition  $W = \bigcup_{i=1}^{p} W_{i,0}$  and  $\delta \ge 0$  given, the overlapping partition  $\{W_{i,\delta}\}$ is obtained defining p partitions  $W_{i,\delta} \supset W_{i,\delta-1}$  by including all the immediate neighboring vertices of the vertices in the partition  $W_{i,\delta-1}$ . Then the restriction operator  $R_{i,\delta}$  from W to  $W_{i,\delta}$  defines the local operator  $A_{i,\delta} = R_{i,\delta}AR_{i,\delta}^T, A_{i,\delta} \in \mathbb{R}^{m_i,\delta \times m_i,\delta}$  20 on  $W_{i,\delta}$ . The AS preconditioning writes:  $M_{AS,\delta}^{-1} = \sum_{i=1}^{p} R_{i,\delta}^T A_{i,\delta}^{-1} R_{i,\delta}$ . Introducing  $\tilde{R}_{i,\delta}$  21 the restriction matrix on a non-overlapping subdomain  $W_{i,0}$ , the Restricted Additive 22

$$u^{k} = u^{k-1} + M_{RAS,\delta}^{-1} \left( f - Au^{k-1} \right), \text{ with } M_{RAS,\delta}^{-1} = \sum_{i=1}^{p} \tilde{R}_{i,\delta}^{T} A_{i,\delta}^{-1} R_{i,\delta}$$
 (1)

The RAS exhibits a faster convergence than the AS, as shown in [5], leading to a 24 better preconditioning that depends of the number of subdomains. When it is applied 25 to linear problems, the RAS has a pure linear rate of convergence/divergence that can 26 be enhanced with optimized boundary conditions giving the ORAS method of [11]. 27 The RAS method's linear convergence allows its acceleration of the convergence by 28 the Aitken's process as done in [8] for the Schwarz method.

In [4] the present authors designed the ARAS2 preconditioning technique based 30 on the Aitken's acceleration of the convergence technique. This paper presents an 31 approach to solve linear systems coming from CFD industrial cases. The choice of an 32

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approximation space based on the Singular Value Decomposition of the interface's 33 solutions of the RAS iterative process presented in [14] is done. This provides a 34 preconditioning technique that depends on the Right Hand Side but with a very low 35 computational time and totally algebraic.

## 2 The ARAS2 Preconditioning Method

In what follows, we write the Aitken Restricted Additive Schwarz (ARAS) iterative 38 process and the associated preconditioner. This preconditioner belongs to the family of the two-level preconditioner techniques (see [10, 13] and references) but the 40 coarse grid operator uses only parts of the artificial interfaces contrary to the patch 41 substructuring method of [7]. In this way, it can be seen as similar as the SchurRAS 42 method of [9] but it differs because the discrete Steklov-Poincaré operator connects 43 the coarse artificial interfaces of all the subdomains.

#### 2.1 The ARAS and ARAS2 Preconditioner's Formulation

Let  $\Gamma_i = W_{i,\delta+1} \setminus W_{i,\delta}$  be the interface associated to  $W_{i,\delta}$  and  $\Gamma = \bigcup_{i=1}^p \Gamma_i$  be the global 46 interface. Then  $u_{|\Gamma} \in \mathbb{R}^n$  is the restriction of the solution  $u \in \mathbb{R}^m$  on the  $\Gamma$  interface 47 and  $e_{|\Gamma}^k = u_{|\Gamma}^k - u_{|\Gamma}^\infty$  is the error of (1) at the interface  $\Gamma$ . Taking into account that 48 there exists a matrix  $P \in \mathbb{R}^{n \times n}$  independent of the iterate k such that  $e^k_{|\Gamma} = Pe^{k-1}_{|\Gamma}$ , 49 we can apply the Aitken's acceleration of the convergence process [8] (if |P| < 1 to 50 ensure existence of  $(I_n - P)^{-1}$  for example) as follows:

$$u_{|\Gamma}^{\infty} = (I_n - P)^{-1} \left( u_{|\Gamma}^k - P u_{|\Gamma}^{k-1} \right). \tag{2}$$

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P can be computed analytically or numerically for a separable operator on separable 52 geometry [8] or numerically approximated in other cases [14]. Using this property 53 on the RAS method, we would like to write a preconditioner which includes the 54 Aitken's acceleration process. We introduce a restriction operator  $R_{\Gamma} \in \mathbb{R}^{n \times m}$  from 55 W to the global artificial interface  $\Gamma$ , with  $R_{\Gamma}R_{\Gamma}^{T}=I_{n}$ .

The Aitken Restricted Additive Schwarz (ARAS) must generate a sequence of 57 solutions on the interface  $\Gamma$ , and accelerate the convergence of the Schwarz process 58 from this original sequence. Then the accelerated solution on the interface replaces 59 the last one. This could be written combining an AS or RAS process Eq. (3a) with 60 the Aitken process written in  $\mathbb{R}^{m \times m}$  Eq. (3b) and substracting the Schwarz solution 61 which is not extrapolated on  $\Gamma$  Eq. (3c). We can write the following approximation 62  $u^*$  of the solution u:

$$u^* = u^{k-1} + M_{RAS,\delta}^{-1}(f - Au^{k-1})$$
(3a)

$$+R_{\Gamma}^{T}\left(I_{n}-P\right)^{-1}\left(u_{|\Gamma}^{k}-Pu_{|\Gamma}^{k-1}\right) \tag{3b}$$

$$-R_{\Gamma}^{T}I_{n}R_{\Gamma}\left(u^{k-1}+M_{RAS,\delta}^{-1}(f-Au^{k-1})\right) \tag{3c}$$

We would like to write  $u^*$  as an iterated solution derived from an iterative process 64 of the form  $u^* = u^{k-1} + M_{ARAS,\delta}^{-1} (f - Au^{k-1})$ , where  $M_{ARAS,\delta}^{-1}$  is the Aitken-RAS 65 preconditioner.

Hence the formulation Eq. (3) leads to an expression of an iterated solution  $u^*$ : 67

$$u^* = u^{k-1} + \left(I_m + R_{\Gamma}^T \left( (I_n - P)^{-1} - I_n \right) R_{\Gamma} \right) M_{RAS,\delta}^{-1} \left( f - Au^{k-1} \right)$$

This iterated solution  $u^*$  can be seen as an accelerated solution of the RAS it- 68 erative process. Drawing our inspiration from the Stephensen's method, we build a 69 new sequence of iterates from the solutions accelerated by the Aitken's acceleration 70 method. Such a process is done in [12]. Then, one considers  $u^*$  as a new  $u^k$  and writes 71 the following ARAS iterative process: 72

$$u^{k} = u^{k-1} + \left(I_{m} + R_{\Gamma}^{T} \left( (I_{n} - P)^{-1} - I_{n} \right) R_{\Gamma} \right) M_{RAS, \delta}^{-1} \left( f - A u^{k-1} \right)$$
 (4)

Then we defined the ARAS preconditioner as

$$M_{ARAS,\delta}^{-1} = \left(I_m + R_{\Gamma}^T \left( (I_n - P)^{-1} - I_n \right) R_{\Gamma} \right) \sum_{i=1}^p \tilde{R}_{i,\delta}^T A_{i,\delta}^{-1} R_{i,\delta}$$
 (5)

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If P is known exactly, the ARAS process written in Eq. (4) needs two steps to 74 converge to the solution u with an initial guess  $u^0 = 0$ . Then we have: 75

**Proposition 1.** *If P is known exactly then we have* 

**Proposition 1.** If 
$$P$$
 is known exactly then we have 
$$A^{-1} = \left(2M_{ARAS,\delta}^{-1} - M_{ARAS,\delta}^{-1}AM_{ARAS,\delta}^{-1}\right) \text{ that leads } \left(I - M_{ARAS,\delta}^{-1}A\right) \text{ to be a nilpo-}$$
77 tent matrix of degree 2.

The previous proposition leads to an approximation of  $A^{-1}$  written from the 2 first 79 iterations of the ARAS iterative process (4). Those 2 iterations compute the Schwarz 80 solutions sequence on the interface needed in order to accelerate the Schwarz method 81 by the Aitken's acceleration. We now write 2 iterations of the ARAS iterative pro- 82 cess (4) for any initial guess and for all  $u^{k-1} \in \mathbb{R}^m$ . 83

$$u^{k+1} = u^{k-1} + \left(2M_{ARAS,\delta}^{-1} - M_{ARAS,\delta}^{-1}AM_{ARAS,\delta}^{-1}\right)\left(f - Au^{k-1}\right)$$

Then we defined the ARAS2 preconditioner as

$$M_{ARAS2.\delta}^{-1} = 2M_{ARAS.\delta}^{-1} - M_{ARAS.\delta}^{-1} A M_{ARAS.\delta}^{-1}$$
 (6)

Hence, if P is known exactly there is no need to use ARAS as a preconditioning technique. Nevertheless, when P is approximated, the Aitken's acceleration of the convergence depends on the local domain solving accuracy, and the cost of the building 87 of an exact P depends on the size n. This is why P is numerically approximated by 88  $P_{\mathbb{U}_q}$ , defining  $q \leq n$  orthogonal vectors  $\mathbb{U}_q \in R^{n imes q}$ , that are able to approximate most 89 of the solution at the interface  $\Gamma$ . Then ARAS( $\mathbb{U}_q$ ) and ARAS2( $\mathbb{U}_q$ ) can be defined 90 as:

$$M_{ARAS(\mathbb{U}_q),\delta}^{-1} = \left(I_m + R_{\Gamma}^T \mathbb{U}_q \left( \left(I_q - P_{\mathbb{U}_q} \right)^{-1} - I_q \right) \mathbb{U}_q^T R_{\Gamma} \right) \sum_{i=1}^p \tilde{R}_{i,\delta}^T A_{i,\delta}^{-1} R_{i,\delta} \tag{7}$$

and

$$M_{ARAS2(\mathbb{U}_q),\delta}^{-1} = 2M_{ARAS(\mathbb{U}_q),\delta}^{-1} - M_{ARAS(\mathbb{U}_q),\delta}^{-1} A M_{ARAS(\mathbb{U}_q),\delta}^{-1}$$
(8)

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As the basis  $\mathbb{U}_q$  can only give an approximation of the searched solution at the interface, it make sense to use  $M_{ARAS(\mathbb{U}_a),\delta}^{-1}$  and  $M_{ARAS2(\mathbb{U}_a),\delta}^{-1}$  as preconditioners.

## 2.2 Orthogonal Basis $\mathbb{U}_q$ Arising from SVD of the Interface's Solutions of Richardson Process

The objective is to compute  $P_{\mathbb{U}_q}$  saving as much computing as possible. The singular 97 value decomposition offers a tool to concentrate the effort only on the main parts of 98 the solution. A singular-value decomposition of a real  $n \times q$  (n > q) matrix Y is its 99 factorization into the product of three matrices  $Y = \mathbb{U}_q \Sigma \mathbb{V}^*$ , where  $\mathbb{U}_q = [U_1, \dots, U_q]$  100 is an  $n \times q$  matrix with orthonormal columns,  $\Sigma$  is an  $n \times q$  nonnegative diagonal 101 matrix with  $\Sigma_{ii} = \sigma_i$ ,  $1 \le i \le q$  and the  $q \times q$  matrix  $\mathbb{V} = [V_1, \dots, V_q]$  is orthogonal. The left  $\mathbb{U}_q$  and right  $\mathbb{V}$  singular vectors are the eigenvectors of  $YY^*$  and  $Y^*Y$  respectively. 103 It readily follows that  $Av_i = \sigma_i u_i$ ,  $1 \le i \le q$ . We are going to recall some properties 104 of the SVD. Assume that the  $\sigma_i$ ,  $1 \le i \le q$  are ordered in decreasing order and there exists an r such that  $\sigma_r > 0$  while  $\sigma_r + 1 = 0$ . Then A can be decomposed in a dyadic 106 decomposition:

$$Y = \sigma_1 U_1 V_1^* + \sigma_2 U_2 V_2^* + \ldots + \sigma_r U_r V_r^*.$$
(9)

This means that SVD provides a way to find optimal lower dimensional approxima- 108 tions of a given series of data. More precisely, it produces an orthonormal basis for 109 representing the data series in a certain least squares optimal sense.

The orthogonal "basis"  $\mathbb{U}_q$  is obtained as follows. q iterations of the Richardson 111 process  $u^k = u^{k-1} + M_{RAS}^{-1} \delta(f - Au^{k-1})$  are performed and  $R_{\Gamma} u^k \in \mathbb{R}^n, 1 \le k \le q$  112 belonging to the interface  $\Gamma$  are stored in a matrix  $Y \in \mathbb{R}^{n \times q}$ . Then the SVD of Y 113 is computed to obtain the matrix  $\mathbb{U}_q$  with an arithmetic cost less than the one of a 114 local solution. It leads to efficiency and low computational cost as illustrated in [1]. 115 Nevertheless, the preconditioner ARAS2( $\mathbb{U}_q$ ) obtained is solution dependent.

# **2.3** Building of the $P_{\mathbb{U}_a}$ Matrix

The matrix  $P_{\mathbb{U}_q}$  can be computed as follows keeping the q+1 first singular values of the SVD greater than a set tolerance, we writes: 119

$$\mathbf{Y}_{1:a,1:a+1} = \Sigma_{1:a,1:a} \mathbb{V}_{1:a,1:a+1}^{T} \tag{10}$$

$$\mathbf{E}_{1:q,1:q+1} = \mathbf{Y}_{1:q,2:q+1} - \mathbf{Y}_{1:q,1:q}$$
(11)

If 
$$\mathbf{E}_{1:q,1:q}$$
 is invertible then (12)

$$P_{\mathbb{U}_q} = \mathbf{E}_{1:q,2:q+1} \, \mathbf{E}_{1:q,1:q}^{-1} \tag{13}$$

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The previous building requires the inversion of the matrix  $\mathbf{E}_{1:q,1:q}$  which can be ill 120 conditioned. It is why the second building of matrix  $P_{\mathbb{U}_q}$  that follows is prefered. 121 Selecting the q first singular values of the SVD greater than a set tolerance, one 122 iteration of the RAS algorithm is applied on the q the homogeneous problems where 123  $U^i$ ,  $1 \le i \le q$  is set as boundary condition on the interface  $\Gamma$ . The result of this RAS 124 iterate with  $M_{RAS,\delta}^{-1}$  on the boundary  $\Gamma$  is the column of  $P_{\mathbb{U}_q}$  associated with the 125 component  $U_i$  of the basis. Let us notice that this q computing can be made in the 126 same time considering the q right hand sides in a matrix form.

# 3 Numerical Experiments on 2D and 3D Industrial Problems from Navier-Stokes Equations

In this section we focus on solving linear systems coming from industrial problems 130 with the ARAS2 preconditioning technique. The sparse matrices correspond to the 131 assemblage of all the elementary Jacobian matrices resulting from the partial first- 132 order derivations with respect to the conservative fluid variables of the discrete steady 133 (real) Reynolds-averaged Navier-Stokes equations. We note here that the Jacobian 134 matrix is non-symmetric and is non positive definite.

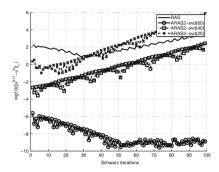
Table 1 summarizes the main features of the linear systems from the two cases 136 solved. Those cases are available in the sparse matrix collection [3]. Turbulence is 137 considered in the 2D and 3D cases. We partition the system with PARMETIS into 138 p subdomains. We must notice that for such problems with non-elliptic operators, 139 the ILU factorization is hazardous. Then, the preconditioner is computed from exact 140 factorization of local operators.

Figure 1 presents for the case PR02 the convergence behaviour of the Richard- 142 son and the GMRES preconditioned by the ARAS2 preconditioner where the  $P_{\mathbb{U}_a}$  143 is approximated by SVD. For this matrix the RAS Richardson process diverges. If 144 the number of singular values kept is not sufficient, the ARAS2 process diverges as 145 well. If we used 60 iterates of RAS Richardson process then the "full"  $P_{\mathbb{U}_q}$  makes 146 the ARAS2 Richardson process converge in one iterate. Nevertheless ARAS2 works 147 quite well in both cases as a preconditioner of the GMRES method. We must notice 148 that here we have an effective gain to use the ARAS2 instead of RAS as Richardson 149 process. The same behavior is also retrieved when ARAS2 is used as preconditioner. 150

For a 3D case the number of non-zero and the band profile increase. Then solving 151 local problems by LU factorization begins to be expensive in terms of memory. A 152 better approach consists of solving subproblems by an iterative method. For the case 153 RM07, we choose to solve subproblems by a GMRES preconditioned by ILU. The 154 idea to save computational time is to approximate the Aitken's acceleration with the 155 basis arising from SVD and solving subproblems with less accuracy for the computing of the preconditioner. Table 2 shows the good strong numerical scalability of the 157 ARAS2 preconditioning compare to the RAS.

	order			nnz
				8 185 136
RM07	381 689	3D	54 527	37 464 962

**Table 1.** Main features of the linear systems with *order* the size of the matrix with real coefficients, dim the dimension of the problem, nn is the number of mesh nodes, nnz is the number of non-zero elements in the matrix



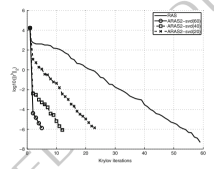


Fig. 1. Solving 2D Navier Stokes equation with turbulence (CASE PR02), PARMETIS partitioning, p = 4, overlap 2, ARAS2 is built with a SVD basis, (*left*) Convergence of Iterative Schwarz Process, (right) convergence of GMRES method preconditioned by RAS and ARAS2

p	RAS	ARAS(36)	ARAS2(36)
3	87 (1.)	77 (1.1299)	53 (1.6415)
6	112 (1.)	93 (1.2043)	63 (1.7778)
12	171 (1.)	124 (1.3790)	84 (2.0357)

Table 2. CASE RM07: Number of GMRES iterations (ratio of iterations with RAS over iterations with ARAS or ARAS2) for a tolerance 1e-10, overlap 1.

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# An Implicit and Parallel Chimera Type Domain **Decomposition Method**

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## 1 Introduction

The Chimera Method developed originally in [1, 19, 20] simplifies the construction 16 of computational meshes about complex geometries. This is achieved by breaking 17 the geometries into components and generating independently a series of different 18 meshes. This enables one a great flexibility on the choice of the type of elements, 19 their orientations and local mesh refinement. The components are further coupled by 20 transmitting information from one mesh to the other to obtain a global solution.

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The Chimera Method is a very efficient tool to treat moving objects [3, 16] as the 22 different meshes can move as rigid bodies in an independent way. Nevertheless, we 23 will focus in this work on fixed subdomains. The main application in this context is 24 optimization analysis, where different configurations can be tested without having to 25 remesh the whole geometry. In order to achieve this, we have developed a versatile 26 strategy based on the Chimera Method.

Usually, in the Chimera Method, the mesh is divided into a background mesh, 28 which covers all the computational domain, and patch (overset) meshes attached to 29 the different components (objects) which are located upon the background mesh. 30 First, we apply a proper preprocessing consisting in removing elements of the background mesh located inside the patch meshes to create apparent interfaces between 32 the background and the patches. The present algorithm requires in addition to smooth 33 the interfaces. This is achieved using a smoothing strategy of the interfaces and the 34 neighboring volume mesh. Then a new coupling algorithm is carried out in order to 35 obtain a "continuous solution" across the interfaces. In the literature, the Chimera 36 coupling has generally been implemented as an iterative algorithm (see [2] for a 37

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Schwarz coupling or [9] for a Dirichlet/Neumann coupling). Here the coupling is 38 implicit. The implementation properties of the proposed coupling facilitate its parallel implementation and makes it a versatile method to be used on general PDE's.

In the following we explain the two basic steps of the Chimera method. The 41 preprocessing step which consists in creating the interfaces between the subdomains. 42 This is a purely geometrical task. We then present the coupling step which couples 43 the solution from the different meshes. Finally we show a numerical examples.

## 2 Interface Creation Process

The first task of the Chimera method is to create apparent interfaces between the 46 background and the patch meshes. This is achieved by the hole cutting step of the 47 Chimera method. As will be explained in next section, our coupling strategy requires 48 smooth interfaces. After the hole cutting, smoothing of the interfaces are also necessary. We now explain these two points.

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2.1 Hole Cutting

The hole cutting tasks consists in removing elements (the hole elements) from the 52 background mesh to form interfaces with the patches. We start by identifying the hole 53 nodes. The hole nodes are those nodes of the background mesh that are located inside 54 the patch mesh. To do this we have used a skd-tree strategy, as explained in [12]. Skd- 55 trees are used to find efficiently the signed shortest distance between a point and a 56 surface. In our case, the surfaces are the patch outer boundaries. In practice we obtain 57 a better efficiency if we use the search algorithm described in [18], which is a slightly 58 modified version of the above reference. Having found the hole nodes, we identify 59 the hole elements which are the background elements of which all nodes are hole 60 nodes. The fringe nodes are defined as the nodes located on the outer boundaries 61 of the hole elements. They are the hole nodes having non-hole neighbor nodes. The 62 fringe nodes are used to form the interface of the backgound with the patches.

2.2 Smoothing 64

The domain decomposition coupling we propose is geometrical, as will be shown in 65 next section. It is therefore important to ensure a minimum regularity of the interfaces 66 and the mesh nearby, as this will affect the quality of the results. Figure 1 (Left) 67 shows an example of typical background interface resulting from the previous hole 68 cutting process. The proposed strategy consists in smoothing first the interface and 69 then the volume mesh in the vicinity.

In this article, we are interested in mesh smoothing techniques that relocate the 71 nodes to improve the mesh without changing its topology. The particular method we 72 consider here is based on local mesh smoothing algorithms, since they have shown 73 to be efficient in repairing distorted elements. The most common smoothing tech- 74 nique is Laplacian smoothing (see [13]), which moves a given node to the barycenter 75 of all its connected nodes. This method is not computationally expensive but does 76 not guarantee an improvement in mesh quality. In addition, it can create invalid elements or poor quality elements resulting in convergence and shrinkage problems. To 78 overcome this shortcoming, different variations of Laplacian smoothing have been 79 proposed like [5, 22].

Optimization-based smoothing algorithms are alternative local smoothing strategies. These algorithms depend on the type of mesh, the optimization method used and a measure of the mesh quality, and require an objective function to be optimized. The objective function should include a good representation of the mesh quality. A good summary of measures for the quality of tetrahedra and a global definition functions described in the above reference, there exist other quality interpretations matrices and matrix norms. This matrix perspective suggests several different objective functions as, for example, the smoothness objective function in terms of the condition number of the Jacobian matrix; see [6].

Our smoothing process consists first of a surface Laplacian-smoothing algorithm 91 based on [21] for the interface. An example is shown in Fig. 1. As a consequence,

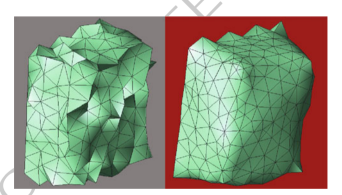


Fig. 1. (Left) Original interface after hole cutting. (Right) Smoothed interface

we need to relocate the volume nodes in order to repair the bad quality elements. To tackle this problem, we have applied a tetrahedra mesh improvement via optimization of the element condition number developed in [6]. This optimization uses a steepest descent method with a modified line search adapted to the geometrical seconstraints of the sub-mesh associated to the node we want to move. The implemented line search satisfies the Armijo rule which guarantees the local convergence of the method. For more details about this issue the reader can refer to [14]. Besides, a structured strategy is applied to perform the line search. The descent direction is obtained using the gradient of the objective function  $f(\mathbf{x})$ , in which the free vertex not (node)  $\mathbf{x}$  is the unknown:  $f(\mathbf{x}) = \|K(\mathbf{x})\|_2 = \left[\sum_{m=0}^{M-1} \kappa_m(\mathbf{x})^2\right]^{1/2}$ , where  $\kappa_m$  represents the condition number associated to the tetrahedron m, the moving node having M

sub-mesh elements. We then compute the steepest descent  $\mathbf{p} = -\nabla f$  and find the position which gives minimum  $f(\mathbf{x})$ . 105

3 DD-Coupling 106

The Chimera method can be viewed as an overlapping domain decomposition technique, where transmission conditions are imposed on the interfaces of the subdomains, see [17]. A key point of the Chimera method is the way the information 109 on the artificial boundaries is transferred, that is, the coupling. The different clas- 110 sical options depends on the type of the transmission conditions imposed on the 111 interfaces. The most typical are Dirichlet/Dirichlet (D/D) coupling, also known as 112 Schwarz' method, Dirichlet/Neumann (D/N) coupling, Dirichlet/Robin (D/R) cou- 113 pling, Robin/Robin(R/R) coupling. In the litterature, the coupled system is usu- 114 ally solved iteratively. In each subdomain  $\Omega_i$  local problems are solved by using 115 as boundary conditions (of Dirichlet or Robin type) the values form its neighbours 116  $\Omega_i$  until convergence is achieved. Relaxation is often needed to obtain this convergence and depends on the local character of the equation. In [8], the equivalence be- 118 tween the one-domain formulation and overlapping domain decomposition methods 119 of Dirichlet/Neumann(Robin) type is shown at the continuous level. The equivalence 120 is no longer true at the discrete level.

We have developed in this work a new way of coupling the subdomains that we 122 refer to as Extension-Dirichlet (Ext+D). The advantage of the method is that it is implicit and parallel. Therefore, no additional iterative loop is introduced and a-fortiori 124 the convergence of the method has no relation with the overlap. The idea consists 125 in extending the subdomains from their interfaces to their neighboring subdomains, 126 and imposing the Dirichlet condition implicitly, by connecting their extension to the 127 nodes of the neighbors. This method is equivalent, in practice, to imposing Dirichlet 128 boundary condition and eliminating it.

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To illustrate the method, let us solve a diffusion equation,  $\Delta u = 0$  using the 130 Galerking method in domain [0, 1] discretized in 4 linear elements, with the boundary conditions, u(0) = 1 and u(1) = 3. The analytical solutions is u = -2x + 1. Fig. 132 ure 2 (Left) shows the two unconnected subdomains and the corresponding assembled global matrix. Then, Fig. 2 (Center) shows, for the same example, the results of 134 an implicit Dirichelt/Dirichlet coupling. To achieve this,  $u_3 - u_5 = 0$  substitutes line 135 3 and  $u_4 - u_2 = 0$  substitutes line 4. The (Ext+D)<sup>2</sup> method we propose is illustrated 136 in Fig. 2 (Right). Starting with the matrix of Fig. 2 (Left), we perform the following: 137

- Extend node 3 shape function to node 6 of the second subdomain. This provides 138 additional terms in the equation for node 3.
- Extend node 4 shape function to node 1 of the second subdomain. This provides additional terms in the equation for node 4.

We can observe that in practice the  $(Ext+D)^2$  method creates new elements. In this 142 example the new elements are 3-6 and 4-1. The element matrices and RHS's are 143

#### An Implicit and Parallel Chimera Type Domain Decomposition Method

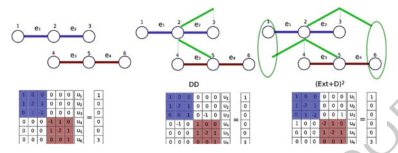


Fig. 2. (Left) Problem statement and domain. (Center) Dirichlet/Dirichlet assembled. (Right)  $(Ext+D)^2$ 

computed as any other elements of the mesh, but only the lines of node 3 and node 4 144 of these matrices and RHS's are assembled into the global matrix, respectively.

The main difficulty of the method is to be able to construct a proper extension 146 from one interface node to the other subdomain. This task is specially complex in 147 the 3D case, mainly due to the restriction that the extension must be closed. In vari- 148 ational terms, this means that the extension has a compact support. We are going to 149 describe the way to create the extensions on the interface  $\Gamma_{ij}$  between subdomain  $\Omega_{i}$  150 and subdomain  $\Omega_i$  in the 2D case. The process, illustrated in Fig. 3, consists in the

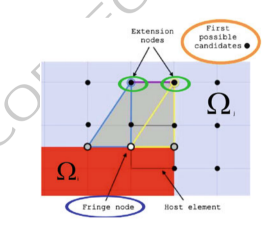


Fig. 3. 2D extensions

following.

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- For a fringe node of  $\Omega_i$ , identify the host element in  $\Omega_i$ .
- The nodes connected to this host element are the possible candidates to create 154 the triangles that form the associated extension. They are the black nodes. 155
- Construct two triangles (blue and yellow) connected to the boundaries of the 156 fringe node. 157

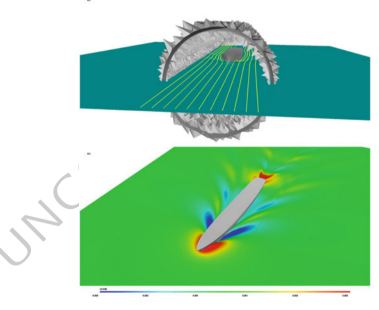
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The choice of the extension nodes (blue and yellow circled) is based on a quality 159 criterion of the resulting triangles [7], among all the possibilities for the previous list. 160 The third node of the triangle is the other node that forms the interface boundary.

## 4 Numerical Example

Figure 4 shows some results obtained for a flow around a boat. The Navier-Stokes 163 equations are solved together with a level set function and one-equation Spalart- 164 Allmaras turbulence model. The space discretization is a variational multiscale finite 165 element method. The complete description of the algorithm can be found in [10, 11, 166] 15] This complex case computed with 256 CPU's reflects the versatile property of 167 our method and its parallel capacity. The first figure shows the extension elements 168 while the second one the velocity module.



**Fig. 4.** (*Top*) Extension elements. (*Bottom*) Level set

5 Conclusions 170

We have devised in this paper a domain decomposition method, referred as  $(Ext+D)^2$  171 which is based on the explicit construction of extension elements assembled almost 172

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as any other element so that the implementation is straightforward. It consists in	17
imposing implicitly Dirichlet transmission conditions and does not introduce any	17
additional iterative loop to the algorithm. Another strength of the method is that it is	17
naturally parallel. However, aspects like conservation should be treated in order to	17
complete the analysis of the method.	17

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# Optimized Schwarz Waveform Relaxation for Porous Media Applications

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1 Introduction 10

Far field simulations of underground nuclear waste disposal involve a number of thallenges for numerical simulations: widely differing lengths and time-scales, 12 highly variable coefficients and stringent accuracy requirements. In the site under 13 consideration by the French Agency for Nuclear Waste Management (ANDRA), the 14 repository would be located in a highly impermeable geological layer, whereas the 15 layers just above and below have very different physical properties (see [1]). It is 16 then natural to use different time steps in the various layers, so as to match the time 17 step with the physics. To do this, we propose to adapt a global in time domain decomposition method, based on Schwarz waveform relaxation algorithms, to problems in heterogeneous media. This method has been introduced and analyzed for 20 linear advection-reaction-diffusion problems with continuous coefficients [2, 6] and 21 extended to discontinuous coefficients [3, 4], with asymptotically optimized Robin 22 transmission conditions in [3]. The method is extended to higher dimension in [4] 23 with convergence results and error estimates for rectangular or strip subdomains.

This method is extended to problems with discontinuous porosity in [5]. A new 25 aproach is proposed to determine optimized transmission conditions for domains 26 with highly variable lengths. In this paper we analyse this approach in 1d. 27

Our model problem for the radionuclide transport is the one dimensional advection 28 diffusion-reaction equation 29

$$\varphi \partial_t u + a \partial_x u - \partial_x (v \partial_x u) + bu = f, \quad \text{on } \mathbb{R} \times (0, T), u(0, x) = u_0(x), \quad x \in \mathbb{R}.$$
 (1)

We focus on a model problem to show the effect of subdomains with widely differing 30 sizes. We consider a decomposition in  $\Omega_1=(-\infty,0)$ ,  $\Omega_2=(0,L)$ ,  $\Omega_3=(L,\infty)$  with 31 L<<1. The reaction coefficient b is taken constant and the coefficients a, v, and  $\varphi$  32 are assumed constant on each  $\Omega_k$ , but may be discontinuous at x=0 and x=L, 33

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$$\varphi = \varphi_k, \quad a = a_k, \quad v = v_k, \quad x \in \Omega_k.$$

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We introduce the notations

$$\mathcal{L}_k v := \varphi_k \partial_t v + a_k \partial_x v - \partial_x (v_k \partial_x v) + bv, \quad \text{on } \Omega_k \times (0, T),$$
  
$$\boldsymbol{\varphi} := (\varphi_1, \varphi_2, \varphi_3), \boldsymbol{a} := (a_1, a_2, a_3), \boldsymbol{v} := (v_1, v_2, v_3).$$

Problem (1) is equivalent to solving problems in subdomains  $\Omega_k$ 

$$\mathcal{L}_k u_k = f$$
, on  $\Omega_k \times (0, T)$ ,  $u_k(0, x) = u_0(x)$ ,  $x \in \Omega_k$ .

with coupling conditions on the interface  $\Gamma_{k,\ell}$  between two neighboring subdomains 36  $\Omega_k$  and  $\Omega_\ell$  given by

$$u_k = u_\ell$$
,  $(v_k \partial_x - a_k) u_k = (v_\ell \partial_x - a_\ell) u_\ell$ , on  $\Gamma_{k,\ell} \times (0,T)$ . (2)

# 2 Domain Decomposition Algorithm

A simple algorithm based on relaxation of the coupling conditions (2) does not converge in general (see [7]). Following previous works [2–4], we introduce the Schwarz waveform relaxation algorithm

$$\mathcal{L}_{k}u_{k}^{n} = f, \quad \text{on } \Omega_{k} \times (0, T),$$

$$u_{k}^{n}(0, x) = u_{0}(x), \quad x \in \Omega_{k},$$

$$(v_{k}\partial_{x} - a_{k}) u_{k}^{n} + \mathcal{L}_{k,\ell}u_{k}^{n} = (v_{\ell}\partial_{x} - a_{\ell}) u_{\ell}^{n-1} + \mathcal{L}_{k,\ell}u_{\ell}^{n-1}, \quad \text{on } \Gamma_{k,\ell} \times (0, T),$$

$$(3)$$

where  $\mathcal{S}_{k,\ell}$  are linear operators in time and space, defined by

$$\mathscr{S}_{k,\ell} \pmb{\psi} = ar{p}_{k,\ell} \pmb{\psi} + ar{q}_{k,\ell} \partial_t \pmb{\psi}.$$

The case  $\bar{q}_{k,\ell} \neq 0$  corresponds to Robin transmission conditions, while the case 43  $\bar{q}_{k,\ell} \neq 0$  corresponds to first order transmission conditions. The well-posedness and 44 convergence have been analyzed for constant porosity in [3] and in higher dimension 45 in [4]. The transmission conditions in (3) imply the coupling conditions (2) at convergence, and lead at the same time to an efficient algorithm, for suitable parameters 47  $\bar{p}_{k,\ell}$  and  $\bar{q}_{k,\ell}$  obtained from an optimization of the convergence factor. 48

Similarly,  $\mathcal{S}_{k,\ell}$  are approximations of the best operators related to transparent 49 boundary operators. They can be found using Fourier analysis in the two half-spaces 50 case. This analysis has been done for discontinuous coefficients [3], and in higher 51 dimension and continuous coefficients [2]. The min-max problem has been analysed 52 in one dimension in [3] with asymptotical Robin parameters. 53

In the field of nuclear waste computations, domains of meter scale are embedded 54 in domains of kilometer scale. The previous optimization of the convergence factor 55 does not take into account the high variability of the domains lengths. Following 56 [5], we determine optimized transmission conditions through the minimization of a 57 convergence factor that takes into account this variability. 58

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## 2.1 Optimal Transmission Conditions

In order to determine the optimal transmission operators  $\mathcal{S}_{k,\ell}$ , we compute the convergence factor of the algorithm. Since the problem is linear, we consider the algorithm (3) on the error (i.e. with f=0 and  $u_0=0$ ). In order to use a Fourier transform for time, we assume that all functions are extended by 0 for t<0.

Let  $e_k^n = u_k^n - u$  be the error in  $\Omega_k$  at iteration k. The operators  $\mathscr{S}_{k,\ell}$  are related to 64 their symbols  $\sigma_{k,\ell}(\omega)$  by

$$\mathscr{S}_{k,\ell}u(t)=rac{1}{2\pi}\int \sigma_{k,\ell}(\omega)\hat{u}(\omega)e^{i\omega t}d\omega.$$

The Fourier transforms  $\hat{e}_k^n$  in time of  $e_k^n$  are solutions of the ordinary differential 66 equation in the x variable 67

$$-v_k\partial_{xx}^2\hat{e}+a_k\partial_x\hat{e}+(i\varphi_k\omega+b)\hat{e}=0.$$

The characteristic roots are

$$r^{\pm}(a_k, v_k, \varphi_k, b, \omega) = \frac{a_k \pm \sqrt{d_k}}{2v_k}, \quad d_k = a_k^2 + 4v_k(i\varphi_k\omega + b). \tag{4}$$

Since  $\Re r^+ > 0$ ,  $\Re r^- < 0$ , and since we look for solutions which do not increase 69 exponentially in |x|, we obtain 70

$$\hat{e}_{1}^{n}(x,\omega) = \alpha_{1}^{n}(\omega)e^{r^{+}(a_{1},v_{1},\phi_{1},b,\omega)x}, \quad \hat{e}_{3}^{n}(x,\omega) = \alpha_{3}^{n}(\omega)e^{r^{-}(a_{3},v_{3},\phi_{3},b,\omega)x}, \\ \hat{e}_{2}^{n}(x,\omega) = \alpha_{2}^{n}(\omega)e^{r^{+}(a_{2},v_{2},\phi_{2},b,\omega)x} + \beta_{2}^{n}(\omega)e^{r^{-}(a_{2},v_{2},\phi_{2},b,\omega)x}.$$
(5)

We set  $\xi^n=(\alpha_1^n,\alpha_2^n,\beta_2^n,\alpha_3^n)^t$ , and  $r_k^\pm=r^\pm(a_k,\nu_k,\phi_k,b,\omega)$ . We define the variables 71  $s_k=s_k(\omega,L),\ 1\leq k\leq 8$ , by

$$s_{1} = \frac{v_{2}r_{2}^{-} - \sigma_{1,2}}{v_{1}r_{1}^{-} - \sigma_{1,2}}, \ s_{2} = \frac{v_{2}r_{2}^{+} - \sigma_{1,2}}{v_{1}r_{1}^{-} - \sigma_{1,2}}, \ s_{3} = \frac{v_{2}r_{2}^{+} - \sigma_{2,3}}{v_{2}r_{2}^{-} - \sigma_{2,3}} \cdot e^{(r_{2}^{-} - r_{2}^{+})L},$$

$$s_{5} = \frac{v_{2}r_{2}^{-} + \sigma_{2,1}}{v_{2}r_{2}^{+} + \sigma_{2,1}}, \ s_{7} = \frac{v_{2}r_{2}^{-} + \sigma_{3,2}}{v_{3}r_{3}^{+} + \sigma_{3,2}} e^{(r_{2}^{+} - r_{3}^{-})L}, \ s_{8} = \frac{v_{2}r_{2}^{+} + \sigma_{3,2}}{v_{3}r_{3}^{+} + \sigma_{3,2}} e^{(r_{2}^{-} - r_{3}^{-})L},$$

$$s_{4} = \frac{v_{1}r_{1}^{-} + \sigma_{2,1}}{v_{2}r_{2}^{+} + \sigma_{2,1}} \cdot \frac{1}{D}, \ s_{6} = \frac{v_{3}r_{3}^{+} - \sigma_{2,3}}{v_{2}r_{3}^{-} - \sigma_{3,2}} \cdot \frac{e^{(r_{3}^{-} - r_{2}^{+})L}}{D}, \ \text{with } D = s_{3}s_{5} - 1.$$

We insert (5) into the transmission conditions in (3), and obtain for  $n \ge 2$ ,

$$\xi^n = M\xi^{n-1},$$

where the matrix  $M = M(\omega, L)$  is defined by

$$M = \begin{pmatrix} 0 & s_1 & s_2 & 0 \\ s_3 s_4 & 0 & 0 & -s_6 \\ -s_4 & 0 & 0 & s_5 s_6 \\ 0 & s_7 & s_8 & 0 \end{pmatrix}.$$

The convergence factor  $\rho(\omega, L)$  for each  $\omega \in \mathbb{R}$  is the spectral radius of M.

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*Remark 1*. The choice for the symbols  $\sigma_{k,\ell}$ 

$$\sigma_{1,2} = v_2 r_2^+, \quad \sigma_{2,1} = -v_1 r_1^-, \quad \sigma_{2,3} = v_3 r_3^+, \quad \sigma_{3,2} = -v_2 r_2^-,$$
 (6)

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leads to  $M^2 = 0$  and thus to optimal convergence in three iterations.

**Proposition 1.** The convergence factor is given by

$$\rho(\omega, L) = \sqrt{\max(|\lambda^-|, |\lambda^+|)},$$

where  $\lambda^{\pm} = \lambda^{\pm}(\omega, L)$  is defined by

$$\lambda^{\pm} = \frac{\alpha + \beta \pm \sqrt{(\alpha - \beta)^2 + 4\gamma\zeta}}{2},$$

with 80

$$\alpha = s_1 s_3 s_4 - s_2 s_4, \; \beta = -s_6 s_7 + s_5 s_6 s_8, \; \gamma = s_3 s_4 s_7 - s_4 s_8, \; \zeta = -s_1 s_6 + s_2 s_5 s_6.$$

This follows from the computation of the roots of the characteristic polynomial of M, which is biquadratic. The corresponding operators to (6) are non-local in time. In the next subsection, we therefore approximate the optimal operators by local ones.

## 2.2 Local Transmission Conditions

We approximate the optimal choice  $\sigma_{k\ell}$  in (6) by polynomials in  $\omega$ :

$$\begin{split} &\sigma_{1,2}^{\mathrm{app}} = \frac{p_{1,2} + a_2}{2} + \frac{q_{1,2}}{2}i\omega, \quad \sigma_{2,1}^{\mathrm{app}} = \frac{p_{2,1} - a_1}{2} + \frac{q_{2,1}}{2}i\omega, \\ &\sigma_{2,3}^{\mathrm{app}} = \frac{p_{2,3} + a_3}{2} + \frac{q_{2,3}}{2}i\omega, \quad \sigma_{3,2}^{\mathrm{app}} = \frac{p_{3,2} - a_2}{2} + \frac{q_{3,2}}{2}i\omega. \end{split}$$

In order to simplify the min-max problem, we will consider the following cases for 87 the choice of  $p_{k,\ell}$  and  $q_{k,\ell}$ :

- 1. (Robin)  $p_{k,\ell} = p, q_{k,\ell} = 0,$
- 2. (Zeroth order)  $p_{1,2} = p_{3,2} = p_1$ ,  $p_{2,1} = p_{2,3} = p_2$ ,  $q_{k\ell} = 0$ ,
- 3. (First order)  $p_{k,\ell} = p, q_{k,\ell} = q,$
- 4. (First order scaled)  $p_{k,\ell} = p$ ,  $q_{1,2} = \varphi_2 q$ ,  $q_{2,1} = \varphi_1 q$ ,  $q_{2,3} = \varphi_3 q$ ,  $q_{3,2} = \varphi_2 q$ .

Then, the parameters are chosen in order to minimize the convergence factor, i.e. we 93 solve, for  $\mathbf{p} = p$  in case 1,  $\mathbf{p} = (p_1, p_2)$  in case 2, and  $\mathbf{p} = (p, q)$  in cases 3 and 4, the 94 min-max problem 95

$$\delta_{m}(L) = \min_{\mathbf{p}} \left( \max_{\omega_{0} \le \omega \le \omega_{\max}} \rho(\omega, \mathbf{p}, \boldsymbol{\varphi}, \boldsymbol{a}, \mathbf{v}, b, L) \right), \tag{7}$$

where  $\rho$  is the spectral radius of M, in which we have replaced  $\sigma_{k,\ell}$  by  $\sigma_{k,\ell}^{\rm app}$ , and m 96 is the order of the approximation. In numerical computations, the frequencies can be 97 restricted to  $\omega_{\rm max} = \frac{\pi}{\Delta t}$ , where  $\Delta t$  is the time step, and  $\omega_0 = \frac{\pi}{T}$ . 98

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**Theorem 1.** We suppose that  $a_k = a$ ,  $\varphi_k = \varphi$  and  $v_k = v$ ,  $1 \le k \le 3$ , thus  $d_k = d$  in 99 (4). Let us consider the Robin case ( $\mathbf{p} = p$ ) and the first order case ( $\mathbf{p} = (p,q)$ ). Then 100 the convergence factor reduces to

$$\rho(\omega, \mathbf{p}, \varphi, a, v, b, L) = \sqrt{\left|\frac{\sigma - \sqrt{d}}{\sigma + \sqrt{d}}\right| \max\left(\left|\frac{\sigma - \mu}{\sigma + \mu}\right|, \left|\frac{\sigma - \eta}{\sigma + \eta}\right|\right)}$$

with

$$\mu = \sqrt{d} \left( \frac{1 + e^{-\frac{\sqrt{d}}{2\nu}L}}{1 - e^{-\frac{\sqrt{d}}{2\nu}L}} \right), \ \eta = \frac{\sqrt{d}}{\mu},$$

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and with  $\sigma = p$  in the Robin case, and  $\sigma = p + qi\omega$  in the first order case. Let L > 0 103 given. Let  $\delta_0(L)$  (resp.  $\delta_1(L)$ ) be the solution of (7) for the Robin case (resp. the first 104 order case). For m = 0 and m = 1, we have  $|\delta_m(L)| < 1$ .

## 3 Numerical Results

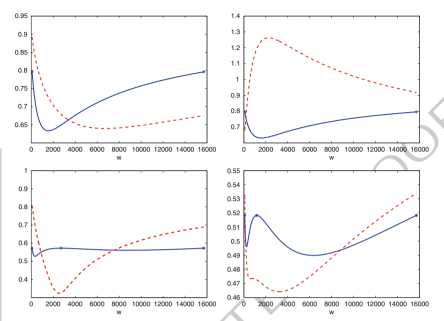
We use the DG-OSWR method in [4] based on a discontinuous Galerkin method in time, with  $\mathbf{P}_1$  finite elements in space in each subdomain. We present an example 108 inspired from nuclear waste simulations, with discontinuous coefficients, and different time and space steps in the subdomains  $\Omega_2 = (0.4954, 0.5047)$  (repository), 110  $\Omega_1 = (0,0.4954)$  and  $\Omega_3 = (0.5047,1)$  (host rock). The parameters for the three 111 subdomains are shown in Table 1. The final time is T=0.04.

<b>/</b>	φ			b		$\Delta t$
$\Omega_1 \cup \Omega_3$	0.06					$T(510^{-3})$
$\Omega_2$	0.1	1	1	0	$510^{-4}$	$T(110^{-3})$

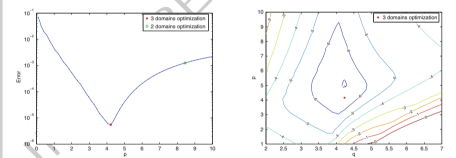
Table 1. Physical and numerical parameters

Let  $p_3^{\star}$  (resp.  $p_2^{\star}$ ) be the parameters derived from a numerical minimization of the three domains convergence factor in (7) (resp. from the two half-spaces convergence factor in [3]). Figure 1 shows  $\rho(\omega, p_3^{\star}, L)$  (solid line) and  $\rho(\omega, p_2^{\star}, L)$  (dashed line) the versus  $\omega$  for  $\Delta t = T(5 \ 10^{-3})$ . We observe that the solution of (7) is characterized the square equioscillation property (at the star marks), as in the two half-spaces case (see fig. [2]). Moreover, for first order transmission conditions, we see that a scaling with the porosity is important only when the parameters are computed from the two half-spaces analysis.

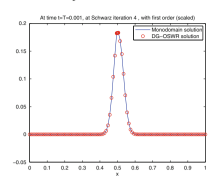
On Fig. 2 we show the error after 20 iterations when running the algorithm on the discretized problem, with  $u_0=f=0$  and random initial guess on the interfaces, for various values of the Robin parameter p (left) and the zeroth order parameters  $p_1, p_2$  123 (right) (in that case, the values obtain with the two half-spaces analysis is not in the range values of the figure).



**Fig. 1.** Convergence factor  $\rho(\omega, \mathbf{p}_3^*, L)$  (*solid line*) and  $\rho(\omega, \mathbf{p}_2^*, L)$  (*dashed line*) versus  $\omega$ : *Top left*: Robin, *top right*: zeroth order, *left bottom*: first order, *right bottom*: first order scaled



**Fig. 2.** Error after 20 iterations: *Left*: for various values of the Robin parameter p (the *lower left star marks*  $p_3^*$  whereas the *upper right circle* shows  $p_2^*$ ), *Right*: the level curves for various values of the zeroth order parameters  $p_1$ ,  $p_2$  (the *star marks* the parameter  $p_3^*$ )



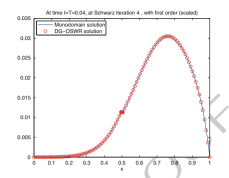
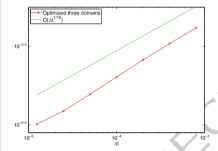
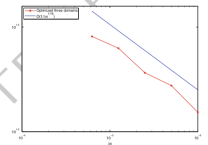


Fig. 3. Evolution of the monodomain solution (solid line) and the OSWR solution at iteration 4 (*circle line*): at t = 0.001 (*left*), t = T = 0.04 (*right*)





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**Fig. 4.** Asymptotic behavior as the mesh is refined: on the *left*  $R(\Delta t)$  and on the *right* where  $\Delta t = O(\Delta x)$ , the rate for the optimized Schwarz waveform relaxation algorithm with optimized first order (scaled) transmission conditions

On Fig. 3, the solution, with first order (scaled) conditions, at iteration 4 is shown 126 for an initial condition equal to 1 in  $\Omega_2$  and 0 elsewhere.

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Figure 4 shows on the left  $R(\Delta t) = 1 - \max_{\pi/T \le \omega \le \pi/\Delta t} \rho(\omega, \mathbf{p}_3^*, L))$  versus  $\Delta t$ , i.e. the convergence factor behaves like  $1 - O(\Delta t)^{1/16}$ , with first order (scaled) optimized transmission conditions. On the right, we run the OSWR algorithm until the 130 error becomes smaller than  $10^{-11}$ , and count the number of iterations. We start with 131  $\Delta t = T/100$  in each subdomain, and repeat this experiment dividing  $\Delta x$  and  $\Delta t$  by 2 132 several times. We observe that the asymptotic result on the left predicts very well the 133 numerical behavior of the algorithm given on the right.

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## **AUTHOR QUERY**

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# On Block Preconditioners for Generalized Saddle **Point Problems**

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1 Introduction 7

We consider a symmetric system of linear equations with a block structure,

$$\mathcal{M} \begin{pmatrix} u \\ p \end{pmatrix} \equiv \begin{pmatrix} A & B^T \\ B & -C \end{pmatrix} \begin{pmatrix} u \\ p \end{pmatrix} = \begin{pmatrix} F \\ G \end{pmatrix}. \tag{1}$$

We assume that A is  $n \times n$  and C is an  $m \times m$  matrix. Many such systems arise from 9 the discretization of (systems of) partial differential equations. For example, Stokes 10 equations discretized with stable finite elements or a mixed finite element method 11 for second order elliptic PDEs lead to a positive definite matrix A and to C=0, so 12 that (1) has a genuine saddle point structure. Certain other PDE problems may result 13 in an indefinite matrix A, or a semidefinite matrix A with a large kernel, which gives 14 (1) the structure of a so called generalized saddle point problem. Linear elasticity 15 equations modelling nearly incompressible materials discretized with mixed finite 16 elements result in both matrices A and C being positive definite, having thus a nature 17 of a penalized saddle point problem. All systems mentioned above have a common 18 feature that the matrix of (1) is indefinite.

The specific structure of (1) makes it possible to design efficient solution methods 20 which intensively exploit the properties of the system, see the recent survey of [4] 21 on the state-of-the-art in this field. Systems derived from the discretization of PDEs 22 are usually very large and sparse, and typically are solved by some iterative method. 23 Unfortunately, these systems are ill-conditioned with respect to the mesh size h, so 24 preconditioning is necessary in order to keep the number of iterations within a rea- 25 sonable limit. Applying a left preconditioner  $\mathscr{P}$ , one then solves a problem with a 26 preconditioned matrix  $\mathscr{P}^{-1}\mathcal{M}$ . We shall consider preconditioners of the form

$$\mathscr{P}_{d} = \begin{pmatrix} I \\ cBA_0^{-1} I \end{pmatrix} \begin{pmatrix} A_0 \\ S_0 \end{pmatrix} \begin{pmatrix} I dA_0^{-1}B^T \\ I \end{pmatrix}$$
 (2)

or

$$\mathscr{P}_{p} = \begin{pmatrix} I dB^{T} S_{0}^{-1} \\ I \end{pmatrix} \begin{pmatrix} A_{0} \\ S_{0} \end{pmatrix} \begin{pmatrix} I \\ c S_{0}^{-1} B I \end{pmatrix}, \tag{3}$$

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where  $A_0$  and  $S_0$  are symmetric, positive (or negative) definite matrices whose inverses are easy to apply and  $c,d \in \{-1,+1\}$ . In accordance with [8], we will refer 30 to  $\mathscr{P}_{d}$  as the family of dual block preconditioners and to  $\mathscr{P}_{p}$  as the family of primal 31 block preconditioners.

Many popular block preconditioners can be formed by choosing appropriate values of c and d in the formulas above. For example, a block diagonal preconditioner, 34 cf. e.g. [2, 6, 9, 13, 19, 21] corresponds to c = d = 0 above. Block triangular preconditioners considered e.g. in [7, 14, 22] and the Bramble-Pasciak preconditioner 36 as well, see [5], are obtained with either c or d equal to zero. The choice c = d = 1 37 in (2) produces a symmetric indefinite preconditioner, see [3, 20, 24, 25], while the 38 same choice in (3) leads to a primal based penalty preconditioner, [1, 8].

It is straightforward that solving a system with  $\mathcal{P}_d$  requires one solve with  $S_0$  and 40 at most two solves with  $A_0$ , while applying  $\mathscr{P}_p$  to a vector takes one solve with  $A_0$  41 and at most two solves with  $S_0$ . When cd = 0, both types of preconditioners require 42 only one solve with  $A_0$  and one with  $S_0$ .

Let us stress that when (1) arises from finite element discretization of PDEs, there 44 is a possibility to use other than block preconditioning approaches. On the other 45 hand, for many types of discretizations and problems, specialized methods based 46 on direct construction of a multigrid or domain decomposition preconditioner— 47 although usually outperforming block preconditioners, [15]—may take a consid-48 erable effort to develop, implement and analyse. Since the block preconditioning 49 approach as discussed here turns out to be based on preconditioners for symmet- 50 ric positive definite matrices, this property makes it a viable and robust alternative to 51 custom methods, as in this case one can efficiently reuse existing theory and software 52 to solve more complex problems. This feature has been recognized in the software 53 package PETSc, see [23], where a family of so called field-splitting preconditioners 54 has recently been implemented.

# 2 Eigenvalue Estimates of the Preconditioned System

Eigenvalue clustering is vital for the convergence of a Krylov method, so it is im- 57 portant to bound the spectrum of  $\mathscr{P}^{-1}\mathcal{M}$ , where  $\mathscr{P}$  stands for either  $\mathscr{P}_d$  or  $\mathscr{P}_p$ . 58 Inspired by the block nature of the problem, which imposes a decomposition of the 59 unknowns into two parts  $(u, p) \in \mathbb{R}^n \times \mathbb{R}^m$ , let us define a block diagonal, symmetric, 60 positive definite matrix

> $\mathscr{J} = \begin{pmatrix} \tilde{A}_0 \\ \tilde{S}_0 \end{pmatrix},$ 62

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where  $A_0$  is either  $A_0$ , if  $A_0$  is positive definite, or  $(-A_0)$ , if  $A_0$  is negative definite; 63  $\hat{S}_0$  is defined in the same way. We assume there exist positive constants  $m_0$  and  $m_1$  64 such that

$$m_0||x||_{\mathscr{I}} \le ||\mathscr{M}x||_{\mathscr{I}^{-1}} \le m_1||x||_{\mathscr{I}} \qquad \forall x \in \mathbb{R}^n \times \mathbb{R}^m,$$

where 67

$$||\binom{u}{p}||_{\mathscr{J}}^2 = ||u||_{\tilde{A}_0}^2 + ||p||_{\tilde{S}_0}^2,$$
 68

This is nothing but a stability and continuity assumption in an appropriate norm, see 69 also [18]. At the same time we suppose there exists a constant  $b_0$  such that for any 70  $u \in \mathbb{R}^n$  and  $p \in \mathbb{R}^m$ ,

$$|p^T B u| \le b_0 ||u||_{\tilde{A}_0} ||p||_{\tilde{S}_0}.$$

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Finally, we assume that for some  $\delta \in \{-1, +1\}$ , the matrix  $\mathcal{H}$  is positive definite, 73 where  $\mathcal{H}$  is equal to either  $\mathcal{H}_d$  or  $\mathcal{H}_p$  (depending on whether we are addressing  $\mathcal{P}_d$ or  $\mathscr{P}_{p}$ ), with

$$\mathcal{H}_{d} = \delta \begin{pmatrix} A_0 - cA \\ S_0 + cdBA_0^{-1}B^T + dC \end{pmatrix},$$
 76

$$\mathcal{H}_{d} = \delta \begin{pmatrix} A_0 - cA \\ S_0 + cdBA_0^{-1}B^T + dC \end{pmatrix},$$

$$\mathcal{H}_{p} = \delta \begin{pmatrix} A_0 + cdB^TS_0^{-1}B - cA \\ S_0 + dC \end{pmatrix}.$$
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It turns out that then both  $\mathcal{H}_d\mathcal{P}_d^{-1}\mathcal{M}$  and  $\mathcal{H}_p\mathcal{P}_p^{-1}\mathcal{M}$  are symmetric and the 79 eigenvalues of the preconditioned matrix are bounded as stated in the following theorem, whose proof appeared in [16]: 81

**Theorem 1.** Suppose the above assumptions are fulfilled. If  $\lambda$  is an eigenvalue of 82  $\mathcal{P}_{d}^{-1}\mathcal{M}$  or of  $\mathcal{P}_{p}^{-1}\mathcal{M}$ , then it is real and satisfies

$$\frac{m_0}{2(1+b_0^2)} \le |\lambda| \le 2m_1(1+b_0^2).$$

Let us mention that earlier Klawonn [12] proved a similar result for block diagonal preconditioning matrices. 86

## 2.1 Example Application: Stabilized Stokes Equations

Theorem 1 relies on the stability of (1) and therefore indicates that block preconditioners can be used also in the case when the inf-sup condition is not satisfied and one 89 uses a so called stabilized method. As a model example let us consider a stabilized 90  $Q_1 - Q_1$  discretization of Stokes equations 91

$$-\Delta u + \nabla p = f,$$
$$\nabla \cdot u = 0.$$

Let  $\mathscr{T}_h$  denote a shape-regular, quasi-uniform triangulation of a polygonal  $\Omega \subset \mathfrak{g}_2$  $R^2$  into quadrilaterals. Define the finite dimensional spaces of bilinear finite elements: 93

$$V_h = \{v \in [H^1_0(\Omega)]^2 : v_{|_{\mathbf{K}}} \in [Q_1(\kappa)]^2 \quad \forall \kappa \in \mathscr{T}_h\}$$
 94

and 95

$$W_h = \{ q \in L_0^2(\Omega) \cap C(\Omega) : q_{|_{\kappa}} \in Q_1(\kappa) \quad \forall \kappa \in \mathcal{T}_h \},$$
 96

where  $Q_1(\kappa)$  denotes the space of bilinear functions on  $\kappa$ . Since  $V_h$  and  $W_h$  do not 97 satisfy the inf-sup condition the following stabilized discretization has been intro- 98 duced in [11]:

$$\begin{cases} (\nabla u_h, \nabla v_h)_{L^2(\Omega)} - (\operatorname{div} v_h, p_h)_{L^2(\Omega)} & \forall v_h \in V_h, \\ -(\operatorname{div} u_h, q_h)_{L^2(\Omega)} - c(p_h, q_h) = -\tau \sum_{\kappa \in \mathscr{T}_h} h_{\kappa}^2(f, \nabla q_h)_{L^2(\kappa)} & \forall q_h \in W_h, \end{cases}$$
(4)

where

$$c(p_h,q_h)= au\sum_{\mathbf{K}\in\mathscr{T}_h}h_{\mathbf{K}}^2(\nabla\,p_h,\,
abla\,q_h)_{L^2(\mathbf{K})}$$
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and  $\tau > 0$  is some prescribed parameter, independent of h. As the above system 102 is stable and continuous in the norm  $\left(||u||_{H_0^1}^2 + ||p||_{L^2}^2\right)^{1/2}$ , one concludes that an 103 optimal preconditioner (with respect to the mesh size h) can be obtained with either 104  $\mathscr{P}_{
m d}$  or  $\mathscr{P}_{
m p}$ , where  $ilde{A}_0$  is spectrally equivalent to the discrete Lapacian operator and 105  $\tilde{S}_0$  is spectrally equivalent to the pressure mass matrix. These operators may require some pre-scaling in order to make either  $\mathcal{H}_d$  or  $\mathcal{H}_d$  positive definite. 107

## **Numerical Experiments**

We confirm the above findings running experiments for a stabilized  $Q_1 - Q_1$  discretization of the Stokes system on a unit square, obtained under MATLAB with the 110 software package IFISS 2.2, see [10].

We investigated the number of iterations of the preconditioned conjugate residual method required to reduce the residual norm by a factor of 10<sup>6</sup>. We experimented 113 with  $\mathcal{P}_d$  having one of the following forms: block diagonal (c = 1, d = 0), upper triangular (c = 0, d = 1) and lower triangular (c = d = 0) (see [17] for implementation 115 details) for varying mesh size h. The results for the case when  $A_0 = A$  and  $S_0 = M$ (as suggested by the above analysis) are provided below, confirming a convergence 117 rate independent of h: 118

n+m	243	867	3,267	12,675	49,923
Lower triangular	17	21	21	22	23
Upper triangular	16	16	16	16	16
Diagonal	32	35	37	39	39

In order to show a more realistic choice of  $A_0$ , we used  $A_0^{-1}$  defined by means of the incomplete Cholesky factorization of A, with drop tolerance  $10^{-3}$ . Since for our 121 model problem the quality of the incomplete Cholesky factorization degrades slowly 122 with increasing size of the system, this is also reflected in an increase of the iteration 123 counts: 124

n+m	243	867	3,267	12,675	49,923
Lower triangular			24	35	113
Upper triangular	17	17	20	33	_
Diagonal	33	38	48	74	132

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It has been observed that (at least in our implementation) the best solution times were obtained mostly for triangular preconditioners.	126 127
3 Conclusions	128
We have presented two classes of block preconditioners for symmetric saddle point	129
problems and provided eigenvalue estimates of the preconditioned system $\mathscr{P}^{-1}\mathcal{M}$	130
under a quite general assumption of the stability and continuity of the problem being	131
solved. In the context of PDEs, based upon this result, an iterative method, optimal	132
with respect to the mesh size $h$ , can be designed, which may reuse existing state-of-	133
the-art preconditioners or fast solvers for certain elliptic problems.	134
Acknowledgments The research has been partially supported in part by Polish Ministry of	135
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# **Optimal Control of the Convergence Rate of Schwarz Waveform Relaxation Algorithms**

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**Summary.** In this study we present a non-overlapping Schwarz waveform relaxation method 13 applied to the one dimensional unsteady diffusion equation. We derive efficient interface conditions using an optimal control approach once the problem is discretized. Those conditions 15 are compared to the usual optimized conditions derived at the PDE level by solving a min-max 16 problem. The performance of the proposed methodology is illustrated by numerical experiments.

1 Introduction 19

Schwarz-like domain decomposition methods are very popular in mathematics, com- 20 putational sciences, and engineering notably for the implementation of coupling 21 strategies. This type of method, originally introduced for stationary problems, can 22 be extended to evolution problems by adapting the waveform relaxation algorithms 23 to provide the so-called Schwarz waveform relaxation method [2, 4]. The idea behind 24 this method is to separate the spatial domain, over which the time-evolution problem 25 is defined, into subdomains. The resulting time-dependent problems are then solved 26 separately on each subdomains. An iterative process with an exchange of boundary 27 conditions at the interface between the subdomains is then applied to achieve the 28 convergence to the solution of the original problem. To accelerate the convergence 29 speed of the iterative process, it is possible to derive efficient interface conditions by 30 solving an optimization problem related to the convergence rate of the method [e.g.; 31 1, 5].

In this study, we specifically address the optimization problem arising from the 33 use of Robin type transmission conditions in the framework of a non-overlapping 34 Schwarz waveform relaxation. For this type of problem, the existing work has been 35 achieved mainly at the PDE level, giving rise to the optimized Schwarz waveform 36

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relaxation algorithm [1, 2, 5]. The objective here is to use the *optimal control theory* 37 paradigm [9] to find parameters optimized at the discrete level, and thus to systematically make a comparison with the parameters determined at the PDE level. This 39 paper is organized as follows: in Sect. 2 we briefly recall the basics of optimized 40 Schwarz methods in the framework of a time evolution problem. Section 3 is dedi- 41 cated to the determination of the optimal control problem that we intend to address. 42 Finally, in Sect. 4 we apply our approach to a diffusion problem.

# 2 Optimization of the Convergence at the PDE Level

## 2.1 Model Problem and Optimized Schwarz Methods

Let us consider  $\Omega$  a bounded open set of  $\mathbb{R}$ . The model problem is to find u such that 46 u satisfies over a time period [0, T]

$$\mathcal{L}u = f,$$
 in  $\Omega \times [0, T],$  (1)  
 $\mathcal{B}u = g,$  on  $\partial \Omega \times [0, T],$  (2)

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$$\mathscr{B}u = g, \quad \text{on } \partial\Omega \times [0, T],$$
 (2)

where  $\mathcal{L}$  and  $\mathcal{B}$  are two partial differential operators, and f the forcing. This problem is complemented by an initial condition

$$u(x,0) = u_0(x), \qquad x \in \Omega. \tag{3}$$

We consider a splitting of the domain  $\Omega$  into two non-overlapping domains  $\Omega_1$  and 50  $\Omega_2$  communicating through their common interface  $\Gamma$ . The operator  $\mathscr L$  introduced 51 previously is split into two operators  $\mathcal{L}_j$  restricted to  $\Omega_j$  (j=1,2). By noting  $\mathcal{F}_1$ , 52  $\mathscr{F}_2$ ,  $\mathscr{G}_1$  and  $\mathscr{G}_2$  the operators defining the interface conditions, the alternating form 53 of the Schwarz waveform relaxation algorithm reads

$$\begin{cases} \mathcal{L}_{1}u_{1}^{k} = f_{1}, & \text{in } \Omega_{1} \times [0,T], \\ u_{1}^{k}(x,0) = u_{o}(x), & x \in \Omega_{1}, \\ \mathcal{B}_{1}u_{1}^{k}(x,t) = g_{1}, & \text{in } [0,T] \times \partial \Omega_{1}, \\ \mathcal{F}_{1}u_{1}^{k}(0,t) = \mathcal{F}_{2}u_{2}^{k-1}(0,t), & \text{in } \Gamma \times [0,T], \end{cases} \begin{cases} \mathcal{L}_{2}u_{2}^{k} = f_{2}, & \text{in } \Omega_{2} \times [0,T], \\ u_{2}^{k}(x,0) = u_{o}(x), & x \in \Omega_{2}, \\ \mathcal{B}_{2}u_{2}^{k}(x,t) = g_{2}, & \text{in } [0,T] \times \partial \Omega_{2}, \\ \mathcal{G}_{2}u_{2}^{k}(0,t) = \mathcal{G}_{1}u_{1}^{k}(0,t), & \text{in } \Gamma \times [0,T], \end{cases}$$

where k = 1, 2, ... is the iteration number, and the initial guess  $u_2^0(0, t)$  must be given. 56 The operators  $\mathscr{F}_i$  and  $\mathscr{G}_i$  must be chosen to impose the desired consistency of the 57 solution on the interface  $\Gamma$ . We consider here the one-dimensional diffusion equation 58 with constant (possibly discontinuous) diffusion coefficients  $\kappa_j$  ( $\kappa_j > 0, j = 1, 2$ ). We 59 define  $\mathcal{L}_{j} = \partial_{t} - \kappa_{j} \partial_{x}^{2}$ ,  $\Omega_{1} = (-L_{1}, 0)$ ,  $\Omega_{2} = (0, L_{2})$   $(L_{1}, L_{2} \in \mathbb{R}^{+})$ , and  $\Gamma = \{x = 0\}$ . 60 In this context, we require the equality of the subproblems solutions and of their 61 normal fluxes on the interface  $\Gamma$ ,

$$u_1(0,t) = u_2(0,t), \qquad \kappa_1 \partial_x u_1(0,t) = \kappa_2 \partial_x u_2(0,t), \qquad t \in [0,T].$$
 (5)

To obtain such a consistency we use mixed boundary conditions of *Robin* type

$$\mathscr{F}_{j} = -\kappa_{j}\partial_{x} + p_{1}, \qquad \mathscr{G}_{j} = \kappa_{j}\partial_{x} + p_{2}, \qquad (j = 1, 2),$$

where  $p_1$  and  $p_2$  are two parameters that can be optimally chosen to improve the 65 convergence speed of the Schwarz method. Algorithm (4) with two-sided Robin 66 conditions (i.e. for  $p_1 \neq p_2$ ) is well-posed for any choice of  $p_1$  and  $p_2$  such that 67  $p_1 + p_2 > 0$ . This result can be shown using a priori energy estimates, as described 68 in [4].

#### 2.2 Optimization of the Convergence Factor

To demonstrate the convergence of algorithm (4) a classical approach [e.g. 6] is to 71 define the error  $e_j^k$  between the exact solution  $u^*$  and the iterates  $u_j^k$ . A Fourier anal-72 ysis enables the transformation of the original PDEs into ODEs that can be solved 73 analytically. The analytical solution on each subdomain is then used to define a convergence factor  $\rho$  of the corresponding *Schwarz algorithm*. For a diffusion problem, 75 defined on subdomains of infinite size (i.e. assuming  $L_1, L_2 \rightarrow \infty$ ), we get

$$\rho(p_1, p_2, \omega) = \left| \frac{(p_2 - \sqrt{i\omega\kappa_2})}{(p_2 + \sqrt{i\omega\kappa_1})} \frac{(p_1 - \sqrt{i\omega\kappa_1})}{(p_1 + \sqrt{i\omega\kappa_2})} \right|, \tag{6}$$

where  $p_1$  and  $p_2$  are two degrees of freedom which can be tuned to accelerate the 77 convergence speed. In (6),  $i = \sqrt{-1}$ , and  $\omega \in \mathbb{R}$  is the angular frequency arising from 78 a Fourier transform in time on  $e_j^k$ . A general approach to choose the Robin parameters 79  $p_1$  and  $p_2$  is to solve a minimax problem [2]

$$\min_{p_1, p_2 \in \mathcal{R}} \left( \max_{\omega \in [\omega_{\min}, \omega_{\max}]} \rho(p_1, p_2, \omega) \right). \tag{7}$$

Because we work in practice on a discrete problem the frequencies allowed by the 81 temporal grid range from  $\omega_{\min} = \pi/T$  to  $\omega_{\max} = \pi/\Delta t$ , where  $\Delta t$  is the time step 82 of the temporal discretization. For the diffusion problem under consideration here, 83 the analytical solution of the optimization problem (7) has been derived in [8] in a 84 general *two-sided* case (i.e. with  $p_1 \neq p_2$ ) with discontinuous coefficients  $\kappa_1 \neq \kappa_2$ . 85 For the sake of simplicity, we consider in the present study the continuous case ( $\kappa_1 = \kappa_2 = \kappa$ ) and we recall the result found in [8] in this case.

**Theorem 1.** Under the assumption  $\kappa_1 = \kappa_2 = \kappa$ , the optimal parameters  $p_1^*$  and  $p_2^*$  88 of the minmax problem (7) are given by

$$p_1^{\star} = \frac{\alpha\sqrt{2\kappa}}{4} \left[ \sqrt{8 + v^2} - v \right], \qquad p_2^{\star} = \frac{\alpha\sqrt{2\kappa}}{4} \left[ \sqrt{8 + v^2} + v \right], \qquad 90$$

where  $\alpha=(\omega_{\min}\omega_{\max})^{1/4}$ ,  $\beta=\alpha^{-1}(\sqrt{\omega_{\min}}+\sqrt{\omega_{\max}})$  and

$$v = \begin{cases} \frac{2\sqrt{\beta - 1}}{\sqrt{2\beta^2 - 12}} & \text{if } \beta \ge 1 + \sqrt{5}, \\ \sqrt{2\beta^2 - 12} & \text{if } \sqrt{6} \le \beta < 1 + \sqrt{5}, \\ 0 & \text{if } 2 < \beta < \sqrt{6}. \end{cases}$$

It is worth mentioning that even if the diffusion coefficients are continuous the 93 *two-sided* case provides a faster convergence than the *one-sided* case studied in [4] 94 (Fig. 1). 95

**General Remarks:** 96

• The usual methodology to optimize the convergence at the continuous level 97 comes with a few assumptions that may lead to inaccuracies once the prob- 98 lem is discretized. For example, as discussed in [7] (Sect. 5), the infinite domain 99 assumption used to determine the convergence factor (6) may lead to apprecia- 100 ble differences in the optimized parameters compared to an approach taking the 101 finiteness of the subdomains into account. We numerically found that the infinite domain assumption is valid as long as the dimensionless Fourier number 103 Fo =  $\kappa_i/(L_i^2\omega)$  (with  $L_i$  the size of subdomain  $\Omega_i$ ) of the problem does not 104 exceed a critical value  $Fo_c = 0.02$ .

• The optimization problem (7) aims at minimizing the maximum value of 106  $\rho(p_1, p_2, \omega)$  over the entire interval  $[\omega_{\min}, \omega_{\max}]$ . This provides a very robust 107 method general enough to deal with the worst case scenario when all the temporal frequencies are present in the error. An even more efficient way to proceed 109 would be to adjust the values of  $p_1$  and  $p_2$  at each iteration so that those parameters are efficiently chosen to "fight" the remaining frequencies in the error.

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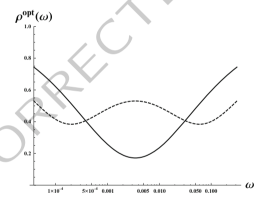


Fig. 1. Convergence factor optimized at the PDE level in the one-sided case (black line) [4] and in the two-sided case (dashed black line) [8], for  $\kappa = 10^{-2}$  m s<sup>-1</sup>,  $\Delta t = 10$  s, and  $T = 2^{13} \Delta t$ 

# 3 Optimal Control of the *Robin* Parameters

To investigate the robustness of the optimized parameters once the problem is discretized, the use of the optimal control theory appears as a natural choice. We aim at 114 controlling the Robin parameter in order to get the best possible convergence speed 115 in the sense of a given cost function  $\mathcal{J}$ . Moreover, following the approach of [3] 116 and the previous discussion, we consider the possibility to use different parameters 117  $p_i$  for different steps of the iterative process. It is easy to check that by choosing 118 different parameters at each iteration we still converge to the solution of the global 119 problem. A first way to choose the parameters is to look, at each iteration k, for  $p_1^k$ and  $p_2^k$  minimizing the error at the interface. In this case the cost function that we 121 intend to minimize at each iteration would be

$$\mathcal{J}(p_1^k, p_2^k) = \frac{w}{2} \int_0^T \left( u_1^k(0, t) - u_2^k(0, t) \right)^2 dt 
+ \frac{\widetilde{w}}{2} \int_0^T \left( \kappa_1 \partial_x u_1^k(0, t) - \kappa_2 \partial_x u_2^k(0, t) \right)^2 dt.$$
(8)

The constants w and  $\widetilde{w}$  must be chosen to balance both terms, depending on the characteristics of the problem (see Sect. 4). The cost function (8) is designed in agreement 124 with the consistency (5) we want to impose at the interface between subdomains. I provides a measure of the "inconsistency" of the solution at each iteration k, and is, 126 thus, directly related to the order of magnitude of the errors  $e_i^k$  of the algorithm (as 127) shown in Fig. 2). An other strategy could be to minimize the error at a given iteration 128 K. The cost function would thus be

$$\mathcal{J}\left((p_1^k, p_2^k)_{k=1,K}\right) = \frac{w}{2} \int_0^T \left(u_1^K(0,t) - u_2^K(0,t)\right)^2 dt 
+ \frac{\widetilde{w}}{2} \int_0^T \left(\kappa_1 \partial_x u_1^K(0,t) - \kappa_2 \partial_x u_2^K(0,t)\right)^2 dt,$$
(9)

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leading to an optimization on 2K parameters. This latter approach is particularly 130 interesting when we intend to obtain the best possible approximation of the exact 131 solution after a number of iterations set in advance. We propose here to lead our 132 study with this kind of approach with K = 5. The *optimal control* approach does not 133 per se reduce the computational cost of the algorithm because many evaluations of 134 the cost function are required during the minimization process (see Algorithm 3). We 135 use this approach as a tool to improve our understanding of the behavior of the Robin 136 parameters in order to find new directions to further accelerate the convergence speed 137 when Robin-type interface conditions are used. We denote by  $\mathbf{p}_1^{\star,\text{num}}$  and  $\mathbf{p}_2^{\star,\text{num}}$ the parameters found numerically by solving the optimal control problem. Those 139 parameters correspond to two vectors of size K. Similarly we will denote by  $p_1^{\star,ana}$ and  $p_2^{\star,ana}$  the parameters found analytically (cf. Theorem 1).

We used Matlab for the computation (Algorithm 3). Note that the well-posedness 142 of the coupling problem (4) is not sufficient to ensure a well-posed optimal control 143 problem. Some additional requirements on the convexity and regularity of the cost 144 function are necessary. We do not provide here such a proof, however we empirically 145 checked that the same solution of the optimal problem is obtained for a wide range 146 of parameter values for the initial guess.

# **4 Numerical Experiments**

We discretized problem (4) using a backward Euler scheme in time and a second 149 order scheme defined on a staggered grid in space (see [8] for more details). We 150

#### **Algorithm 3** Optimal control

```
%== Robin parameters found analytically: p1ana, p2ana
%== Solution of the optimal control problem: plopt, p2opt
%== Initial guess ==%
x_0(1:2:2*K-1)=p1ana;
x_0(2:2:2*K) = p2ana;
%== Solve the optimal control problem ==%
%== the CalcJ function proceeds to K iterations of the
%== Schwarz algorithm using 2K Robin parameters,
%== and computes the associated cost function (9)
x = \text{fminsearch}(\text{@CalcJ}, x_0);
%== Retrieve the optimized parameters
p1opt(1:K)=x(1:2:2*K-1);
p2opt(1:K)=x(2:2:2*K);
```

decompose the domain  $\Omega$  into two non-overlapping subdomains  $\Omega_1 = [-H, 0]$  and 151  $\Omega_2 = [0, H]$  with H = 500 m. The diffusion coefficient is  $\kappa = 10^{-2}$  m<sup>2</sup> s<sup>-1</sup> and the total simulation time is  $T=2^{13}\Delta t$  with  $\Delta t=10$  s. The parameter values lead to a 153 dimensionless Fourier number smaller than 0.02 so that the infinite domain assump- 154 tion is valid. We simulate directly the error equations, i.e.  $f_1 = f_2 = 0$  in (4) and 155  $u_0(x) = 0$ . We start the iteration with a random initial guess  $u_2^0(0,t)$   $(t \in [0,T])$  so 156 that it contains a wide range of the temporal frequencies that can be resolved by 157 the computational grid. This is done to allow a fair comparison as the parameters 158 optimized at the PDE level are optimized assuming that the full range  $[\omega_{\min}, \omega_{\max}]$  159 is present in the error. We first perform the Optimized non-overlapping Schwarz 160 Method (referred as to OSM case) using  $p_1^{\star,ana}$  and  $p_2^{\star,ana}$  and then using an optimal 161 control of the *Robin* parameters with K = 5 (referred as to OptCon case). We first 162 check that the minimization of cost function  $\mathcal{J}$  consistently implies the reduction of 163 the errors  $||e_i||_{\infty}$  of the associated algorithm (Fig. 2). For our experiments, we chose w=1 and  $\widetilde{w}=H/\kappa$  in (9). We notice that in the OptCon case the convergence speed 165 is significantly improved compared to the OSM case. Indeed, nine iterations of the 166 OSM are required to obtain the same accuracy than the OptCon case after only five 167 iterations. In order to have more insight on the way the parameters  $\mathbf{p}_1^{\star,\text{num}}$  and  $\mathbf{p}_2^{\star,\text{num}}$ evolve throughout the iterations we plot, in Fig. 3, the corresponding convergence 169 factor (6) at each iteration. It is striking to realize that the optimal convergence is 170 obtained through a combination of 2-point (equivalent to the *one-sided* case) and 3-171 point (equivalent to the two-sided case) equioscillations sometimes shifted along the 172 ω-axis to adapt to the temporal frequencies still present in the error. The first two 173 iterations aim at working mainly on the high-frequency components while the last 174 three iterations are optimized to work on the low-frequency component. The adap- 175 tivity of the Robin parameters from one iteration to the other brings more flexibility 176 to the method enabling more scale selectivity.

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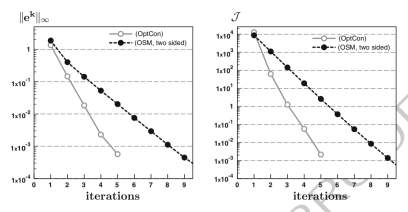


Fig. 2. Evolution of the  $\mathcal{L}^{\infty}$ -norm of the error (*left*) and of the cost function  $\mathcal{J}$  (*right*) with respect to the iterates k in the OSM and OptCon cases

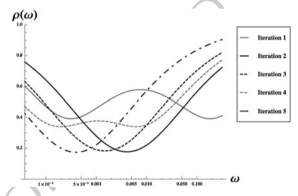


Fig. 3. Sequence of convergence factors  $\rho(\omega)$  resulting from the optimal control of the Robin parameters determined to get the best possible convergence after K = 5 iterations

5 Conclusion 178

Due to its simplicity, the use of *Robin-type transmission conditions* is very attractive 179 when one wants to couple unsteady problems defined on non-overlapping subdo- 180 mains. Once the Robin parameters are properly chosen one can achieve a fast con- 181 vergence [2]. In the present study we showed that there is still room for improvement 182 in the design of the Robin conditions. If the Robin parameters are adjusted from one 183 iteration to the other we showed, thanks to an optimal control approach, that we can 184 significantly improve the convergence speed. It is important to emphasize that the 185 optimal control paradigm proposed in this study is general enough to be used with 186 any type of PDE and an arbitrary number of subdomains.

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# A New Distributed Optimization Approach for Solving **CFD Design Problems Using Nash Game Coalition** and Evolutionary Algorithms

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#### 1 Introduction

For decades, domain decomposition methods (DDM) have provided a way of solv- 9 ing large-scale problems by distributing the calculation over a number of processing 10 units. In the case of shape optimization, this has been done for each new design 11 introduced by the optimization algorithm. This sequential process introduces a bottleneck.

Shape optimization is often done using gradient-based approaches because of 14 their superior efficiency. Adjoint methods provide a mathematical approach of com- 15 puting the gradients [4] using calculus of variations. Methods that combine the gov- 16 erning PDEs, their adjoints and shape parameters into one large system of equations 17 are called *one-shot methods* [1, 6]. The optimal shape can be acquired by solving the system of equations only once. Evidently, this approach has several drawbacks. If 19 the objective function is not unimodal, the method does not guarantee capturing the 20 global optimal solution. Also, if the geometry changes are large, mesh deformation 21 is no longer possible and the mesh has to be regenerated which makes this approach 22 costly.

In this paper, a "distributed one-shot" method is introduced. It is based on ideas 24 originating from the fields of game theory, domain decomposition, and evolutionary 25 computing. The aim is to speed up convergence on one hand by decreasing compu- 26 tational time by intelligent parallelism using Nash game strategies and on the other 27 hand by eliminating the bottleneck caused by sequential "state-costate - gradient" 28 chain processing. The evolutionary approach allows the method to be used in global 29 or non-smooth optimization.

#### 1.1 Nash Games in Geometry and Domain Decomposition

Competitive Nash games were introduced by J. Nash [5]. In a competitive game the 32 players maximize their payoff by taking into account the opponents' strategies. Nash 33 games converge into a Nash equilibrium. For simplicity, let us consider a two-player 34

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game. Let  $S_1$  and  $S_2$  be the sets of available strategies of Players 1 and 2 and  $J_1$  and 35  $J_2$  their payoff functions. A strategy pair  $(\bar{x}_1, \bar{x}_2) \in (S_1, S_2)$  is a Nash equilibrium if 36 and only if

$$J_{1}(\bar{x}_{1}, \bar{x}_{2}) = \inf_{\substack{x_{1} \in S_{1} \\ x_{2} \in S_{2}}} J_{1}(x_{1}, \bar{x}_{2})$$

$$J_{2}(\bar{x}_{1}, \bar{x}_{2}) = \inf_{\substack{x_{2} \in S_{2} \\ x_{2} \in S_{2}}} J_{2}(\bar{x}_{1}, x_{2})$$
(1)

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The above definition can be easily generalized to a Nash game with N players.

Nash games can also be applied to single-objective optimization. If the objective function J is additively separable, i.e.  $J(\mathbf{x}) = \sum_{i=1}^{N} J_i(\mathbf{x}_i)$  and  $\min_{\mathbf{x}} J(\mathbf{x}) = \sum_{i=1}^{N} J_i(\mathbf{x})$  $\min_{\mathbf{x}_i} \sum_{i=1}^N J_i(\mathbf{x}_i) = \mathbf{0}$ , a "virtual" Nash game can be formed [3]. Since there are no 41 true conflicts between the criteria, the global Nash equilibrium is located at the global 42 optimum.

The Nash approach is well suited for inverse problems. The geometry can often 44 be decomposed into smaller subgeometries which can be optimized concurrently 45 [11]. Similarly, a domain decomposition problem for solving a partial differential 46 equation can be considered as an inverse problem with a Nash game approach where 47 the objective function is to minimize the discrepancy between the local overlapped 48 subdomain solutions,

$$JF_{1}(g_{1},\bar{g}_{2}) = \int_{\Omega_{1,2}} |\varphi_{1}(g_{1},\bar{g}_{2}) - \varphi_{2}(g_{1},\bar{g}_{2})|^{2}$$

$$JF_{2}(\bar{g}_{1},g_{2}) = \int_{\Omega_{1,2}} |\varphi_{2}(\bar{g}_{1},g_{2}) - \varphi_{1}(\bar{g}_{1},g_{2})|^{2}$$
(2)

where  $|\cdot|$  is the  $L^2$  norm,  $\varphi_i$  is the solution in the subdomain  $\Omega_i$  and  $g_i$  is the vector of 50 values of  $\varphi_i$  on the subdomain interface boundary  $\Gamma_{i,j}$ .  $\Omega_{1,2}$  is the overlapping region 51 (cf. Fig. 1).

In [3, 7], a hierarchical leader-follower Stackelberg game consisting of a pair of 53 Nash games was implemented for nozzle shape reconstruction. The shape players 54 reconstructed the target geometry using a "leader" Nash game, and the flow play- 55 ers reconstructed the flow using a "follower" Nash game. For each new geometry 56 candidate produced by the shape players, a Nash game was run between the flow 57 players. In this paper, a new Nash evolutionary approach is introduced. It replaces 58 the computationally expensive hierarchical game by a single parallel global Nash 59 game coalition.

#### 1.2 Global Nash Game Coalition Algorithm (GNGCA)

The proposed method operates as follows. The geometry of the configuration is di- 62 vided into subgeometries allocated to shape players whose task is to optimize the 63 shape (or reconstruct the target geometry). Similarly, the flow players minimize the 64 deviation of local solutions on the overlapped region of subdomains. Each shape and 65 flow player evaluate deviation of local solutions or shape optimization with his own 66 Evolutionary Algorithm (EA). After some frequency period, for example a single 67 generation, shape and flow players exchange the elite values among each other. This 68 means the flow is reconstructed along with the geometry making this a "distributed 69 one-shot" method.

This new method is inherently parallel and therefore especially suitable for distributed parallel environments. At the higher level, the flow and shape players operate separately. Depending on the methods used, the optimization process can also be distributed. If an optimizer is used in flow reconstruction, it too can be parallelized. By reducing dimensionality of the geometry problem, algorithmic convergence can be significantly improved. For example, in the case of multi-modal problems splitting the territory can reduce the number of local optima. However, the efficiency of virtual Nash approach is highly dependent on the selected geometry decomposition. Non-optimal splitting can lead in reduced efficiency of the algorithm [11].

### 2 Test Case Description

The method is validated using a simple position reconstruction problem from the 81 field of computational fluid dynamics. The geometry of the problem consists of a 82 large disk element (radius  $\frac{1}{2}$  units) surrounded by  $N \ge 2$  smaller disk elements (radii 83  $\frac{1}{8}$  units). The smaller elements are allowed to move in an area constrained by the 84 number of elements: using radial coordinates,  $r_k = 2.0^{+0.5}_{-1.3675}$  and  $\theta_k = -k\frac{2\pi}{N} - \frac{\pi}{N} \pm$  85  $\frac{\pi}{M}$  (see Fig. 1).

This geometry allows the study of a wide variety of different domain and geometry decompositions (cf. Fig. 1 for a 3 element case). The test case can be made more challenging for example by deforming the shapes of the elements. In this paper, 2 and 6 element cases were studied.

The flow is described by the steady compressible potential flow,

$$\nabla \cdot \rho \nabla \varphi_{k} = 0 \quad \text{in } \Omega_{k}$$

$$\varphi_{k} = \mathbf{v}_{\infty} \text{ on } \Gamma_{\infty}$$

$$\frac{\partial \varphi_{k}}{\partial \mathbf{n}} = 0 \quad \text{on } \Gamma_{1,\dots,n}$$

$$\varphi_{k} = \varphi_{j} \text{ on } \Gamma_{j}$$

$$\varphi_{k} = \varphi_{\ell} \text{ on } \Gamma_{\ell}$$
(3)

where k is the index of the subdomain, and  $j,\ell$  the right and left side neighbor 92 domain indexes. Free-flow velocity  $\mathbf{v}_{\infty} = (v_x, v_y) = (v_{\infty} \cos \alpha, v_{\infty} \sin \alpha), \ |v_{\infty}| = 1$ . 93 The angle of attack  $\alpha = 0.0^{\circ}$ . The density  $\rho$  is calculated using the formula  $\rho = 94$   $\left\{1 + \frac{\gamma - 1}{2}M_{\infty}^2\left(1 - |v|^2\right)\right\}^{\frac{1}{\gamma - 1}}$ . The constant  $\gamma = 1.4$  is the ratio of specific heats for 95 air. With a free flow Mach number  $M_{\infty} = .3$  the flow is subsonic in the whole domain. 96 The objective is to reconstruct the original positions of the elements by minimizing the  $L^2$  norm of pressure difference between the computed and target surface 98 pressures:  $JS_k(\mathbf{x}_k) = \frac{1}{n_{p_k}} \sum_{i=1}^{n_{p_k}} \left| p_{k_i} - p_{k_i}^{target} \right|^2$  where  $\mathbf{x}_k = (r_k, \theta_k)$  is the decomposed 99 design vector and  $n_{p_k}$  is the number of pressure points in the region of the decomposed geometry. The vector  $p_k$  includes the relevant surface pressure values. The 101 global objective function is the sum of local functions. The objective function for the 102 flow players is the  $L^2$  norm of the discrepancy on the overlapped subregion (Eq. 2).

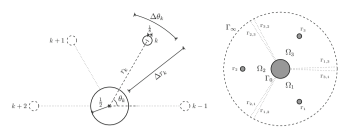


Fig. 1. Test case geometry and example decomposition

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### 3 Test Setting

A variant of the popular Differential Evolution (DE) algorithm is used as the opti- 105 mization platform. The algorithm, differential evolution with adaptive control param- 106 eters (jDE) is described in detail in the original paper [2]. The difference compared to the standard differential evolution is that the two control parameters, mutation factor 108 F and crossover rate CR are not kept fixed. Instead, each member of the population 109 has individual values which are allowed to change between given ranges. When a 110 new individual is formed, the offspring inherits the values from its progenitor, or 111 new random values are generated with probability of  $\tau_1$  for F and  $\tau_2$  for CR. In this 112 work the population size  $NP = 10n_{dim}$  was used where  $n_{dim}$  is the number of dimensions in the decomposed design vector, i.e. each instance of algorithm uses an equal 114 number of individuals in order to make comparing them fair. Mutation factor is al- 115 lowed to vary within the range F = [0.1, 1.0] and crossover rate CR = [0.0, 1.0]. The control parameter replacement probabilities are set to  $\tau_{1,2} = 0.1$ . The algorithms end when the stopping criteria  $JS_k = 10^{-5}$  is reached.

Because the algorithms work in parallel, a generational approach would cause 119 bottlenecks because of the non-constant fitness function computation times. Instead, 120 a non-generational approach is used where the older individuals are replaced imme- 121 diately if the offspring is superior. In addition, the elite information exchange is done 122 asynchronously.

Three different approaches are tested. In the first one, the jDE algorithm is run 124 traditionally using full domain and design vector. For the second approach, a "geom- 125 etry decomposition" approach introduced in [9] is used ("Nash-jDE"). The design 126 vector  $\mathbf{x} = (r_1, \theta_1, \dots, r_N, \theta_N)$  is divided between the elements  $(\mathbf{x}_k = (r_k, \theta_k), k =$  $\{1,\ldots,N\}$ ), which are then optimized using several jDE algorithms operating on 128 separate subpopulations. After each generation, the global design vector is updated 129 using elite values from each subpopulation. The proposed GNGCA algorithm is used 130 in the third case. For flow reconstruction, since the flow is subsonic, the additive 131 Schwarz domain decomposition algorithm is sufficient. The overlapped regions of 132 subdomains are made of one strip. The computational domain is divided radially so 133 that each subdomain contains one element (Fig. 1).

The FreeFEM++ v3.18 software is used as the solver [8]. The flow is computed 135 using finite element method with a preconditioned conjugate gradient algorithm. 136

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**Table 1.** Performance of the algorithms. The symbol  $n_{sl}$  refers to the number of (shape player) slave processes, t is the wall-clock time in seconds and  $n_{it}$  to the number of objective function evaluations required by the algorithm in order to reach the target precision.

		jDE		Nash-	jDЕ	GNG	CA	spee	d-up
case	$n_{sl}$	t	$n_{it}$	t	$n_{it}$	t	$n_{it}$	jDE	N-jDE
	2	1155.00s	815	390.83s	279	306.57s	514	3.77×	1.27×
2 elements	4	332.05s	474	210.97s	302	194.74s	652	$1.70 \times$	1.08×
	6	190.42s	412	132.62s	279	174.60s	888	1.09×	0.76×
	6	3632.85s	4387	971.17s	1175	171.61s	1894	21.17×	5.66×
6 elements	12	1742.23s	4226	333.90s	809	115.87s	2502	$15.04 \times$	$2.88 \times$
	18	1201.11s	4369	244.53s	880	114.08s	3743	$10.53 \times$	$2.14 \times$

Since the flow is nonlinear, Eq. 3 is solved iteratively until the threshold value of 137  $\varepsilon_0 = 10^{-10}$  for density is reached. The algorithms are run on a computer containing 64 Intel Xeon CPU cores clocked at 2.67 GHz.

The mesh is constructed using Triangle v1.6 Delaunay mesh generator [10]. Nu- 140 merical noise is minimized using mesh regeneration with the Laplacian. In order to 141 avoid inverse elements and maintain mesh quality, the mesh is regenerated over certain intervals ( $\delta r_k = 0.1, \delta \theta_k = 10^\circ$ ). An example decomposed mesh is illustrated in 143 Fig. 3. Computing one subdomain gives speed-ups ranging from  $3.2 \times$  to  $14.0 \times$ .

### 4 Results and Discussion

The elapsed wall-clock times and the number of objective function evaluations re- 146 quired by each of the algorithm are listed on Table 1. Convergence curves of the 147 algorithms are shown in Fig. 2. Final mesh and reconstructed global pressure field 148 are compared to the reference in Fig. 3.

The results demonstrate that the geometry decomposition method using virtual 150 Nash games can be used to increase algorithmic efficiency in geometry reconstruction problems. The proposed global Nash game approach shows that reconstructing 152 geometry and flow simultaneously the wall-clock time can be reduced dramatically, 153 provided the difference in the size of global and decomposed domains is sufficiently 154 large. In the case of six domains, the speed-up compared to the original method is 155 massive, over 20×. The increase compared to the pure geometry decomposition ap- 156 proach is also notable, over  $5 \times$ . If the algorithms are compared a bit more fairly, i.e. 157 the flow players are considered equal to the shape players, the speed-ups are  $10 \times 158$ and  $2\times$ .

The efficiency of flow reconstruction is critical for the success of the proposed algorithm. Finding the correct geometry in an incompletely reconstructed flow field is 161 not possible, which is evident in the large number of shape player objective function 162 iterations needed. Unlike in the case of the other methods, increasing the number of 163 slave processes brought only limited speed-ups for GNGCA. This was due the fact 164

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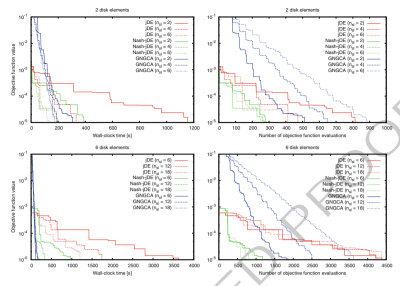


Fig. 2. Convergence curves of the tested algorithms. The onvergence according to the wallclock time spent is on the left and the algorithmic convergence based on the required number of iterations is on the right

the flow players did not feed the shape players with accurate flow information fast 165 enough resulting in an increased number of shape player iterations and correspond- 166 ingly reduced efficiency improvement.

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Algorithmic convergence can be improved by reducing the complexity of the 168 problem. A classical method where the boundary nodes are used as shape design 169 variables may be problematic due to a large number of variables. The situation can 170 be improved using parallel algorithms and Bézier spline parametrization. In cases 171 involving highly compressible potential flows where the flow is locally supersonic 172 the domain reconstruction has to be augmented with an optimizer. The flow can be 173 reconstructed using fast gradient methods on linearized equations coupled by DDM, 174 or analogously to the shape presentation, the number of variables on interface bound- 175 ary can be reduced using parametrization and the nonlinear flow can be reconstructed 176 with evolutionary algorithms (cf. [3]).

### **5** Conclusion and Future

In this paper first results for a new "distributed one-shot" method that applies vir- 179 tual Nash games, domain and geometry decomposition methods, are presented and 180 discussed. The feasibility of the method is validated using an academic test case 181 consisting of position reconstruction in a subsonic nonlinear flow.

In the forthcoming step, the Schwarz domain decomposition algorithm will be 183 replaced with more robust methods. The simple compressible potential flow equa- 184

#### New Distributed Optimization Approach

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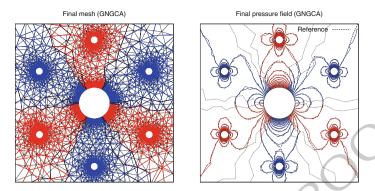


Fig. 3. Example final mesh and pressure field (GNGCA) compared to the reference

tion will be replaced with nonlinear systems of equations including Euler, Navier- 185 Stokes, and Maxwell equations. Further tests involve complex geometries such as 186 multi-element airfoils. The implementation of GPUs is also being studied. The ulti- 187 mate target is to extend the method to speed up the capture of solutions of complex 188 large scale problems which are frequently met in particular in 3D industrial detailed 189 design.

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# A Neumann-Dirichlet Preconditioner for FETI-DP Method for Mortar Discretization of a Fourth Order **Problems in 2D**

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1 Introduction a

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This study focuses on a construction of a parallel preconditioner for a FETI-DP 9 (dual primal Finite Element Tearing and Interconnecting) method for a mortar Hsieh- 10 Clough-Tocher (HCT) discretization of a model fourth order problem with discontinuous coefficients.

FETI-DP methods were introduced in [8]. They form a class of fast and efficient 13 iterative solvers for algebraic systems of equations arising from the finite element 14 discretizations of elliptic partial differential equations of second and fourth order, 15 cf. [8, 10, 11, 16] and references therein. In a one-level FETI-DP method one has 16 to solve a linear system for a set of dual variables formulated by eliminating all 17 primal unknowns. The FETI-DP system contains in itself a coarse problem, while 18 the preconditioner is usually fully parallel and constructed only from local problems. 19

There are many works investigating iterative solvers for mortar method for sec- 20 ond order problem, e.g. cf. [1-3] and references therein. There have also been a few 21 FETI-DP type algorithms developed for mortar discretization of second order prob- 22 lems, cf. e.g. [6, 7, 9]. But there is only a small number of studies focused on fast 23 solvers for mortar discretizations of fourth order elliptic problems, cf. [12, 15, 17]. 24 In this study we follow the approach of [9] which considers the case of a FETI-DP 25 method for mortar discretization of a second order problem.

In this paper we first present the construction of mortar discretization of a fourth 27 order elliptic problem which locally utilizes Hsieh-Clough-Tocher finite elements 28 in the subdomains. Next we introduce a FETI-DP problem and then a Neumann- 29 Dirichlet parallel preconditioner for a FETI-DP problem is proposed. Finally, we 30 present the almost optimal bounds of the condition number, namely, a bound which 31 grows like  $C(1 + \log(H/h))^2$ , where H is the maximal diameter of subdomains and 32 h is a fine mesh parameter.

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### 2 Discrete Problem

In this section we focus on a mortar Hsieh-Clough-Tocher (HCT) finite element discretization for a model fourth order elliptic problem with discontinuous coefficients. 36

Let  $\Omega$  be a polygonal domain in the plane. We assume that there exists a partition of  $\Omega$  into disjoint polygonal subdomains  $\Omega_k$  such that  $\overline{\Omega} = \bigcup_{k=1}^N \overline{\Omega}_k$  with  $\overline{\Omega}_k \cap \overline{\Omega}_l$  38 being an empty set, an edge or a vertex (crosspoint). We also assume that these 39 subdomains form a coarse triangulation of the domain which is shape regular in 40 the sense of [5]. We introduce a global interface  $\Gamma = \bigcup_i \overline{\partial \Omega_i} \setminus \overline{\partial \Omega}$  which plays an 41 important role in our study.

Our model differential problem is to find  $u^* \in H_0^2(\Omega)$  such that

$$a(u^*, v) = \int_{\Omega} f v \, dx \quad \forall v \in H_0^2(\Omega), \tag{1}$$

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where  $f \in L^2(\Omega)$ ,  $H_0^2(\Omega) = \{u \in H^2(\Omega) : u = \partial_n u = 0 \text{ on } \partial\Omega\}$  and a(u,v) = 44  $\sum_{k=1}^N \int_{\Omega_k} \rho_k [u_{x_1x_1}v_{x_1x_1} + 2u_{x_1x_2}v_{x_1x_2} + u_{x_2x_2}v_{x_2x_2}] dx$ . The coefficients  $\rho_k$  are positive 45 and constant. Here  $u_{x_kx_l} := \frac{\partial^2 u}{\partial x_k\partial x_l}$  for k,l=1,2 and  $\partial_n u$  is a unit normal derivative of u.

In each subdomain  $\Omega_k$  we introduce a quasiuniform triangulation  $T_h(\Omega_k)$  made 48 of triangles with the parameter  $h_k = \max_{\tau \in T_h(\Omega_k)} \operatorname{diam}(\tau)$ , cf. e.g. [4]. We can now

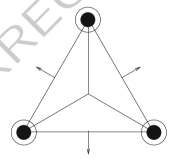


Fig. 1. Degrees of freedom of HCT element

introduce local finite element spaces. Let  $X_h(\Omega_k)$  be the Hsieh-Clough-Tocher (HCT) 50 macro finite element space defined as follows:

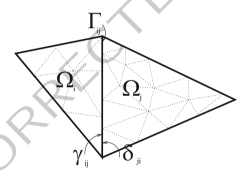
$$X_h(\Omega_k) = \{u \in C^1(\Omega_k) : u \in P_3(\tau_i), \ \tau_i \in T_h(\Omega_k), \text{ for the subtriangles } \tau_i, \ i = 1, 2, 3, \text{ formed by connecting the vertices of any } \tau \in T_h(\Omega_k) \text{ to its centroid, and} \ u = \partial_n u = 0 \text{ on } \partial\Omega_k \cap \partial\Omega\},$$

where  $P_3(\tau_i)$  is the function space of cubic polynomials defined over  $\tau_i$ . The degrees 52 of freedom of a function  $u \in X_h(\Omega_k)$  over  $\tau \in T_h(\Omega_k)$  are defined as:  $\{u(p_k), \nabla u(p_k), 53 \partial_n u(m_j)\}_{k,j=1,2,3}$ , where  $p_k$  is a vertex and  $m_j$  is a midpoint of an edge of  $\tau$ , cf. Fig. 1. 54

Next a global space  $X_h(\Omega)$  is defined as  $X_h(\Omega) = \prod_{i=1}^N X_h(\Omega_k)$ . We also introduce  $\widetilde{X}_h(\Omega)$  – a subspace of  $X_h(\Omega)$  formed by all functions in  $X_h(\Omega)$ , which has 56 all degrees of freedom continuous at the crosspoints, i.e. the common vertices of 57 substructures.

Let  $\Gamma_{kl}$  denote the interface between two subdomains  $\Omega_k$  and  $\Omega_l$  i.e. the open edge 59 that is common to these subdomains. Note that each interface  $\Gamma_{kl}$  inherits two one 60 dimensional triangulations made of segments that are edges of elements of  $T_h(\Omega_k)$  61 and  $T_h(\Omega_l)$ , respectively. Thus there are two independent 1D triangulations on  $\Gamma_{kl}$ : 62  $T_{h,k}(\Gamma_{kl})$  related to  $\Omega_k$  and another one associated with  $\Omega_l$  -  $T_{h,l}(\Gamma_{lk})$ , cf. Fig. 2. Let 63  $\gamma_{kl}$  be a mortar, i.e. the side corresponding to  $\Omega_k$  if  $\rho_k \geq \rho_l$  and then let  $\delta_{lk}$  be the other side of  $\Gamma_{lk}$  associated to  $\Omega_l$  called a slave (nonmortar).

For each interface  $\Gamma_{kl}$  we introduce two test spaces associated with its slave triangulation  $T_{h,l}(\delta_{lk})$  (cf. [13, 14]): let  $M_t^h(\delta_{lk})$  be the space formed by  $C^1$  smooth 67 piecewise cubic functions on the slave triangulation of  $\delta_{lk}$ , which are piecewise linear in the two end elements, and let  $M_n^h(\delta_{lk})$  be the space of continuous piecewise 69 quadratic functions on the elements of this triangulation, which are piecewise linear 70 in the two end elements.



**Fig. 2.** Independent meshes on an interface  $\Gamma_{ij}$ 

We also define a space  $M = \prod_{\delta_{lk} \subset \Gamma} M_{lk}$  with  $M_{lk} = M_t^l(\delta_{lk}) \times M_n^l(\delta_{lk})$  and a 72 bilinear form  $b(u, \psi)$ : let  $u = (u_k)_{k=1}^N \in \widetilde{X}_h(\Omega)$  and  $\psi = (\psi_{lk})_{\delta_{lk}} = (\psi_{lk,t}, \psi_{lk,n})_{\delta_{lk}} \in \tau_3$ M, then  $b(u, \psi) = \sum_{\delta_{lk} \subset \Gamma} \sum_{s \in \{t,n\}} b_{lk,s}(u, \psi_{lk,s})$  with

$$b_{lk,t}(u, \psi_{lk,t}) = \int_{\delta_{lk}} (u_k - u_l) \psi_{lk,t} ds,$$
  
$$b_{lk,n}(u, \psi_{lk,n}) = \int_{\delta_{lk}} (\partial_n u_k - \partial_n u_l) \psi_{lk,n} ds.$$

Further we will use the same notation for a function and for the vector with the values 75 of degrees of freedom of this function.

We introduce discrete problem as the saddle point problem: find a pair  $(u_h^*, \lambda^*) \in$ 77  $\widetilde{X}_h(\Omega) \times M$  such that 78

$$a(u_h^*, v) + b(v, \lambda^*) = f(v) \quad \forall v \in \widetilde{X}_h(\Omega),$$
 (2)

$$b(u_h^*, \phi) = 0 \qquad \forall \phi \in M, \tag{3}$$

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where  $a_h(u, v) = \sum_{k=1}^{N} a_k(u, v)$  for

$$a_k(u,v) = \int_{\Omega_k} \rho_k[u_{x_1x_1}v_{x_1x_1} + 2u_{x_1x_2}v_{x_1x_2} + u_{x_2x_2}v_{x_2x_2}] dx.$$

This problem has a unique solution and error bounds are established, e.g. cf. [14].

### 3 Matrix Form of Mortar Conditions

Note that (3) is equivalent to two mortar conditions on each  $\delta_{lk} = \gamma_{kl} = \Gamma_{kl}$ :

$$\int_{\delta_{lk}} (u_k - u_l) \phi \, ds = 0 \quad \forall \phi \in M_t^l(\delta_{lk}), \tag{4}$$

$$\int_{\delta_{lk}} (\partial_n u_k - \partial_n u_l) \psi \, ds = 0 \quad \forall \psi \in M_n^l(\delta_{lk}). \tag{5}$$

We introduce the following splitting of two vectors representing the tangential 84 and normal traces  $u_{\delta_{lk}}$  and  $\partial_n u_{\delta_{lk}}$ :  $u_{\delta_{lk}} = u_{\delta_{lk}}^{(r)} + u_{\delta_{lk}}^{(c)}$  and  $\partial_n u_{\delta_{lk}} = \partial_n u_{\delta_{lk}}^{(r)} + \partial_n u_{\delta_{lk}}^{(c)}$  on a 85 slave  $\delta_{lk} \subset \partial \Omega_l$ , where superscript (c) refers to degrees of freedom related to crosspoints (ends of this edge) and superscript (r) refers to degrees of freedom related to 87 remaining nodes (vertices and midpoints) on this edge. We can now rewrite (4) and 88 (5) in a matrix form on each interface  $\Gamma_{kl} \subset \Gamma$ :

$$B_{t,\delta_{lk}}^{(c)} u_{\delta_{lk}}^{(c)} + B_{t,\delta_{lk}}^{(r)} u_{\delta_{lk}}^{(r)} = B_{t,\delta_{lk}}^{(c)} u_{\delta_{lk}}^{(c)} + B_{t,\gamma_{kl}}^{(r)} u_{\gamma_{kl}}^{(c)} + B_{t,\gamma_{kl}}^{(r)} u_{\gamma_{kl}}^{(r)}, B_{n,\delta_{lk}}^{(c)} \partial_{n} u_{\delta_{lk}}^{(c)} + B_{n,\delta_{lk}}^{(r)} \partial_{n} u_{\delta_{lk}}^{(r)} = B_{n,\gamma_{kl}}^{(c)} \partial_{n} u_{\gamma_{kl}}^{(c)} + B_{n,\gamma_{kl}}^{(r)} \partial_{n} u_{\gamma_{kl}}^{(r)},$$
(6)

where the matrices  $B_{t,\delta_{lk}} = [B_{t,\delta_{lk}}^{(c)}, B_{t,\delta_{lk}}^{(r)}]$  and  $B_{n,\delta_{lk}} = [B_{n,\delta_{lk}}^{(c)}, B_{n,\delta_{lk}}^{(r)}]$  are mass matri-90 ces obtained by substituting the traces of standard nodal basis functions of  $X_h(\Omega_l)$  91 and nodal basis functions of  $M_t^l(\delta_{lk}), M_n^l(\delta_{lk})$ , respectively, into (4). The matrices 92  $B_{t,\gamma_{kl}} = [B_{t,\gamma_{kl}}^{(c)}, B_{t,\gamma_{kl}}^{(r)}]$  and  $B_{n,\gamma_{kl}} = [B_{n,\gamma_{kl}}^{(c)}, B_{n,\gamma_{kl}}^{(r)}]$  are constructed analogously but utilizing traces onto  $\gamma_{kl}$  of standard nodal basis functions of  $X_h(\Omega_k)$ . Note that  $B_{t,\delta_{lk}}^{(r)}, B_{n,\delta_{lk}}^{(r)}$  94 are positive definite square matrices, but that all other matrices in (6) are rectangular 95 in general.

### **4 FETI-DP Problem**

Let  $K_l$  be a matrix of  $a_l(\cdot,\cdot)$  in the standard basis of  $X_h(\Omega_l)$ . Then let  $\tilde{K}$  be the matrix 98 obtained from a block diagonal matrix  $K := \operatorname{diag}(K_l)_{l=1}^N$  by taking into account the 99 continuity of the degrees of freedom at crosspoints. We can partition  $\tilde{K}$  into

$$\tilde{K} = \begin{pmatrix} K_{ii} & K_{ic} & K_{ir} \\ K_{ci} & K_{cc} & K_{cr} \\ K_{ri} & K_{rc} & K_{rr} \end{pmatrix},$$

where the superscript (i) refer to the degrees of freedom associated with nodal points interior to subdomain, (c) to the degrees of freedom related to crosspoints, and (r) to the degrees of freedom associated the remaining nodes on masters and slaves. Then the matrix formulation of (2) and (3) is the following:

$$\begin{pmatrix} K_{ii} & K_{ic} & K_{ir} & 0 \\ K_{ci} & K_{cc} & K_{cr} & (B^{(c)})^T \\ K_{ri} & K_{rc} & K_{rr} & (B^{(r)})^T \\ 0 & B^{(c)} & B^{(r)} & 0 \end{pmatrix} \begin{pmatrix} u^{(i)} \\ u^{(c)} \\ u^{(r)} \\ \lambda^* \end{pmatrix} = \begin{pmatrix} f_i \\ f_c \\ f_r \\ 0 \end{pmatrix}.$$
(7)

Here  $B^{(c)}$  is the matrix built from  $B^{(c)}_{t,\delta_{lk}}, B^{(c)}_{n,\delta_{lk}}, B^{(c)}_{t,\gamma_{kl}}, B^{(c)}_{n,\gamma_{kl}}$  for all  $\Gamma_{kl} = \gamma_{kl} = \delta_{lk} \subset \Gamma$  and  $B^{(r)} := \mathrm{diag}([-B^{(r)}_{\gamma_{kl}}, B^{(r)}_{\delta_{lk}}])_{\Gamma_{kl} \subset \Gamma}$  is the block diagonal matrix with

$$B_{\gamma_{kl}}^{(r)} := \begin{pmatrix} B_{t,\gamma_{kl}}^{(r)} & 0\\ 0 & B_{n,\gamma_{kl}}^{(r)} \end{pmatrix}, \qquad B_{\delta_{lk}}^{(r)} := \begin{pmatrix} B_{t,\delta_{lk}}^{(r)} & 0\\ 0 & B_{n,\delta_{lk}}^{(r)} \end{pmatrix}. \tag{8}$$

Next we eliminate the unknowns related to the interior nodes and crosspoints i.e. 108  $u^{(i)}$ ,  $u^{(c)}$  in (7) and we get 109

$$\tilde{S}u^{(r)} + \tilde{B}^T \lambda^* = \tilde{f}_r, 
\tilde{B}u^{(r)} + \tilde{S}_{cc} \lambda^* = \tilde{f}_c,$$
(9)

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119

where the respective matrices are defined as follows:

$$\tilde{S} := K_{rr} - (K_{ri} K_{rc}) (\tilde{K}^{(ic)})^{-1} \begin{pmatrix} K_{ir} \\ K_{cr} \end{pmatrix},$$

$$\tilde{B} := B^{(r)} - (0 B^{(c)}) (\tilde{K}^{(ic)})^{-1} \begin{pmatrix} K_{ir} \\ K_{cr} \end{pmatrix},$$
111
113

$$ilde{B}:=B^{(r)}-(0\,B^{(c)})( ilde{K}^{(ic)})^{-1}\left(rac{K_{ir}}{K_{cr}}
ight),$$
 113

and 
$$\tilde{S}_{cc} := -(0 \ B^{(c)})(\tilde{K}^{(ic)})^{-1} \begin{pmatrix} 0 \\ (B^{(c)})^T \end{pmatrix}$$
 with the nonsingular matrix  $\tilde{K}^{(ic)} := 114 \begin{pmatrix} K_{ii} \ K_{ic} \ K_{ci} \ K_{cc} \end{pmatrix}$ .

Eliminating  $u^{(r)}$  we obtain the following FETI-DP problem: find  $\lambda^* \in M$  such that 117

$$F(\lambda^*) = d,\tag{10}$$

where  $d := \tilde{f}_c - \tilde{B}\tilde{S}^{-1}\tilde{f}_r$  and  $F := \tilde{S}_{cc} - \tilde{B}\tilde{S}^{-1}\tilde{B}^T$ 118

### 5 Parallel Preconditioner

Let  $W_r=\{w^{(r)}:w\in\widetilde{X}_h(\Omega)\}$  i.e.  $W_r$  is the space of vectors representing all degrees 120 of freedom of functions from  $X_h(\Omega)$  associated with nodes (vertices and midpoints) 121 on  $\Gamma$  but are *not* associated with crosspoints. 122 We can decompose any vector  $w^{(r)} \in W_r$  into vectors related to masters and 123 slaves:

 $w^{(r)} = \left(w_{\Gamma}^{(r)}, w_{\Delta}^{(r)}\right)^T, \tag{125}$ 

where  $w_{\Gamma}^{(r)}$  is the vector with the values of degrees of freedom which are associated with the nodes on the masters and  $w_{\Delta}^{(r)}$  is the vector with the values of degrees of 127 freedom which are related to the nodes on the slaves. We then introduce  $W_{\Delta} = \{w_{\Delta}^{(r)}: 128\}$   $w_{\Delta}^{(r)} \in W_{r}$  i.e. the space formed by vectors in  $W_{r}$  which have only entries related to 129 the degrees of freedom which are associated with the nodes on the slaves. It is very 130 important to note that

$$\dim M = \dim W_{\Lambda}.$$

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Let  $S_{\Delta}$  be the matrix obtained by restricting  $\tilde{S}: W_r \to W_r$  to  $W_{\Delta}$ .

Note that this matrix is can be represented as a block diagonal matrix with non-singular diagonal blocks  $S_{k,\Delta}$ , i.e.

$$S_{\Delta} := \operatorname{diag}(S_{k,\Delta})_k,$$
 136

where the subscript k runs over all subdomains that have at least one edge on  $\Gamma$  as a slave. Naturally, we could also partitioned this matrix with respect to the slaves.

Define nonsingular block diagonal matrix  $B_{\Delta}: W_{\Delta} \to W_{\Delta}$ :

$$B_{\Delta} := \mathrm{diag}(B_{\delta_{lk}}^{(r)})_{\delta_{lk} \subset \Gamma},$$
 140

where  $B_{\delta_{lk}}^{(r)}$  are block diagonal matrices (with two nonsingular blocks) defined in (8). 141 Then we introduce our parallel preconditioner:

$$\mathscr{M}_{DN}^{-1} := B_{\Lambda}^{-T} S_{\Delta} B_{\Lambda}^{-1},$$

which is nonsingular, or equivalently its inverse:  $\mathcal{M}_{DN} := B_{\Delta} S_{\Delta}^{-1} B_{\Delta}^{T}$ . Note that  $S_{\Delta}$  and thus  $\mathcal{M}_{DN}$  are dependent on the discontinuous coefficients  $\rho_{k}$ .

#### 6 Condition Number Bounds

The main result of this paper is the following theorem which yields the bound of the condition number of preconditioned problem: 147

**Theorem 1.** It holds that

$$\langle \mathscr{M}_{DN}\lambda,\lambda \rangle \leq \langle F\lambda,\lambda \rangle \leq C \left(1+\log\left(rac{H}{\underline{h}}
ight)
ight)^2 \langle \mathscr{M}_{DN}\lambda,\lambda 
angle \qquad orall \lambda \in M,$$
 149

where  $H = \max_k h_k$ ,  $\underline{h} = \min_k h_k$ , and C a positive constant independent of the coefficients, or the parameters  $H_k$  and  $h_k$ . Here  $\langle \cdot, \cdot \rangle$  is the standard  $l_2$  inner product.

	as a direct consequence of this theorem we see that the condition number of	152
$\mathcal{M}_{\mathrm{DN}}^{-1}$	F is bounded by $C\left(1+\log\left(\frac{H}{\underline{h}}\right)\right)^2$ .	153
T	The lower bound in the theorem is obtained by purely algebraic arguments. And	154
	et the upper bound by using several technical results of which the most important	
_	s the estimate of special trace norms of jumps of tangential and normal traces	
over	an interface $\Gamma_{kl} \subset \Gamma$ .	15
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## A DG Space-Time Domain Decomposition Method

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Summary. In this paper we present a hybrid domain decomposition approach for the parallel 6 solution of linear systems arising from a discontinuous Galerkin (DG) finite element approximation of initial boundary value problems. This approach allows a general decomposition of 8 the space-time cylinder into finite elements, and is therefore applicable for adaptive refinements in space and time. 10

### 1 A Space-Time DG Finite Element Method

As a model problem we consider the transient heat equation

$$\partial_t u(x,t) - \Delta u(x,t) = f(x,t) \quad \text{for } (x,t) \in Q := \Omega \times (0,T),$$

$$u(x,t) = 0 \quad \text{for } (x,t) \in \Sigma := \partial \Omega \times (0,T),$$

$$u(x,0) = u_0(x) \quad \text{for } (x,t) \in \Omega \times \{0\}$$

$$(3)$$

$$u(x,t) = 0$$
 for  $(x,t) \in \Sigma := \partial \Omega \times (0,T)$ , (2)

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$$u(x,0) = u_0(x) \quad \text{for } (x,t) \in \Omega \times \{0\}$$
 (3)

where  $\Omega \subset \mathbb{R}^n$ , n = 1, 2, 3, is a bounded Lipschitz domain, and T > 0. Let  $\mathscr{T}_N$  be 13 a decomposition of the space-time cylinder  $Q = \Omega \times (0,T) \subset \mathbb{R}^{n+1}$  into simplices 14  $\tau_k$  of mesh size h. For simplicity we assume that the space time cylinder Q has a 15 polygonal (n = 1), a polyhedral (n = 2), or a polychoral (n = 3) boundary  $\partial Q$ . With  $\mathcal{I}_N$  we denote the set of all interfaces (interior facets) e between two neighboring 17 elements  $\tau_k$  and  $\tau_\ell$ . For an admissible decomposition the interior facets are edges 18 (n = 1), triangles (n = 2), or tetrahedrons (n = 3). 19

With respect to an interior facet  $e \in \mathcal{I}_N$  we define for a function v the jump

$$[v]_e(x,t) := v_{|\tau_k}(x,t) - v_{|\tau_\ell}(x,t)$$
 for all  $(x,t) \in e$ ,

the average 22

$$\langle v \rangle_e(x,t) := \frac{1}{2} \left[ v_{|\tau_k}(x,t) + v_{|\tau_\ell}(x,t) \right] \quad \text{for all } (x,t) \in e,$$

and the upwind in time direction by

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$$\{v\}_e^{\mathrm{up}}(x,t) := \begin{cases} v_{|\tau_k}(x,t) & \text{for } n_t \ge 0, \\ v_{|\tau_\ell}(x,t) & \text{for } n_t < 0 \end{cases} \quad \text{for all } (x,t) \in e,$$

where  $\mathbf{n} = (\mathbf{n}_x, n_t)$  is the normal vector of the interior facet e.

For a decomposition  $\mathcal{T}_N$  of the space–time cylinder Q we introduce the discrete 27 function space of piecewise polynomials of order p 28

$$S_{h,0}^p(\mathscr{T}_N):=\left\{v:v_{| au_k}\in\mathbb{P}_p( au_k) ext{ for all } au_k\in\mathscr{T}_N ext{ and }v_{|\Sigma}=0
ight\}.$$

The proposed space—time approach is based on the use of an interior penalty Galerkin approximation of the Laplace operator and an upwind scheme for the approximation of the time derivative, see, e.g., [3, 5]. Hence we have to find  $u_h \in S_{h,0}^p(\mathscr{T}_N)$  such that 32

$$a_{\mathrm{DG}}(u_{h}, v_{h}) := -\sum_{k=1}^{N} \int_{\tau_{k}} u_{h} \partial_{t} v_{h} dx dt + \int_{\Sigma_{T}} u_{h} v_{h} dx$$

$$+ \sum_{e \in \mathscr{I}_{N}} \int_{e} n_{t} \left\{ u_{h} \right\}_{e}^{\mathrm{up}} [v_{h}]_{e} ds_{(x,t)} + \sum_{k=1}^{N} \int_{\tau_{k}} \nabla_{x} u_{h} \cdot \nabla_{x} v_{h} dx dt$$

$$- \sum_{e \in \mathscr{I}_{N}} \int_{e} [\langle \mathbf{n}_{x} \cdot \nabla_{x} u_{h} \rangle_{e} [v_{h}]_{e} - \varepsilon [u_{h}]_{e} \langle \mathbf{n}_{x} \cdot \nabla_{x} v_{h} \rangle_{e}] ds_{(x,t)}$$

$$+ \frac{\sigma}{h} \sum_{e \in \mathscr{I}_{N}} \int_{e} |\mathbf{n}_{x}|^{2} [u_{h}]_{e} [v_{h}]_{e} ds_{(x,t)}$$

$$= \int_{O} f v_{h} dx dt + \int_{\Sigma_{0}} u_{0} v_{h} dx =: F(v_{h})$$

$$(4)$$

is satisfied for all  $v_h \in S_{h,0}^p(\mathscr{T}_N)$ . The parameters  $\sigma$  and  $\varepsilon$  have to be chosen appropriately. For  $v_h \in S_{h,0}^p(\mathscr{T}_N)$  and  $\sigma > 0$  the related energy norm is given by

$$\|v_h\|_{\mathrm{DG}}^2 := \|v_h\|_A^2 + \|v_h\|_B^2,$$
 35

where 36

$$\begin{aligned} \|v_h\|_A^2 &:= \sum_{k=1}^N \|\nabla_x v_h\|_{\tau_k}^2 + \frac{\sigma}{h} \sum_{e \in \mathscr{I}_N} \||\mathbf{n}_x| [v_h]_e\|_{L_2(e)}^2, \\ \|v_h\|_B^2 &:= h \sum_{k=1}^N \|\partial_t v_h\|_{\tau_k}^2 + \frac{1}{2} \|v_h\|_{L_2(\Sigma_0 \cup \Sigma_T)}^2 + \frac{1}{2} \sum_{e \in \mathscr{I}_N} \|\sqrt{|n_t|} [v_h]_e\|_{L_2(e)}^2. \end{aligned}$$

The unique solvability of the variational formulation (4) is based on the following 37 stability result.

**Lemma 1.** Let  $\varepsilon \in \{-1,0,1\}$  and  $\sigma > 0$ . For  $\varepsilon \in \{-1,0\}$  let  $\sigma$  be sufficient large. 39 *Then the stability estimate* 

$$\sup_{0 \neq v_h \in S_{h,0}^p(\mathscr{T}_N)} \frac{a_{\mathrm{DG}}(u_h, v_h)}{\|v_h\|_{\mathrm{DG}}} \ge c_1^A \|u_h\|_{\mathrm{DG}} \quad \text{for all } u_h \in S_{h,0}^p(\mathscr{T}_N)$$

is satisfied where the constant  $c_1^A$  depends on the shape of the finite elements, and 42 on the stabilization parameter  $\sigma$ . However, for a sufficient large choice of  $\sigma$  we can 43 ensure  $c_1^A = \frac{1}{2}$ .

*Proof.* The proof follows as in [5], by using the technique as in [2]; see also [3].  $\Box$ 

By using standard arguments we can then conclude the energy error estimate

$$||u - u_h||_{DG} \le ch^{\min\{s, p+1\}-1} |u|_{H^s(O)}$$

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when assuming  $u \in H^s(Q)$  for some  $s \le p+1$ , and, by applying the Aubin–Nitsche 47 trick, for  $\varepsilon = -1$ ,

$$||u - u_h||_{L_2(\Omega)} \le ch^{\min\{s, p+1\}} |u|_{H^s(Q)}.$$
 (5)

To illustrate the proposed DG finite element method in space and time as well as the 49 given error estimates we consider a first numerical example for the initial boundary 50 value problem (1)–(3) for n = 1 and  $\Omega = (0,1)$ , T = 1. This implies  $Q = (0,1)^2$ . The 51 given data f and  $u_0$  are chosen such that the solution is given as

$$u(x,t) = \sin(\pi x)(1-t)^{3/4} \in H^{1,25-\bar{\epsilon}}(Q)$$
 with  $\bar{\epsilon} > 0$ .

Starting from a triangulation of  $Q=(0,1)^2$  into four triangles we consider a sequence of several uniform refinement steps to analyze the convergence behavior of 55 the presented method. Using piecewise linear basis functions, i.e. p=1,  $\varepsilon=-1$  and 56  $\sigma=10$ , the numerical results are given in Table 1 which confirm the convergence 57 rate of 1.25 as predicted by the error estimate (5).

level	elements	dof	$  u - u_h  _{L_2(Q)}$	eoc
0	4	8	2.2679 - 1	_
1	16	40	5.1354 - 2	2.14
2	64	176	1.3107 - 2	1.97
3	256	736	3.4813 - 3	1.91
4	1024	3008	9.7383 - 4	1.84
5	4096	12160	3.0406 - 4	1.68
6	16384	48896	1.0923 - 4	1.48
7	65536	196096	4.3315 - 5	1.33
8	262144	785408	1.7935 - 5	1.27
9	1048576	3143680	7.5278 - 6	1.25
10	4194304	12578816	3.1694 - 6	1.25
11	16777216	50323456	1.3345 - 6	1.25

**Table 1.** Numerical results for p = 1,  $\varepsilon = -1$  and  $\sigma = 10$ .

### 2 A Hybrid Space-Time Domain Decomposition Method

The presented space–time method (4) results in a large linear system of algebraic 60 equations. For its iterative solution we introduce a hybrid formulation as in [1, 2]. 61 Therefore we subdivide the space–time domain Q into P non–overlapping subdomains  $Q_i$ , i = 1, ..., P, 63

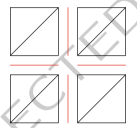
$$\overline{Q} = \bigcup_{i=1}^{P} \overline{Q}_i, \quad Q_i \cap Q_j = \emptyset \quad \text{for } i \neq j.$$

59

By 65

$$\Gamma := \bigcup_{i=1}^{P} \Gamma_{i} \quad \text{with } \Gamma_{i} := \overline{\partial Q_{i} \setminus \partial Q}$$

we denote the interface of the space–time domain decomposition, see Fig. 1.



**Fig. 1.** Space–time decomposition of Q and the interface  $\Gamma$ 

With respect to the interface  $\Gamma$  we introduce the discrete function space of piecewise 68 polynomials of order p, 69

$$S_h^p(\Gamma) := \left\{ v \in L_2(\Gamma) : v_{|e} \in \mathbb{P}_p(e) \text{ for all } e \in \mathscr{I}_N \text{ with } e \subseteq \Gamma \right\}.$$

For the solution of the local partial differential equations in all subdomains  $Q_i$  we 71 apply the space–time method as described by the variational formulation (4). For this 72 we denote by  $a_{\mathrm{DG}}^{(i)}(\cdot,\cdot)$  the restriction of the bilinear form  $a_{\mathrm{DG}}(\cdot,\cdot)$  on the subdomain 73  $Q_i$ ,  $i=1,\ldots,P$ , i.e.

$$\begin{split} a_{\mathrm{DG}}^{(i)}(u_h,v_h) &:= -\sum_{k=1}^N \int_{\tau_k \cap Q_i} u_h \, \partial_t v_h \, dx dt + \int_{\Sigma_T \cap \partial Q_i} u_h v_h \, dx \\ &+ \sum_{e \in \mathscr{I}_N} \int_{e \cap Q_i} n_t \, \{u_h\}_e^{\mathrm{up}} [v_h]_e \, ds_{(x,t)} + \sum_{k=1}^N \int_{\tau_k \cap Q_i} \nabla_x u_h \cdot \nabla_x v_h \, dx dt \\ &- \sum_{e \in \mathscr{I}_N} \int_{e \cap Q_i} [\langle \mathbf{n}_x \cdot \nabla_x u_h \rangle_e [v_h]_e - \varepsilon \, [u_h]_e \, \langle \mathbf{n}_x \cdot \nabla_x v_h \rangle_e] \, ds_{(x,t)} \\ &+ \frac{\sigma}{h} \sum_{e \in \mathscr{I}_N} \int_{e \cap Q_i} |\mathbf{n}_x|^2 \, [u_h]_e [v_h]_e ds_{(x,t)}. \end{split}$$

Accordingly, the restriction of the linear form  $F(\cdot)$  on a subdomain  $Q_i$  is given by

$$F^{(i)}(v_h) := \int_{Q_i} f v_h dx dt + \int_{\Sigma_0 \cap \partial Q_i} u_0 v_h dx.$$
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For the coupling of the local fields we first introduce a new unknown on the interface, 77

$$\lambda := \langle u \rangle_e = \frac{1}{2} \left[ u_{|\tau_k} + u_{|\tau_\ell} \right] \quad \text{on } \Gamma \cap e.$$

With this we can rewrite the jump of a function as

$$[u]_e=u_{| au_k}-u_{| au_\ell}=2\left(u_{| au_k}-\lambda
ight)=2\left(\lambda-u_{| au_\ell}
ight)\quad ext{on }\Gamma\cap e.$$

Therefore we obtain for the coupling terms related to the Laplace operator

$$\begin{split} &\sum_{e \in \mathscr{I}_N} \int_{e \cap \Gamma} \langle \mathbf{n}_x \cdot \nabla_x u \rangle_e [v]_e \, ds_{(x,t)} = \sum_{k=1}^N \int_{\partial \tau_k \cap \Gamma} \mathbf{n}_{k,x} \cdot \nabla_x u (v - \mu) \, ds_{(x,t)}, \\ &\sum_{e \in \mathscr{I}_N} \int_{e \cap \Gamma} [u]_e \, \langle \mathbf{n}_x \cdot \nabla_x v \rangle_e \, ds_{(x,t)} = \sum_{k=1}^N \int_{\partial \tau_k \cap \Gamma} (u - \lambda) \, \mathbf{n}_{k,x} \cdot \nabla_x v \, ds_{(x,t)}, \\ &\sum_{e \in \mathscr{I}_N} \int_{e \cap \Gamma} |\mathbf{n}_x|^2 \, [u]_e \, [v]_e \, ds_{(x,t)} = 2 \sum_{k=1}^N \int_{\partial \tau_k \cap \Gamma} |\mathbf{n}_{k,x}|^2 \, (u - \lambda) (v - \mu) \, ds_{(x,t)}. \end{split}$$

For the classical solution u of the transient heat equation (1)–(3) there obviously 82 holds for an interior facet  $e \in \mathscr{I}_N$  83

$$\lambda = \langle u \rangle_e = \frac{1}{2} \left[ u_{|\tau_k} + u_{|\tau_\ell} \right] = u_{|\tau_k} = u_{|\tau_\ell} \quad \text{on } e.$$

Therefore the upwind in time can be written as

$$\{u\}_e^{\mathrm{up}} = \begin{cases} u_{\mid \tau_k} & \text{for } n_t \geq 0, \\ u_{\mid \tau_t} & \text{for } n_t < 0 \end{cases} = \begin{cases} u_{\mid \tau_k} & \text{for } n_{k,t} \geq 0, \\ \lambda & \text{for } n_{k,t} < 0 \end{cases} =: \{u/\lambda\}_{\partial \tau_k}^{\mathrm{up}} & \text{on } \Gamma \cap e.$$

The coupling containing the upwind part can now be expressed by

$$\sum_{e \in \mathscr{I}_N} \int_{e \cap \Gamma} n_t \{u\}_e^{\text{up}}[v]_e \, ds_{(x,t)} = \sum_{k=1}^N \int_{\partial \tau_k \cap \Gamma} n_{k,t} \{u/\lambda\}_{\partial \tau_k}^{\text{up}}(v-\mu) \, ds_{(x,t)}. \tag{88}$$

With respect to each subdomain  $Q_i$  we therefore can define the bilinear form

$$\begin{split} c^{(i)}(u_h,\lambda_h;v_h,\mu_h) &:= \sum_{\substack{k=1\\\tau_k\subseteq Q_i}}^N \int_{\partial \tau_k\cap\Gamma} n_{k,t} \left\{ u_h/\lambda_h \right\}_{\partial \tau_k}^{\mathrm{up}}(v_h-\mu_h) \, ds_{(x,t)} \\ &- \sum_{\substack{k=1\\\tau_k\subseteq Q_i}}^N \int_{\partial \tau_k\cap\Gamma} \left[ \mathbf{n}_{k,x} \cdot \nabla_x u_h \left( v_h - \mu_h \right) - \varepsilon (u_h - \lambda_h) \, \mathbf{n}_{k,x} \cdot \nabla_x v_h \right] \, ds_{(x,t)} \\ &+ \frac{2\sigma}{h} \sum_{\substack{k=1\\\tau_h\subset O_i}}^N \int_{\partial \tau_k\cap\Gamma} |\mathbf{n}_{k,x}|^2 \left( u_h - \lambda_h \right) (v_h - \mu_h) \, ds_{(x,t)}. \end{split}$$

Hence we can write the discrete hybrid space-time variational formulation to find 90  $u_h \in S_{h,0}^p(\mathscr{T}_N)$  and  $\lambda_h \in S_h^p(\Gamma)$  satisfying

$$\sum_{i=1}^{P} \left[ a_{\text{DG}}^{(i)}(u_h, v_h) + c^{(i)}(u_h, \lambda_h; v_h, \mu_h) \right] = \sum_{i=1}^{P} F^{(i)}(v_h)$$
 (6)

for all  $v_h \in S_{h,0}^p(\mathscr{T}_N)$  and  $\mu_h \in S_h^p(\Gamma)$ . As in [2] we can prove unique solvability of the 92 hybrid scheme (6). Moreover, related error estimates as derived for the DG scheme 93 remain valid.

The discrete variational formulation (6) is equivalent to the solution of the linear 95 equations

$$\begin{pmatrix} A_{II}^{(1)} & A_{I\Gamma}^{(1)} \\ A_{II}^{(2)} & A_{I\Gamma}^{(2)} \\ & \ddots & \vdots \\ & A_{II}^{(P)} & A_{I\Gamma}^{(P)} \\ A_{\Gamma I}^{(1)} & A_{\Gamma I}^{(2)} & \cdots & A_{\Gamma I}^{(P)} & A_{\Gamma \Gamma} \end{pmatrix} \begin{pmatrix} \mathbf{u}_{I}^{(1)} \\ \mathbf{u}_{I}^{(2)} \\ \vdots \\ \mathbf{u}_{I}^{(P)} \\ \lambda_{\Gamma} \end{pmatrix} = \begin{pmatrix} \mathbf{f}_{I}^{(1)} \\ \mathbf{f}_{I}^{(2)} \\ \vdots \\ \mathbf{f}_{I}^{(P)} \\ \mathbf{f}_{\Gamma} \end{pmatrix}$$

$$(7)$$

where the local block matrices  $A_{II}^{(i)}$  correspond to the local bilinear forms  $a_{\mathrm{DG}}^{(i)}(\cdot,\cdot)$  97 and  $c^{(i)}(\cdot,0;\cdot,0)$ , while the remaining block matrices describe the coupling across the interface. For an appropriate choice of the DG parameters, see Lemma 1, the local 99 matrices  $A_{II}^{(i)}$  are invertible. Hence we obtain the Schur complement system 100

$$\left[ A_{\Gamma\Gamma} - \sum_{i=1}^{P} A_{\Gamma I}^{(i)} \left( A_{II}^{(i)} \right)^{-1} A_{I\Gamma}^{(i)} \right] \lambda_{\Gamma} = \mathbf{f}_{\Gamma} - \sum_{i=1}^{P} A_{\Gamma I}^{(i)} \left( A_{II}^{(i)} \right)^{-1} \mathbf{f}_{I}^{(i)},$$

$$\mathbf{u}_{I}^{(i)} = \left( A_{II}^{(i)} \right)^{-1} \left[ \mathbf{f}_{I}^{(i)} - A_{I\Gamma}^{(i)} \lambda_{\Gamma} \right] \quad \text{for } i = 1, \dots, P.$$
(8)

with

$$\mathbf{u}_{I}^{(i)} = \left(A_{II}^{(i)}\right)^{-1} \left[\mathbf{f}_{I}^{(i)} - A_{I\Gamma}^{(i)} \lambda_{\Gamma}\right] \quad \text{for } i = 1, \dots, P.$$

The inversion of the local matrices  $A_{II}^{(i)}$  can be done in parallel either by using some 102 appropriate direct approach, or suitable iterative schemes. For the solution of the 103 global Schur complement system (8) we can use, for example the GMRES method.

## 3 Numerical Examples

To illustrate the hybrid domain decomposition approach we consider for n=3 the spatial domain  $\Omega = (0,1)^3$  and T = 1, i.e.  $Q = (0,1)^4$ . As initial triangulation for the space-time domain we use 96 pentatopes of the same size, see also [4]. The initial 108 triangulation is used as a partition of the space-time domain into P = 96 subdomains, 109 which we keep fixed for all computations. As exact solution of the transient heat 110 equation (1) we now consider the smooth function

$$u(x,t) = \sin(\pi x_1)\sin(\pi x_2)\sin(\pi x_3)t^2.$$

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For the iterative solution of the Schur complement system (8) we use the GMRES 113 method without preconditioning with a relative error reduction of  $\varepsilon_{\text{GMRES}} = 10^{-8}$ . In 114 the Tables 2 and 3 we present the iteration numbers of the GMRES method for different levels of a uniform refinement of the space—time mesh for p=1 and p=2. We 116 observe that the number of required iterations grows slightly indicating the need of 117 using an appropriate preconditioner. The results also show the optimal convergence 118 rates for the error in the  $L_2(Q)$  norm when using linear and quadratic basis functions. 119

level	elements	$\mathrm{dof}~\mathbf{u}_{I}^{(i)}$	$dof  \lambda_{\varGamma}$	iter.	$  u-u_h  _{L_2(Q)}$	eoc
0	96	192	768		6.120 - 2	E
1	1536	5376	6144	143	3.821 - 2	0.68
2	24576	104448	49152	197		
3	393216	1818624	393216	294	4.024 - 3	1.75
4	6291456	30277632	3145728	475	1.111 - 3	1.86

**Table 2.** Numerical results with 96 subdomains for p = 1,  $\varepsilon = -1$  and  $\sigma = 10$ .

level	elements	$\operatorname{dof} \mathbf{u}_{I}^{(i)}$	$\mathrm{dof}\lambda_{\varGamma}$	iter.	$  u-u_h  _{L_2(Q)}$	eoc
0	96	720	1920	404	4.199 - 2	_
1					7.492 - 3	
2	24576	322560	122880	900	1.005 - 3	2.90
3	393216	5529600	983040	1131	1.293 - 4	2.96

**Table 3.** Numerical results with 96 subdomains for p = 2,  $\varepsilon = -1$  and  $\sigma = 10$ .

4 Conclusions 120

In this paper we have presented a hybrid DG domain decomposition approach for the parallel solution of initial boundary value problems. Numerical examples for one—and three—dimensional spatial domains indicate the accuracy and applicability of the proposed method. However, the numerical results also indicate the need to use an appropriate global preconditioner for the Schur complement system (8). Moreover, when solving the coupled system (7) iteratively, suitable local preconditioners are mandatory as well. A possible choice is to use space-time multigrid methods. Although we have only considered uniform refinements in this paper, the proposed approach is also applicable to non—uniform and adaptive refinements, see, for example, [4]. For this we need to use suitable a posteriori error estimators, and the solution algorithms need to be robust with respect to adaptive refinements. Although we have

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only considered the simple model problem of the transient heat equation, the proposed approach can be extended to more complicated problems, see, e.g., [4] for a first example for the transient Navier-Stokes system.	
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## **Parallel Adaptive Deflated GMRES**

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**Summary.** Many scientific libraries are currently based on the GMRES method as a Krylov 7 subspace iterative method for solving large linear systems. The restarted formulation known as 8 GMRES(*m*) has been extensively studied and several approaches have been proposed to reduce 9 the negative effects due to the restarting procedure. A common effect in GMRES(*m*) is a slow 10 convergence rate or a stagnation in the iterative process. In this situation, it is less attractive 11 as a general solver in industrial applications. In this work, we propose an adaptive deflation 12 strategy which retains useful information at time of restart to avoid stagnation in GMRES(*m*) 13 and improve its convergence rate. We give a parallel implementation in the PETSc package. 14 The provided numerical results show that this approach can be effectively used in the hybrid 15 direct/iterative methods to solve large-scale systems.

1 Introduction

The GMRES method due to [11] is widely used, thanks to its monotonic convergence properties, as a Krylov subspace method for solving large and sparse linear systems. Due to memory and computational requirements, the restarted GMRES (noted as GMRES(m)) is generally used. At the time of restart, information from the previous Krylov subspace is discarded and the orthogonality between successive Krylov subspaces is not preserved. The worst case is when the successive generated Krylov subspaces are very close. As a result, there is no significant reduction in the residual norm and the iterative process may stagnate. Deflation techniques are a class of acceleration strategies that collects useful information at the time of restart mainly to 26 avoid this stagnation and improve the convergence rate. The main idea behind these methods is to remove the smallest eigencomponents from the residual vector as they are known to slow down the convergence of GMRES.

In a practical use of a deflation strategy, it is necessary to define the number 30 of eigenvalues to deflate. As the deflation process induces additional operations to 31 GMRES(m), it is interesting as well to know a priori if the deflation will be beneficial. In this work, we propose an adaptive deflated GMRES(m) which aims at enhancing the convergence of GMRES(m) by adaptively extracting the spectral information 34

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needed to speedup the convergence. The adaptive strategy is based on a (near) stagnation test which defines if the deflation process is needed or not and if more accurate 36 spectral information are required. Although we use a stagnation test similar to that 37 in [12], our approach is different since we assume that the restart length m is fixed. 38 This work is motivated by the convergence behavior of GMRES when it is used with 39 a Schwarz preconditioner. As the number of subdomains increases, the eigenvalues 40 are less and less clustered. The restarting may have the disadvantage to discard the 41 smallest eigenvalues before their convergence. The proposed adaptive strategy will 42 thus keep these spectral values in the Krylov subspace until their convergence.

The remaining part of this report is organized as follows: in Sect. 2, we first recall 44 the basis of the deflation technique applied as a preconditioner and we derive the 45 adaptive strategy. In Sect. 3, we discuss on the parallel implementation. Section 4 46 is focused on numerical experiments to show the benefits of this scheme on a real 47 industrial CFD test case.

### 2 Adaptive Preconditioner for the Deflated GMRES(m)

We are interested in the solution of the linear system

$$Ax = b \tag{1}$$

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The GMRES method is among the best methods to solve this system when the co-51 efficient matrix A is nonsingular and nonsymmetric. For large linear systems, the 52 restarted version should always be used to reduce the memory and computational 53 requirements. The deflated GMRES has been proposed to reduce the negative ef- 54 fects of the restarting procedure. The general idea behind these methods is to add 55 to the Krylov subspace an approximation of the invariant subspace associated to the 56 smallest eigenvalues. In [7], this is carried out by defining a preconditioner that is 57 equal to the projected matrix onto the approximated invariant subspace and is taken 58 as the identity on the orthogonal subspace. Hence, given  $U = [u_1, \dots, u_r] \in \mathbb{R}^{n \times r}$  the 59 r-dimensional basis of the invariant subspace associated to the eigenvalues to deflate, 60 the preconditioner is defined as

$$M_D^{-1} \equiv I_n + U(|\lambda_n|T_r^{-1} - I_r)U^T, \quad T = U^T B U,$$
 (2)

where  $\lambda_n$  is the largest eigenvalue in magnitude,  $I_n$  and  $I_r$  are the identity matrices 62 and B the initial preconditioned matrix. Since  $M_D^{-1}$  is nonsingular, the eigenvalues of 63 the resulted matrix  $M_D^{-1}B$  or  $BM_D^{-1}$  are  $\lambda_{r+1},\ldots,\lambda_n,|\lambda_n|$  with a multiplicity at least r. 64 It is therefore expected to get a faster convergence rate with this preconditioner since 65 the r smallest eigencomponents that slow down the convergence are deflated. This 66 assumes that U is a good approximation of the basis of the selected invariant sub- 67 space. For large matrices however, the cost of accurately computing U (as suggested 68 in [7] and later in [4]) may induce a significant overhead. This process should be 69 carried out only when it is necessary, for instance to avoid stagnation.

#### **Algorithm 4** DGMRES(m,k,r): Restarted GMRES with adaptive deflation

```
1: input (m, itmax, \varepsilon, k, smv, bgv, rmax);
 2: Set B \equiv AM^{-1}, M^{-1} is any external preconditioner
 3: r_0 = b - Ax_0; U = []; M_D = I; it = 0; r = 0;
 4: while (||r_0|| > \varepsilon)
        Arnoldi process on B to get BM_D^{-1}V_m = V_{m+1}\bar{H}_m. See [11]
        x_m = x_0 + M_D^{-1} M^{-1} V_m y_m, y_m solution of min \|\beta e_1 - \bar{H}_m y_m\|_2;
 6:
 7:
         r_m = b - Ax_m, it \leftarrow it + m;
 8:
         If (||r_m|| > \varepsilon \text{ and } it < itmax) then
            Iter = m * log(\frac{\varepsilon}{\|r_m\|}) / log(\frac{\|r_m\|}{\|r_0\|});
 9:
10:
            If (Iter > smv * (itmax - it)) and r < rmax) then
               Compute k Schur vectors of B noted X. See [7]
11:
12:
               Orthogonalize X against U
               Compute T = \begin{bmatrix} U & X \end{bmatrix}^T B \begin{bmatrix} U & X \end{bmatrix} \equiv \begin{pmatrix} U^T B U & U^T B X \\ X^T B U & X^T B X \end{pmatrix}
13:
14:
                Increase U by X; r \leftarrow r + k;
15:
                If(Iter > bgv * (itmax - it)) then
                   Improve U as indicated in [4, Sect. 3]
16:
17:
                                   Set M_D^{-1} \equiv I_n + U(|\lambda_n|T)
                Factorize T
18:
19:
            End If
20:
         End If
21:
         x_0 = x_m
22: end while
```

We thus propose here an adaptive strategy that detects a near-stagnation in the 71 iterative process or a slow reduction in the residual norm. This approach is based 72 upon the work by Sosonkina et al. [12] in which the Krylov subspace is adaptively 73 increased along the cycles of GMRES(m); Here, we find it natural to enrich the sub- 74 space with the eigencomponents that slow down the convergence. The main steps are 75 given in Algorithm 4. First, m steps of the Arnoldi process are performed to compute 76 the orthonormal basis  $V_m$ . It also creates an upper Hessenberg matrix  $H_m = V_m^T B V_m$  77 which is the restriction of B onto the m-dimensional Krylov subspace. Then, a leastsquares problem is solved to minimize the residual norm in the Krylov subspace. At 79 the time of restart, if the desired residual norm is not achieved, a stagnation test 80 is computed to determine if a deflation process could be beneficial to accelerate the 81 convergence. This test considers the convergence rate over the previous restart cycles 82 and evaluates the number of iterations (*Iter*) needed to achieve the desired accuracy. 83 If Iter is greater than the remaining number of steps (bounded by a small multi- 84 ple smv of the number of iterations allowed), then data are computed to update the 85 preconditioner associated to the deflation process. This test is therefore used to re- 86 duce the iteration counts in GMRES(m). To detect a near-stagnation, we use another 87 test which considers a large multiple bgv of the remaining number of steps. In this 88 case, a harmonic projection is carried out to accurately compute the eigenvalues and 89 continuously update the previous estimation of U.

### 3 Implementation Notes

We now give some details about the implementation of Algorithm 4 on distributed- 92 memory computers. The programming model is SPMD (Single Program Multiple 93 Data) and communications are done using the message-passing interface (MPI). The 94 adjacency graph of the input sparse matrix is first built. PARMETIS is then used 95 to partition the vertices of the graph into D disjoint vertices. From this partition- 96 ing, the matrix is distributed such that each processor holds a contiguous chunk of 97 rows corresponding to the vertices it owns. The right hand side and all other vectors 98 (Krylov basis, invariant basis) are distributed accordingly. Note that the goal of this 99 data distribution is to get a good load balance and to minimize communication during 100 matrix-vector multiply and preconditioning steps. When the additive Schwarz pre- 101 conditioner is used, an overlapping partitioning can be defined by taking recursively 102 adjacent vertices from the initial disjoint partitions.

The main parallel operations in Algorithm 4 so far are the matrix-vector multiply, scalar products, and the application of  $M^{-1}$  and  $M_D^{-1}$ .  $M^{-1}$  can be any parallel 105 preconditioner as long as it implements the basic operation  $v_i \leftarrow M^{-1}v_i$ . In our tests, 106 the restricted additive Schwarz has been used as defined in [5]. It is then necessary 107 in the setup phase to factorize in each process the block matrices  $A_p$  corresponding 108 to the restriction of A onto the defined subdomains.  $M_D^{-1}$  is applied to a distributed 109 vector  $v_i$  in a straightforward manner given the data distribution described above. 110 This implies r all-to-all communications to compute the projection onto the invariant subspace. There is no additional communication for the other terms since the  $r \times r$  112 dense matrix T is owned by each process.

We provide an implementation of this method using the PETSc package (see 114 [3]). The original implementation of the built-in KSP GMRES has been modified to 115 provide the data needed for the deflation and to apply the resulting preconditioner 116 to generate the Krylov basis. Although the current presentation does not discuss the 117 choice of side of preconditioning, the implementation does define left and right pre- 118 conditioning. Note that the current adaptive preconditioning can be associated with 119 any other preconditioner available in the package or defined by the end user since we 120 provide generic interface similar to the other Krylov subspace methods in the pack- 121 age. The resulted KSP module (named as DGMRES) is available in PETSc release 122 3.2.

## **4 Numerical Experiments**

This section presents some numerical results to prove the efficiency of the proposed 125 approaches. The test problem arises from design optimization in computational fluid 126 dynamics. The physical model is a 3D flow simulation in a jet engine compressor 127 rotor. The physical equations are the Reynolds-Averaged Navier-Stokes for com- 128 pressible flows, discretized using the finite volume method as presented by Aubert 129 et al. [2]. The matrices have been extracted from the software Turb'Opty $^{TM}$  designed by the FLUOREM company. They are also available in the University of 131

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Florida sparse matrix collection (see [6]) under the name RM07R in the FLUO- 132 REM group. The matrix is nonsymmetric and indefinite with a size 272,635 and 133 37,355,908 nonzero entries. Other test cases can be found in [8].

With this test case so far, previous studies have shown the limits of some existing 135 solvers in terms of memory usage and numerical accuracy (see [9]). Pacull et al. [10] 136 have proved as well the instability of the ILU factorization to approximate the solution of linear subsystems. In our hybrid approach, we therefore rely on a direct solver 138 within each subdomain, such as MUMPS [1].

### 4.1 Benefits of the Deflated Restarting

We now give the main benefits of using the deflated GMRES with the additive 141 Schwarz method (ASM). It is known that one level ASM is a weak preconditioner 142 when the number of subdomains D gets large. The size of the Krylov subspace m 143 could then be increased to enhance the robustness of the global method. However, 144 choosing a good size m of the Krylov subspace is a trial-and-error process. With the 145 adaptive deflation, we show experimentally that the method is robust for various val- 146 ues of m and D. Moreover, using a large number of subdomains reduces the memory 147 required to handle the submatrices by the direct solver. Hence it is expected that the 148 time to factorize these matrices and the memory required will get smaller as D in- 149 creases. This is reported in the last column of Table 1. We also report the number of 150 matrix-vector multiplies and the global CPU time with respect to the number of sub- 151 domains D. We then compare the restarted version (GMRES(m)) with the deflated 152 version (DGMRES(m,k)), where m=48 and 64. A dash in a field means that the relative residual norm of  $10^{-8}$  is not reached after 2500 iterations. It can be observed 154 that DGMRES provides reliable and faster convergence than the classical restarted 155 GMRES. It also gives a faster method since significantly fewer iterations are needed. 156 Furthermore, the method reveals a substantial acceleration as the number of processors increases. Note that without the deflation, this acceleration will not be obtained 158 since the number of matrix-vector multiplies increases hugely with the subdomains. 159 For instance, this behavior can be seen with GMRES(64) when using D = 16 and 160 D = 32.

Table 1. RM07R: Benefits of using DGMRES with an additive Schwarz preconditioner and an overlap of 1. The deflation process reduces the total number of iterations and helps to use a large number of subdomains and thus a large number of processors. Here, the number of processors is indeed equal to the number of subdomains.

D	GMRES	DGMRES(47,1)			GMRES	S(64)	DGMRES(63,1)			
D	Matvecs	Time	Matvecs	Time	r	Matvecs	Time	Matvecs	Time	r
16	551	230	212	173.4	3	355	193.8	208	168.9	2
32	-	-	533	109.2	4	2217	244.6	455	94.6	7
64	-	-	410	56.8	4	-	-	453	50.8	7
128	-	-	791	51.5	15	-	-	638	44.3	8

161

t1.1 t1.2 t1.3

t1.5 t1.6

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### 4.2 Adaptive DGMRES and Full GMRES

From the robustness standpoint, the full GMRES approach is more reliable than the 163 restarted version even with the deflation process. However as the size of the basis 164 grows, it should be more sensitive to round-off errors. To illustrate this behavior, 165 we consider two formulations of the Arnoldi process, namely the classical Gram- 166 Schmidt (CGS) and the modified Gram-Schmidt (MGS) algorithms. The former is 167 sometimes preferred since it provides good kernel operations in parallel environ- 168 ments. In the PETSc package, for instance, it is used by default in the GMRES im- 169 plementation as the orthogonalization method with a possible iterative refinement 170 strategy. In Fig. 1, the residual history is displayed with respect to the number of 171 matrix-vector products. The method stops when the relative residual norm is  $10^{-10}$ . 172 It can then be noticed that with CGS, stagnation occurs in the full GMRES (in solid 173 line) due to severe cancellation in the algorithm and consequently a loss of orthogonality. This does not happen when the basis is small since the round-off errors are 175 not propagated very far and DGMRES (dash-dotted line) converges at the desired 176 accuracy even with CGS. Note that although good accuracy is finally achieved in

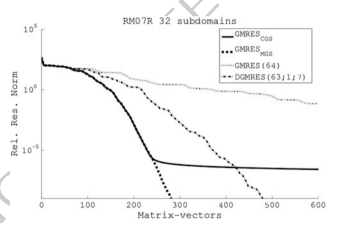


Fig. 1. Convergence of full GMRES, GMRES(m) and DGMRES(m,k,r) with classical Gram-Schmidt(CGS) and modified Gram-Schmidt (MGS) orthogonalization scheme. k is the number of eigenvalues to extract at each detected stagnation and r is the total number of eigenvalues extracted at the convergence. Thirty two subdomains are used in the additive Schwarz method with a 1-overlap

full GMRES with MGS (dashed line), it will require much more memory to store all 178 the vectors of the growing Krylov basis (265 vectors in this case). In DGMRES, the 179 Krylov basis is stored just for one cycle. Only the invariant basis U is stored over 180 the restart cycles together with vectors  $M^{-1}AU$  to reduce the matrix-vector counts. 181 Thus in this example, only  $63 + 7 \times 2 = 77$  vectors are stored. Note also that this 182 number can be further reduced by using a smaller Krylov basis since convergence is 183 still good, as shown in Table 1.

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5 Conclusion 185

We have designed an adaptive deflation strategy that can be used for preconditioned 186 GMRES. We show in this paper that the proposed algorithm can be used to improve 187 the robustness and reduce both CPU time and memory required by hybrid solvers 188 based on a one level additive Schwarz method. We have implemented this method in 189 the new module DGMRES of the PETSc library.

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# **Quasi-optimality of BDDC Methods for MITC Reissner-Mindlin Problems**

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1 Introduction 13

The goal of this paper is to improve a condition number bound proven in [5] for a 14 Balancing Domain Decomposition Method by Constraints (BDDC) for the Reissner- 15 Mindlin plate bending problem discretized with MITC elements. This BDDC pre- 16 conditioner is based on selecting the plate rotations and deflection degrees of freedom 17 at the subdomain vertices as primal continuity constraints. In [5], we proved that the resulting BDDC algorithm is scalable in the number of subdomains N and independent of the plate thickness t and that the condition number  $\kappa$  of the preconditioned 20 Reissner-Mindlin plate problem is bounded by

$$\kappa \leq C(H/h),$$

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with C a constant independent of the plate thickness t, the mesh size h and the subdomain size H. In the present contribution, we prove the improved quasi-optimal 24 result

$$\kappa \le C(1 + \log^3\left(H/h\right)). \tag{26}$$

We remark that the MITC discretization of Reissner-Mindlin problems can lead to 27 very ill-conditioned discrete system, with condition number 28

$$\kappa_{no} \sim Ch^{-2}t^{-2}$$
.

Introduced in [11] and analyzed in [17, 21, 22], BDDC methods have evolved from 30 previous domain decomposition work on Balancing Neumann-Neumann methods. 31 BDDC algorithm have been extended in recent years from scalar elliptic problems 32 to almost incompressible elasticity [12, 24], the Stokes system [18], flow in porous 33

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media [28], and spectral element discretizations [15, 23, 24]. BDDC and overlapping 34 Schwarz methods for Reissner-Mindlin plate problems discretized with Falk-Tu elements have been studied in the recent Ph.D. thesis [16], while multigrid method for 36 plates have been studied in [26]. Among the several finite element works for plates, 37 we mention [2, 3, 7–10, 13, 14, 19, 20, 27]. 38

### 2 The MITC Reissner-Mindlin Plate Bending Problem

**Continuous problem.** Let  $\Omega$  be a polygonal domain in  $\mathbb{R}^2$  representing the midsurface of the plate, for simplicity assumed to be clamped on the whole boundary  $\partial \Omega$ . 41 The Reissner-Mindlin plate bending problem (see [1, 7]) reads

$$\begin{cases} \text{Find } \boldsymbol{\theta}^{ex} \in [H_0^1(\Omega)]^2, u^{ex} \in H_0^1(\Omega) \text{ such that} \\ a(\boldsymbol{\theta}^{ex}, \boldsymbol{\eta}) + \mu k t^{-2} (\boldsymbol{\theta}^{ex} - \nabla u^{ex}, \boldsymbol{\eta} - \nabla v) = (f, v) \quad \forall \boldsymbol{\eta} \in [H_0^1(\Omega)]^2, v \in H_0^1(\Omega), \end{cases}$$

with  $\mu$  the shear modulus, k is the shear correction factor, t the plate thickness,  $u^{ex}$  43 the deflection,  $\boldsymbol{\theta}^{ex}$  the rotation of the normal fibers and f the applied scaled normal 44 load. Moreover,  $(\cdot, \cdot)$  stands for the standard scalar product in  $L^2(\Omega)$  and  $a(\cdot, \cdot)$  is the 45 bilinear form

$$a(\boldsymbol{\theta}^{ex}, \boldsymbol{\eta}) = (\mathbb{C}\varepsilon(\boldsymbol{\theta}^{ex}), \varepsilon(\boldsymbol{\eta})),$$
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with  $\mathbb C$  the positive definite tensor of bending moduli and  $\varepsilon(\cdot)$  the symmetric gradient 48 operator. Introducing the scaled shear stresses  $\gamma^{ex} = \mu k t^{-2} (\theta^{ex} - \nabla \mu^{ex})$ , problem (1) 49 can be written in terms of the following mixed variational formulation, where for 50 simplicity we have assumed  $\mu k = 1$ : 51

$$\begin{cases} \text{Find } \boldsymbol{\theta}^{ex} \in [H_0^1(\Omega)]^2, u^{ex} \in H_0^1(\Omega), \boldsymbol{\gamma}^{ex} \in [L^2(\Omega)]^2 \text{ such that} \\ a(\boldsymbol{\theta}^{ex}, \boldsymbol{\eta}) + (\boldsymbol{\gamma}^{ex}, \boldsymbol{\eta} - \nabla v) = (f, v) \quad \forall \boldsymbol{\eta} \in [H_0^1(\Omega)]^2, v \in H_0^1(\Omega) \\ (\boldsymbol{\theta}^{ex} - \nabla u^{ex}, \boldsymbol{s}) - t^2(\boldsymbol{\gamma}, \boldsymbol{s}) = 0 \quad \forall \boldsymbol{s} \in [L^2(\Omega)]^2 . \end{cases}$$
 (2)

**Discrete problem.** We discretize the plate problem by MITC (Mixed Interpolation 52 of Tensorial Components) elements; see e.g. [1, 7, 8] for more details on this family 53 of elements. Let  $\tau_h$  denote a triangular or quadrilateral conforming finite element 54 mesh on  $\Omega$ , of characteristic mesh size h. Let  $\boldsymbol{\Theta}$ , U and  $\boldsymbol{\Gamma}$  be the discrete spaces for 55 rotations, deflections and shear stresses, respectively and define  $\mathbf{X} = \boldsymbol{\Theta} \times U$ . Then the 56 Reissner-Mindlin plate bending problem (2) discretized with MITC elements reads

$$\begin{cases} \text{Find } (\boldsymbol{\theta}, u) \in \mathbf{X}, \ \boldsymbol{\gamma} \in \boldsymbol{\Gamma} \text{ such that} \\ a(\boldsymbol{\theta}, \boldsymbol{\eta}) + (\boldsymbol{\gamma}, \boldsymbol{\Pi} \ \boldsymbol{\eta} - \nabla v) = (f, v) \quad \forall (\boldsymbol{\eta}, v) \in \mathbf{X} \\ (\boldsymbol{\Pi} \ \boldsymbol{\theta} - \nabla u, s) - t^2(\boldsymbol{\gamma}, s) = 0 \quad \forall s \in \boldsymbol{\Gamma} \end{cases}$$
(3)

where  $\Pi:([H^1(\Omega)]^2+\mathbf{\Gamma})\longrightarrow \mathbf{\Gamma}$  is the MITC reduction operator. Using the second 58 equation of (3), shear stresses can be eliminated to obtain the following positive 59 definite discrete formulation: 60

$$\begin{cases} \text{Find } (\boldsymbol{\theta}, u) \in \mathbf{X} \text{ such that} \\ b((\boldsymbol{\theta}, u), (\boldsymbol{\eta}, v)) = (f, v) \quad \forall (\boldsymbol{\eta}, v) \in \mathbf{X} \end{cases}$$
(4)

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where we have defined  $b((\boldsymbol{\theta}, u), (\boldsymbol{\eta}, v)) := a(\boldsymbol{\theta}, \boldsymbol{\eta}) + t^{-2}(\Pi \boldsymbol{\theta} - \nabla u, \Pi \boldsymbol{\eta} - \nabla v)$ . In 61 this paper, we address directly the positive definite problem (4), in the spirit of [4.5], 62 instead of the mixed formulation (3). For the convergence analysis of the MITC 63 elements, see e.g. [3, 8, 13, 25]. The MITC elements perform optimally with respect 64 to the polynomial degree and regularity of the solution, and their rate of convergence 65 is independent of the thickness parameter t.

### 3 Iterative Substructuring and BDDC Preconditioning

Subspace decomposition and Schur complement. We decompose the domain  $\Omega$  68 into N open, nonoverlapping subdomains  $\Omega_i$  of characteristic size H forming a 69 shape-regular finite element mesh  $\tau_H$ . This coarse triangulation  $\tau_H$  is further refined 70 into a finer triangulation  $\tau_h$  of characteristic size h; both meshes will typically be 71 composed of triangles or quadrilaterals. In the sequel, we assume that the material 72 tensor  $\mathbb{C}$  is constant on the whole domain.

As it is standard in iterative substructuring methods, we first reduce the problem 74 to the interface  $\Gamma = (\bigcup_{i=1}^N \partial \Omega_i) \setminus \partial \Omega$ , by implicitly eliminating the interior degrees 75 of freedom. In variational form, this process consists in a suitable decomposition of 76 the discrete space  $\mathbf{X} = \boldsymbol{\Theta} \times U$ . More precisely, let us define  $\mathbf{W} = \mathbf{X}_{|\Gamma}$ , i.e. the space 77 of the traces of functions in **X**, as well as the local spaces  $\mathbf{X}_i = \mathbf{X} \cap [H_0^1(\Omega_i)]^3$ . The 78 space **X** can be decomposed as  $\mathbf{X} = \bigoplus_{i=1}^{N} \mathbf{X}_{i} \oplus \overline{\mathscr{H}}(\mathbf{W})$ . Here  $\overline{\mathscr{H}} : \mathbf{W} \longrightarrow \mathbf{X}$  is the 79 discrete "plate-harmonic" extension operator defined by solving the problem

$$\begin{cases} \operatorname{Find} \overline{\mathscr{H}}(\boldsymbol{w}_{\Gamma}) \in \mathbf{X} \text{ such that } \overline{\mathscr{H}}(\boldsymbol{w}_{\Gamma})|_{\Gamma} = \boldsymbol{w}_{\Gamma} \text{ and} \\ b(\overline{\mathscr{H}}(\boldsymbol{w}_{\Gamma}), \boldsymbol{v}_{I}) = 0 \qquad \forall \boldsymbol{v}_{I} \in \mathbf{X}_{i} \quad i = 1, 2, \dots, N. \end{cases}$$

Defining the Schur complement bilinear form  $s(\mathbf{w}_{\Gamma}, \mathbf{v}_{\Gamma}) = b(\overline{\mathcal{H}}(\mathbf{w}_{\Gamma}), \overline{\mathcal{H}}(\mathbf{v}_{\Gamma}))$ , the 81 Schur complement system reads  $s(\boldsymbol{u}_{\Gamma}, \boldsymbol{v}_{\Gamma}) = \langle \tilde{\boldsymbol{f}}, \boldsymbol{v}_{\Gamma} \rangle \ \forall \boldsymbol{v}_{\Gamma} \in \boldsymbol{W}$ , for a suitable 82 right-hand side f.

The BDDC Reissner-Mindlin plate preconditioner. BDDC preconditioners, introduced in [11] and analyzed in [21], can be regarded as an evolution of Balancing 85 Neumann-Neumann preconditioners for the Schur complement system. In this sec- 86 tion, we briefly recall the BDDC preconditioner of [5].

Define  $\Gamma_i:=\partial\Omega_i$ , and  $\Gamma_{ij}=\partial\Omega_i\cap\partial\Omega_j$ ,  $i,j\in\{1,2,\ldots,N\}$ , the common edge 88 between two adjacent subdomains  $\Omega_i$  and  $\Omega_j$ . The local spaces  $\overline{W}_i$  are the spaces 89 of discrete functions defined by  $\overline{\mathbf{W}}_i = \mathbf{W}_{|\Gamma_i}$ , i = 1, 2, ..., N. Let  $\overline{\mathcal{H}}_i : \overline{\mathbf{W}}_i \longrightarrow \mathbf{X}|_{\Omega_i}$ , 90  $i=1,2,\ldots,N$ , represent the restriction of the operator  $\overline{\mathcal{H}}$  to the subdomain  $\Omega_i$ 91

$$\begin{cases} \text{Find } \overline{\mathscr{H}}_i(\mathbf{w}_i) \in \mathbf{X}|_{\Omega_i} \text{ such that } \overline{\mathscr{H}}(\mathbf{w}_i)|_{\Gamma_i} = \mathbf{w}_i \text{ and} \\ b_i(\overline{\mathscr{H}}_i(\mathbf{w}_i), \mathbf{v}_i) = 0 \qquad \forall \, \mathbf{v}_i \in \mathbf{X}_i, \end{cases}$$

where the  $b_i(\cdot,\cdot)$  are given by restricting the integrals in  $\underline{b(\cdot,\cdot)}$  to the domain  $\Omega_i$ , 92  $i=1,2,\ldots,N$ . The local bilinear forms are  $s_i(\boldsymbol{w}_i,\boldsymbol{v}_i)=b_i(\mathcal{H}_i\boldsymbol{w}_i,\mathcal{H}_i\boldsymbol{v}_i), \forall \boldsymbol{w}_i,\boldsymbol{v}_i\in$  93  $\overline{\boldsymbol{W}}_i$ . Let  $R_i^T$ ,  $i=1,2,\ldots,N$  be the prolongation operators which extend any function 94 of  $\overline{\boldsymbol{W}}_i$  to the function of  $\boldsymbol{W}$  which is zero at all the nodes not on  $\Gamma_i$ . Note that for 95  $\boldsymbol{w},\boldsymbol{v}\in\boldsymbol{W}, \sum_{i=1}^N s_i(R_i\boldsymbol{w},R_i\boldsymbol{v})=s(\boldsymbol{w},\boldsymbol{v})$ . For  $x\in\Gamma$ , we also define the weight  $N_x=96$   $\#\{j\in\mathbb{N}\mid x\in\partial\Omega_j\}$  and the weighted counting operators  $\delta_i:\overline{\boldsymbol{W}}_i\longrightarrow\overline{\boldsymbol{W}}_i$  (and their 97 inverses  $\delta_i^\dagger$ ) by

$$\delta_i \mathbf{v}_i(x) = N_x \mathbf{v}_i(x), \qquad \delta_i^{\dagger} \mathbf{v}_i(x) = N_x^{-1} \mathbf{v}_i(x), \quad \forall x \text{ node of } \Gamma_i \cap \Gamma.$$

Let  $C_i: \overline{W}_i \to \mathbb{R}^{3cc_i}$  be local constraint operators that read function values at the 100 corners of the subdomain  $\Omega_i$ , with  $cc_i$  the number of corners of the subdomain. Then 101 we define the local constrained spaces 102

$$oldsymbol{W}_i = \{oldsymbol{w}_i \in \overline{oldsymbol{W}}_i \, | \, C_i oldsymbol{w}_i = oldsymbol{0}\},$$
 103

and a global coarse space  $\mathbf{W}_0 \subset \mathbf{W}$  associated with the function values at the subdomain vertices. Given the number m of such subdomain vertices, let  $w_c \in \mathbb{R}^{3m}$  be a vector representing the respective nodal values. Then the space  $\mathbf{W}_0$  is defined by

$$\boldsymbol{W}_{0} = \{ \sum_{i=1}^{N} R_{i}^{T} \delta_{i}^{\dagger} \boldsymbol{w}_{0,i} \mid C_{i} \boldsymbol{w}_{0,i} = R_{i}^{C} w_{c}, w_{c} \in \mathbb{R}^{3m}, s_{i}(\boldsymbol{w}_{0,i}, \boldsymbol{w}_{0,i}) \rightarrow \min \},$$

with  $R_i^C$  the operator extracting the vertex values for the subdomain  $\Omega_i$  from the 107 global vector  $\boldsymbol{w}_c$  of all the subdomain vertex values. Any element  $\boldsymbol{w} \in \boldsymbol{W}$  can be 108 uniquely decomposed as  $\boldsymbol{w} = \boldsymbol{w}_0 + \sum_{i=1}^N \boldsymbol{w}_i$ , with  $\boldsymbol{w}_0 \in \boldsymbol{W}_0$ ,  $\boldsymbol{w}_i \in \boldsymbol{W}_i$  for  $i = 1, \dots, N$ . 109 We use inexact bilinear forms defined by

$$\tilde{s}_i(\boldsymbol{w}_i, \boldsymbol{v}_i) = s_i(\boldsymbol{\delta}_i \boldsymbol{w}_i, \boldsymbol{\delta}_i \boldsymbol{v}_i) \qquad \forall \boldsymbol{w}_i, \boldsymbol{v}_i \in \boldsymbol{W}_i, \ i = 1, 2, \dots, N, \\
\tilde{s}_0(\boldsymbol{w}_0, \boldsymbol{v}_0) = \sum_{i=1}^N s_i(\boldsymbol{w}_{0,i}, \boldsymbol{v}_{0,i}) \qquad \forall \boldsymbol{w}_0, \boldsymbol{v}_0 \in \boldsymbol{W}_0.$$

Finally, we define the coarse operator  $P_0: \mathbf{W} \longrightarrow \mathbf{W}_0$  by

$$\tilde{s}_0(P_0\boldsymbol{u},\boldsymbol{v}_0) = s(\boldsymbol{u},\boldsymbol{v}_0) \ \forall \boldsymbol{v}_0 \in \boldsymbol{W}_0,$$

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and the local operators  $P_i = R_i^T \tilde{P}_i : \mathbf{W} \longrightarrow R_i^T \mathbf{W}_i$  by

$$\tilde{s}_i(\tilde{P}_i \boldsymbol{u}, \boldsymbol{v}_i) = s(\boldsymbol{u}, R_i^T \boldsymbol{v}_i) \ \forall \boldsymbol{v}_i \in \boldsymbol{W}_i.$$

Then, our BDDC method is defined by the preconditioned operator

$$P = \sum_{i=0}^{N} P_i . \tag{5}$$

The matrix form of P and the associated preconditioner can be found in [5].

## 4 A Quasi-optimal BDDC Convergence Bound

We start by recalling the following assumption from [5], using the same notations.

**Assumption 1** Given any  $\Gamma_i$ , i = 1, 2, ..., N, let  $\mathcal{E}_i$  represent the set of the edges of  $\Gamma_i$ . Then, we assume that there exist two positive constants  $k_*, k^*$  and a boundary 120 seminorm  $|\cdot|_{\tau(\Gamma_i)}$  on  $\overline{\boldsymbol{W}}_i$ ,  $i=1,2,\ldots,N$ , such that 121

$$|\mathbf{w}_i|_{\tau(\Gamma_i)}^2 \le k^* s_i(\mathbf{w}_i, \mathbf{w}_i) \quad \forall \mathbf{w}_i \in \overline{\mathbf{W}}_i,$$
 (6)

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$$|\mathbf{w}_i|_{\tau(E)}^2 \ge k_* s_i(\mathbf{w}_i, \mathbf{w}_i) \quad \forall \mathbf{w}_i \in \mathbf{W}_i, \tag{7}$$

$$|\boldsymbol{w}_i|_{\tau(\Gamma_i)}^2 = \sum_{e \in \mathscr{E}_i} |\boldsymbol{w}_i|_{\tau(e)}^2 \quad \forall \boldsymbol{w}_i \in \overline{\boldsymbol{W}}_i,$$
 (8)

where  $|\cdot|_{\tau(e)}$  is a given seminorm on the edge e.

We notice that we cannot adopt the obvious choice  $|\mathbf{w}_i|_{\tau(F_i)} = s_i(\mathbf{w}_i, \mathbf{w}_i)$ , since it 123 can be shown that it does not satisfy (8), not even with a bound including a uniform 124 constant. We have the following main result. 125

**Theorem 2.** If Assumption 1 holds, then the condition number  $\kappa$  of the Reissner- 126 Mindlin BDDC preconditioned operator P in (5) satisfies the bound 127

$$\kappa(P) \le C \left( 1 + \log^3 \left( H/h \right) \right), \tag{128}$$

with the constant C depending only on the material constants and mesh regularity, 129 and not on the plate thickness t. 130

Here we can only outline the main steps of the proof; full details can be found 131 in [6]. The proof proceeds by showing that Assumption 1 holds for the MITC plate 132 bending problem (4) and by establishing the respective upper and lower bounds for 133 the constants  $k_*, k^*$  in (6), (7). These bounds in turn will prove Theorem 2 since 134  $\kappa(P) \le C(1 + 5k_*^{-1}k^*)$ , see [5, 21] for a proof. Upper bound (6). The upper bound is established exactly as in [5, Sect. 5.2]. 136

**Lower bound** (7). To prove the lower bound, we note that the local spaces  $\mathbf{W}_i$ , 137  $i = 1, 2, \dots, N$ , are composed of rotation and deflection parts, which we denote by 138  $\overline{W}_i = \overline{\Theta}_i \times \overline{U}_i$ . Accordingly, we denote the rotation and deflection parts of the constrained space by  $\mathbf{W}_i = \mathbf{\Theta}_i \times U_i$ , where the functions of  $\mathbf{\Theta}_i$  and  $U_i$  vanish at the 140 subdomain corner nodes. We work with the following seminorm defined in [5]: 141  $|\mathbf{w}_i|_{\tau(\Gamma_i)}^2 = \sum_{e \in \mathscr{E}_i} |\mathbf{w}_i|_{\tau(e)}^2 \quad \forall \mathbf{w}_i = (\boldsymbol{\theta}_i, u_i) \in \overline{\mathbf{W}}_i$ , where for all edges  $e \in \mathscr{E}_i$ 

$$|\mathbf{w}_i|_{\tau(e)}^2 = |\mathbf{\theta}_i|_{\gamma(e)}^2 + ht^{-2}||\Pi|\mathbf{\theta}_i \cdot \mathbf{\tau} - u_i'||_{L^2(e)}^2,$$
 143

$$|oldsymbol{ heta}_i|_{\gamma(e)} := \inf_{oldsymbol{\psi} \in [H^1(\Omega_i)]^2, oldsymbol{\psi}|_e = oldsymbol{ heta}_i|_e} ||oldsymbol{arepsilon}(oldsymbol{\psi})||_{L^2(\Omega_i)},$$

**7** is the tangent unit vector at the boundary and the apex indicates the derivative, 146 in the direction of  $\tau$ , for functions defined on the (one dimensional) boundary. We 147

now improve the lower bound proved in [5] by introducing a splitting of the plate 148 rotation variable. Consider  $\mathbf{w}_i = (\mathbf{\theta}_i, u_i) \in \mathbf{W}_i$  and define the splitting  $\mathbf{\theta}_i^{(2)} \in \mathbf{\Theta}_i^{(2)} := 148 \operatorname{span} \left\{ B_l^i \mathbf{\tau} \right\}_{l \in \Gamma}$ , by

$$\int_{e} \boldsymbol{\theta}_{i}^{(2)} \cdot \boldsymbol{\tau} = \int_{e} \boldsymbol{\theta}_{i} \cdot \boldsymbol{\tau} - u_{i}' \quad \forall e \in \mathscr{E}_{i},$$
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and let  $\boldsymbol{\theta}_i^{(1)} = \boldsymbol{\theta}_i - \boldsymbol{\theta}_i^{(2)}$  so that  $\boldsymbol{\theta}_i = \boldsymbol{\theta}_i^{(1)} + \boldsymbol{\theta}_i^{(2)}$ . By construction, it holds

$$\int_{e} u_{i}' - \boldsymbol{\theta}_{i}^{(1)} \cdot \boldsymbol{\tau} = 0 \quad \forall e \in \mathscr{E}_{i}.$$

We introduce also the related splitting of  $\mathbf{w}_i$ 

$$\mathbf{w}_i = \mathbf{w}_i^{(1)} + \mathbf{w}_i^{(2)}, \qquad \mathbf{w}_i^{(1)} = (u_i, \mathbf{\theta}_i^{(1)}), \qquad \mathbf{w}_i^{(2)} = (0, \mathbf{\theta}_i^{(2)}).$$
 154

An improved lower bound can be obtained by estimating the split terms in the following two lemmas; see [6] for complete proofs.

**Lemma 1.** There exists a constant C > 0 independent of h such that for all edges e 157 of all subdomains  $\Omega_i$  158

$$|\mathbf{w}_i|_{\tau(e)} = |(u_i, \mathbf{\theta}_i)|_{\tau(e)} \ge C(|(u_i, \mathbf{\theta}_i^{(1)})|_{\tau(e)} + |(0, \mathbf{\theta}_i^{(2)})|_{\tau(e)}).$$
 159

This lemma follows from the inequality  $||(0, \boldsymbol{\theta}_i^{(2)})||_{\tau(e)} \leq C||\boldsymbol{w}_i||_{\tau(e)}$ , that is derived 160 in [6] from the definition of  $\boldsymbol{\theta}_i^{(2)}$ , a scaling argument and an inverse inequality. A similar argument applied to the extension of  $\boldsymbol{\theta}_i^2$  by zero inside  $\Omega_i$  leads to the following 162 lemma.

**Lemma 2.** There exists a constant C > 0 independent of h such that

$$s_i(\boldsymbol{w}_i^{(2)}, \boldsymbol{w}_i^{(2)}) \le C |\boldsymbol{w}_i^{(2)}|_{\tau(F)}^2.$$
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The main step in the proof of Theorem 2 is the bound of the following proposition, 166 obtained by considering an auxiliary rotated Stokes problem with boundary data  $\boldsymbol{\theta}_i^{(1)}$  167 and several technical estimates, see [6, Proposition 5.5].

**Proposition 1.** There exists a constant C > 0 independent of h such that

$$s_i(\boldsymbol{w}_i^{(1)}, \boldsymbol{w}_i^{(1)}) \le C (1 + \log^3(H/h)) |\boldsymbol{w}_i^{(1)}|_{\tau(F)}^2.$$
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The upper bound then follows by combining the three previous results. Indeed, first 171 recalling the splitting  $\mathbf{w}_i = \mathbf{w}_i^{(1)} + \mathbf{w}_i^{(2)}$  and using a triangle inequality, then applying 172 Lemma 2 and Proposition 1, finally using Lemma 1 yields 173

$$s_i(\mathbf{w}_i, \mathbf{w}_i) \le 2\left(s_i(\mathbf{w}_i^{(1)}, \mathbf{w}_i^{(1)}) + s_i(\mathbf{w}_i^{(2)}, \mathbf{w}_i^{(2)})\right)$$
 174

$$\leq C\Big((1+\log^3{(H/h)})|\boldsymbol{w}_i^{(1)}|_{\tau(\Gamma_i)}^2+|\boldsymbol{w}_i^{(2)}|_{\tau(\Gamma_i)}^2\Big)\leq C(1+\log^3{(H/h)})|\boldsymbol{w}_i|_{\tau(\Gamma_i)}^2. \tag{176}$$

Bound (7) is therefore proved with  $k_*^{-1} = C(1 + \log^3(H/h))$ , with the constant C 177 depending only on the material constants and mesh regularity.

We remark that an extensive set of numerical tests, also including jump in the 179 coefficients, which are in complete accordance with Theorem 2, can be found in [5]. 180

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# **Penalty Robin-Robin Domain Decomposition Schemes** for Contact Problems of Nonlinear Elastic Bodies

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1 Introduction 8

Many domain decomposition techniques for contact problems have been proposed 9 on discrete level, particularly substructuring and FETI methods [1, 4].

Domain decomposition methods (DDMs), presented in [2, 10, 11, 16] for unilateral two-body contact problems of linear elasticity, are obtained on continuous level. 12 All of them require the solution of nonlinear one-sided contact problems for one or 13 both of the bodies in each iteration.

In works [6, 14, 15] we have proposed a class of penalty parallel Robin-Robin 15 domain decomposition schemes for unilateral multibody contact problems of linear 16 elasticity, which are based on penalty method and iterative methods for nonlinear 17 variational equations. In each iteration of these schemes we have to solve in a parallel 18 way some linear variational equations in subdomains.

In this contribution we generalize domain decomposition schemes, proposed in 20 [6, 14, 15] to the solution of unilateral and ideal contact problems of nonlinear elastic 21 bodies. We also present theorems about the convergence of these schemes.

## 2 Formulation of Multibody Contact Problem

Consider a contact problem of N nonlinear elastic bodies  $\Omega_{\alpha} \subset \mathbb{R}^3$  with sectionally smooth boundaries  $\Gamma_{\alpha}$ ,  $\alpha = 1, 2, ..., N$  (Fig. 1). Denote  $\Omega = \bigcup_{\alpha=1}^{N} \Omega_{\alpha}$ .

A stress-strain state in point  $\mathbf{x} = (x_1, x_2, x_3)^{\top}$  of each body  $\Omega_{\alpha}$  is defined by the 26 displacement vector  $\mathbf{u}_{\alpha} = u_{\alpha i} \mathbf{e}_i$ , the tensor of strains  $\hat{\boldsymbol{\varepsilon}}_{\alpha} = \varepsilon_{\alpha i j} \mathbf{e}_i \mathbf{e}_j$  and the tensor 27 of stresses  $\hat{\sigma}_{\alpha} = \sigma_{\alpha ij} \mathbf{e}_i \mathbf{e}_j$ . These quantities satisfy Cauchy relations, equilibrium 28 equations and nonlinear stress-strain law [8]: 29

$$\sigma_{\alpha ij} = \lambda_{\alpha} \, \delta_{ij} \, \Theta_{\alpha} + 2 \, \mu_{\alpha} \, \varepsilon_{\alpha ij} - 2 \, \mu_{\alpha} \, \omega_{\alpha}(e_{\alpha}) \, e_{\alpha ij}, \ i, j = 1, 2, 3 \,, \tag{1}$$

where  $\Theta_{\alpha}=arepsilon_{\alpha\,11}+arepsilon_{\alpha\,22}+arepsilon_{\alpha\,33}$  is the volume strain,  $\lambda_{\alpha}(\mathbf{x})>0,\ \mu_{\alpha}(\mathbf{x})>0$  are 30 bounded Lame parameters,  $e_{\alpha ij} = \varepsilon_{\alpha ij} - \delta_{ij} \Theta_{\alpha}/3$  are the components of the strain 31

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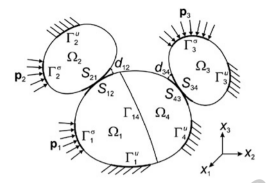


Fig. 1. Contact of several bodies

deviation tensor,  $e_{\alpha} = \sqrt{2g_{\alpha}}/3$  is the deformation intensity,  $g_{\alpha} = (\varepsilon_{\alpha 11} - \varepsilon_{\alpha 22})^2 + 32(\varepsilon_{\alpha 22} - \varepsilon_{\alpha 33})^2 + (\varepsilon_{\alpha 33} - \varepsilon_{\alpha 11})^2 + 6(\varepsilon_{\alpha 12}^2 + \varepsilon_{\alpha 23}^2 + \varepsilon_{\alpha 31}^2)$ , and  $\omega_{\alpha}(z)$  is nonlinear differentiable function, which satisfies the following properties:

$$0 \le \omega_{\alpha}(z) \le \partial \left(z \,\omega_{\alpha}(z)\right) / \partial z < 1, \ \partial \left(\omega_{\alpha}(z)\right) / \partial z \ge 0. \tag{2}$$

On the boundary  $\Gamma_{\alpha}$  let us introduce the local orthonormal basis  $\boldsymbol{\xi}_{\alpha}$ ,  $\boldsymbol{\eta}_{\alpha}$ ,  $\mathbf{n}_{\alpha}$ , 35 where  $\mathbf{n}_{\alpha}$  is the outer unit normal to  $\Gamma_{\alpha}$ . Then the vectors of displacements and 36 stresses on the boundary can be written in the following way:  $\mathbf{u}_{\alpha} = u_{\alpha\xi} \boldsymbol{\xi}_{\alpha} + 37 u_{\alpha\eta} \boldsymbol{\eta}_{\alpha} + u_{\alpha\eta} \mathbf{n}_{\alpha}$ ,  $\boldsymbol{\sigma}_{\alpha} = \hat{\boldsymbol{\sigma}}_{\alpha} \cdot \mathbf{n}_{\alpha} = \boldsymbol{\sigma}_{\alpha\xi} \boldsymbol{\xi}_{\alpha} + \sigma_{\alpha\eta} \boldsymbol{\eta}_{\alpha} + \sigma_{\alpha\eta} \mathbf{n}_{\alpha}$ .

 $u_{\alpha\eta} \, \eta_{\alpha} + u_{\alpha n} \, \mathbf{n}_{\alpha}, \, \sigma_{\alpha} = \hat{\boldsymbol{\sigma}}_{\alpha} \cdot \mathbf{n}_{\alpha} = \sigma_{\alpha\xi} \, \boldsymbol{\xi}_{\alpha} + \sigma_{\alpha\eta} \, \boldsymbol{\eta}_{\alpha} + \sigma_{\alpha n} \, \mathbf{n}_{\alpha}.$  38 Suppose that the boundary  $\Gamma_{\alpha}$  of each body consists of four disjoint parts:  $\Gamma_{\alpha} = 39$   $\Gamma_{\alpha}^{u} \cup \Gamma_{\alpha}^{\sigma} \cup \Gamma_{\alpha}^{l} \cup S_{\alpha}, \, \Gamma_{\alpha}^{u} \neq \emptyset, \, \Gamma_{\alpha}^{u} = \overline{\Gamma_{\alpha}^{u}}, \, \Gamma_{\alpha}^{l} \cup S_{\alpha} \neq \emptyset, \, \text{where } S_{\alpha} = \bigcup_{\beta \in B_{\alpha}} S_{\alpha\beta}, \, \text{and } 40$   $\Gamma_{\alpha}^{l} = \bigcup_{\beta' \in I_{\alpha}} \Gamma_{\alpha\beta'}. \, \text{Surface } S_{\alpha\beta} \, \text{is the possible unilateral contact area of body } \Omega_{\alpha} \, \text{with } 41$  body  $\Omega_{\beta}$ , and  $B_{\alpha} \subset \{1, 2, \dots, N\}$  is the set of the indices of all bodies in unilateral 42 contact with body  $\Omega_{\alpha}. \, \text{Surface } \Gamma_{\alpha\beta'} = \Gamma_{\beta'\alpha} \, \text{is the ideal contact area between bodies } 43$   $\Omega_{\alpha} \, \text{and } \Omega_{\beta'}, \, \text{and } I_{\alpha} \subset \{1, 2, \dots, N\} \, \text{is the set of the indices of all bodies which have } 44$  ideal contact with  $\Omega_{\alpha}.$ 

We assume that the areas  $S_{\alpha\beta}\subset \Gamma_{\alpha}$  and  $S_{\beta\alpha}\subset \Gamma_{\beta}$  are sufficiently close  $(S_{\alpha\beta}\approx 46\ S_{\beta\alpha})$ , and  $\mathbf{n}_{\alpha}(\mathbf{x})\approx -\mathbf{n}_{\beta}(\mathbf{x}')$ ,  $\mathbf{x}\in S_{\alpha\beta}$ ,  $\mathbf{x}'=P(\mathbf{x})\in S_{\beta\alpha}$ , where  $P(\mathbf{x})$  is the projection 47 of  $\mathbf{x}$  on  $S_{\alpha\beta}$  [12]. Let  $d_{\alpha\beta}(\mathbf{x})=\pm \|\mathbf{x}-\mathbf{x}'\|_2$  be a distance between bodies  $\Omega_{\alpha}$  and 48  $\Omega_{\beta}$  before the deformation. The sign of  $d_{\alpha\beta}$  depends on a statement of the problem. 49

We consider homogenous Dirichlet boundary conditions on the part  $\Gamma_{\alpha}^{u}$ , and Neumann boundary conditions on the part  $\Gamma_{\alpha}^{\sigma}$ :

$$\mathbf{u}_{\alpha}(\mathbf{x}) = 0, \ \mathbf{x} \in \Gamma_{\alpha}^{u}; \ \mathbf{\sigma}_{\alpha}(\mathbf{x}) = \mathbf{p}_{\alpha}(\mathbf{x}), \ \mathbf{x} \in \Gamma_{\alpha}^{\sigma}.$$
 (3)

On the possible contact areas  $S_{\alpha\beta}$ ,  $\beta \in B_{\alpha}$ ,  $\alpha = 1, 2, ..., N$  the following nonlinear unilateral contact conditions hold:

$$\sigma_{\alpha n}(\mathbf{x}) = \sigma_{\beta n}(\mathbf{x}') \le 0, \ \sigma_{\alpha \xi}(\mathbf{x}) = \sigma_{\beta \xi}(\mathbf{x}') = \sigma_{\alpha \eta}(\mathbf{x}) = \sigma_{\beta \eta}(\mathbf{x}') = 0, \quad (4)$$

$$u_{\alpha n}(\mathbf{x}) + u_{\beta n}(\mathbf{x}') \le d_{\alpha \beta}(\mathbf{x}),$$
 (5)

$$\left(u_{\alpha n}(\mathbf{x}) + u_{\beta n}(\mathbf{x}') - d_{\alpha \beta}(\mathbf{x})\right) \sigma_{\alpha n}(\mathbf{x}) = 0, \ \mathbf{x} \in S_{\alpha \beta}, \ \mathbf{x}' = P(\mathbf{x}) \in S_{\beta \alpha}.$$
 (6)

On ideal contact areas  $\Gamma_{\alpha\beta'} = \Gamma_{\beta'\alpha}$ ,  $\beta' \in I_{\alpha}$ ,  $\alpha = 1, 2, ..., N$  we consider ideal 56 mechanical contact conditions:

$$\mathbf{u}_{\alpha}(\mathbf{x}) = \mathbf{u}_{\beta'}(\mathbf{x}), \ \boldsymbol{\sigma}_{\alpha}(\mathbf{x}) = -\boldsymbol{\sigma}_{\beta'}(\mathbf{x}), \ \mathbf{x} \in \Gamma_{\alpha\beta'}.$$
 (7)

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### 3 Penalty Variational Formulation of the Problem

For each body  $\Omega_{\alpha}$  consider Sobolev space  $V_{\alpha} = [H^1(\Omega_{\alpha})]^3$  and the closed subspace 59  $V_{\alpha}^0 = \{\mathbf{u}_{\alpha} \in V_{\alpha} : \mathbf{u}_{\alpha} = 0 \text{ on } \Gamma_{\alpha}^u\}$ . All values of the elements  $\mathbf{u}_{\alpha} \in V_{\alpha}$ ,  $\mathbf{u}_{\alpha} \in V_{\alpha}^0$  on 60 the parts of boundary  $\Gamma_{\alpha}$  should be understood as traces [9].

the parts of boundary  $\Gamma_{\alpha}$  should be understood as traces [9].

Define Hilbert space  $V_0 = V_1^0 \times \ldots \times V_N^0$  with the scalar product  $(\mathbf{u}, \mathbf{v})_{V_0} = 62$   $\sum_{\alpha=1}^N (\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha})_{V_{\alpha}} \text{ and norm } \|\mathbf{u}\|_{V_0} = \sqrt{(\mathbf{u}, \mathbf{u})_{V_0}}, \, \mathbf{u}, \mathbf{v} \in V_0. \text{ Introduce the closed consex set of all displacements in } V_0, \text{ which satisfy nonpenentration contact conditions}$ 

$$K = \left\{ \mathbf{u} \in V_0 : \ u_{\alpha n} + u_{\beta n} \le d_{\alpha \beta} \text{ on } S_{\alpha \beta}, \ \mathbf{u}_{\alpha'} = \mathbf{u}_{\beta'} \text{ on } \Gamma_{\alpha' \beta'} \right\}, \tag{8}$$

where 
$$\{\alpha,\beta\} \in Q$$
,  $Q = \{\{\alpha,\beta\} : \alpha \in \{1,2,\ldots,N\}, \beta \in B_{\alpha}\}, \{\alpha',\beta'\} \in Q^I, Q^I = \{\{\alpha',\beta'\} : \alpha' \in \{1,2,\ldots,N\}, \beta' \in I_{\alpha}\}, \text{ and } d_{\alpha\beta} \in H_{00}^{1/2}(\Xi_{\alpha}), \Xi_{\alpha} = \operatorname{int}(\Gamma_{\alpha} \setminus \Gamma_{\alpha}^{u}).$ 

Let us introduce bilinear form  $A(\mathbf{u}, \mathbf{v}) = \sum_{\alpha=1}^{N} a_{\alpha}(\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha}), \mathbf{u}, \mathbf{v} \in V_0$ , which represents the total elastic deformation energy of the system of bodies, linear form 69  $L(\mathbf{v}) = \sum_{\alpha=1}^{N} l_{\alpha}(\mathbf{v}_{\alpha}), \mathbf{v} \in V_0$ , which is equal to the external forces work, and non-70 quadratic functional  $H(\mathbf{v}) = \sum_{\alpha=1}^{N} h_{\alpha}(\mathbf{v}_{\alpha}), \mathbf{v} \in V_0$ , which represents the total nonlinear deformation energy:

$$a_{\alpha}(\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha}) = \int_{\Omega_{\alpha}} \left[ \lambda_{\alpha} \Theta_{\alpha}(\mathbf{u}_{\alpha}) \Theta_{\alpha}(\mathbf{v}_{\alpha}) + 2 \mu_{\alpha} \sum_{i,j} \varepsilon_{\alpha i j}(\mathbf{u}_{\alpha}) \varepsilon_{\alpha i j}(\mathbf{v}_{\alpha}) \right] d\Omega, \quad (9)$$

$$l_{\alpha}(\mathbf{v}_{\alpha}) = \int_{\Omega_{\alpha}} \mathbf{f}_{\alpha} \cdot \mathbf{v}_{\alpha} \, d\Omega + \int_{\Gamma_{\alpha}^{\sigma}} \mathbf{p}_{\alpha} \cdot \mathbf{v}_{\alpha} \, dS \,, \tag{10}$$

$$h_{\alpha}(\mathbf{v}_{\alpha}) = 3 \int_{\Omega_{\alpha}} \mu_{\alpha} \int_{0}^{e_{\alpha}(\mathbf{v}_{\alpha})} z \, \omega_{\alpha}(z) \, dz \, d\Omega \,, \tag{11}$$

where  $\mathbf{p}_{\alpha} \in [H_{00}^{-1/2}(\Xi_{\alpha})]^3$ , and  $\mathbf{f}_{\alpha} \in [L_2(\Omega_{\alpha})]^3$  is the vector of volume forces.

Using [12], we have shown that the original contact problem has an alternative 76 weak formulation as the following minimization problem on the set K: 77

$$F(\mathbf{u}) = A(\mathbf{u}, \mathbf{u})/2 - H(\mathbf{u}) - L(\mathbf{u}) \to \min_{\mathbf{u} \in K}.$$
 (12)

Bilinear form A is symmetric, continuous with constant  $M_A > 0$  and coercive 78 with constant  $B_A > 0$ , and linear form L is continuous. Nonquadratic functional H is 79 doubly Gateaux differentiable in  $V_0$ :

$$H'(\mathbf{u}, \mathbf{v}) = \sum_{\alpha} h'_{\alpha}(\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha}), \ H''(\mathbf{u}, \mathbf{v}, \mathbf{w}) = \sum_{\alpha} h''_{\alpha}(\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha}, \mathbf{w}_{\alpha}), \ \mathbf{u}, \mathbf{v}, \mathbf{w} \in V_0, \quad (13)$$

$$h'_{\alpha}(\mathbf{u}_{\alpha}, \mathbf{v}_{\alpha}) = 2 \int_{\Omega_{\alpha}} \mu_{\alpha} \, \omega_{\alpha}(e_{\alpha}(\mathbf{u}_{\alpha})) \sum_{i,j} e_{\alpha ij}(\mathbf{u}_{\alpha}) \, e_{\alpha ij}(\mathbf{v}_{\alpha}) \, d\Omega. \tag{14}$$

Moreover, we have proved that the following conditions hold:

$$(\exists C > 0) (\forall \mathbf{u} \in V_0) \{ (1 - C) A(\mathbf{u}, \mathbf{u}) \ge 2H(\mathbf{u}) \}, \tag{15}$$

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$$(\forall \mathbf{u} \in V_0) (\exists R > 0) (\forall \mathbf{v} \in V_0) \left\{ \left| H'(\mathbf{u}, \mathbf{v}) \right| \le R \|\mathbf{v}\|_{V_0} \right\}, \tag{16}$$

$$(\exists D > 0) (\forall \mathbf{u}, \mathbf{v}, \mathbf{w} \in V_0) \left\{ \left| H''(\mathbf{u}, \mathbf{v}, \mathbf{w}) \right| \le D \|\mathbf{v}\|_{V_0} \|\mathbf{w}\|_{V_0} \right\}, \tag{17}$$

$$(\exists B > 0) (\forall \mathbf{u}, \mathbf{v} \in V_0) \left\{ A(\mathbf{v}, \mathbf{v}) - H''(\mathbf{u}, \mathbf{v}, \mathbf{v}) \ge B \|\mathbf{v}\|_{V_0}^2 \right\}. \tag{18}$$

From these properties, it follows that there exists a unique solution  $\bar{\mathbf{u}} \in K$  of minimization problem (12), and this problem is equivalent to the following variational 87 inequality, which is nonlinear in  $\mathbf{u}$ :

$$A(\mathbf{u}, \mathbf{v} - \mathbf{u}) - H'(\mathbf{u}, \mathbf{v} - \mathbf{u}) - L(\mathbf{v} - \mathbf{u}) \ge 0, \ \forall \mathbf{v} \in K, \ \mathbf{u} \in K.$$

To obtain a minimization problem in the whole space  $V_0$ , we apply a penalty method [3, 7, 9, 13] to problem (12). We use a penalty in the form

$$J_{\theta}(\mathbf{u}) = \frac{1}{2\theta} \sum_{\{\alpha,\beta\} \in Q} \left\| \left( d_{\alpha\beta} - u_{\alpha n} - u_{\beta n} \right)^{-} \right\|_{L_{2}(S_{\alpha\beta})}^{2} + \frac{1}{2\theta} \sum_{\{\alpha',\beta'\} \in Q'} \left\| \mathbf{u}_{\alpha'} - \mathbf{u}_{\beta'} \right\|_{[L_{2}(\Gamma_{\alpha'\beta'})]^{3}}^{2},$$
(20)

where  $\theta > 0$  is a penalty parameter, and  $y^- = \min\{0, y\}$ .

Now, consider the following unconstrained minimization problem in  $V_0$ :

$$F_{\theta}(\mathbf{u}) = A(\mathbf{u}, \mathbf{u})/2 - H(\mathbf{u}) - L(\mathbf{u}) + J_{\theta}(\mathbf{u}) \to \min_{\mathbf{u} \in V_0}.$$
 (21)

The penalty term  $J_{\theta}$  is nonnegative and Gateaux differentiable in  $V_0$ , and its differential  $J'_{\theta}(\mathbf{u}, \mathbf{v}) = -\frac{1}{\theta} \sum_{\{\alpha, \beta\} \in \mathcal{Q}} \int_{S_{\alpha\beta}} \left( d_{\alpha\beta} - u_{\alpha n} - u_{\beta n} \right)^{-} \left( v_{\alpha n} + v_{\beta n} \right) dS + \frac{1}{\theta} \sum_{\{\alpha', \beta'\} \in \mathcal{Q}^{I}} \int_{\Gamma_{\alpha'\beta'}} \left( \mathbf{u}_{\alpha'} - \mathbf{u}_{\beta'} \right) \cdot \left( \mathbf{v}_{\alpha'} - \mathbf{v}_{\beta'} \right) dS$  satisfy the following properties [15]: 95

$$(\forall \mathbf{u} \in V_0)(\exists \tilde{R} > 0)(\forall \mathbf{v} \in V_0) \left\{ \left| J'_{\theta}(\mathbf{u}, \mathbf{v}) \right| \le \tilde{R} \left\| \mathbf{v} \right\|_{V_0} \right\}, \tag{22}$$

$$(\exists \tilde{D} > 0)(\forall \mathbf{u}, \mathbf{v}, \mathbf{w} \in V_0) \left\{ \left| J_{\theta}'(\mathbf{u} + \mathbf{w}, \mathbf{v}) - J_{\theta}'(\mathbf{u}, \mathbf{v}) \right| \le \tilde{D} \|\mathbf{v}\|_{V_0} \|\mathbf{w}\|_{V_0} \right\}, \tag{23}$$

$$(\forall \mathbf{u}, \mathbf{v} \in V_0) \left\{ J'_{\theta} \left( \mathbf{u} + \mathbf{v}, \mathbf{v} \right) - J'_{\theta} \left( \mathbf{u}, \mathbf{v} \right) \ge 0 \right\}. \tag{24}$$

Using these properties and the results in [3], we have shown that problem (21) 98 has a unique solution  $\bar{\mathbf{u}}_{\theta} \in V_0$  and is equivalent to the following nonlinear variational 99 equation in the space  $V_0$ :

$$F'_{\theta}(\mathbf{u}, \mathbf{v}) = A(\mathbf{u}, \mathbf{v}) - H'(\mathbf{u}, \mathbf{v}) + J'_{\theta}(\mathbf{u}, \mathbf{v}) - L(\mathbf{v}) = 0, \ \forall \mathbf{v} \in V_0, \ \mathbf{u} \in V_0.$$
 (25)

Using the results of works [7, 13], we have proved that  $\|\bar{\mathbf{u}}_{\theta} - \bar{\mathbf{u}}\|_{V_0} \to 0$ .

### 4 Iterative Methods for Nonlinear Variational Equations

In arbitrary reflexive Banach space  $V_0$  consider an abstract nonlinear variational 103 equation 104

$$\Phi(\mathbf{u}, \mathbf{v}) = L(\mathbf{v}), \ \forall \mathbf{v} \in V_0, \ \mathbf{u} \in V_0$$
(26)

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where  $\Phi: V_0 \times V_0 \to \mathbb{R}$  is a functional, which is linear in  $\mathbf{v}$ , but nonlinear in  $\mathbf{u}$ , 105 and L is linear continuous form. Suppose that this variational equation has a unique 106 solution  $\bar{\mathbf{u}}_* \in V_0$ .

For the numerical solution of (26) we use the next iterative method [5, 6, 15]:

$$G(\mathbf{u}^{k+1}, \mathbf{v}) = G(\mathbf{u}^{k}, \mathbf{v}) - \gamma \left[ \Phi(\mathbf{u}^{k}, \mathbf{v}) - L(\mathbf{v}) \right], \ \forall \mathbf{v} \in V_{0}, \ k = 0, 1, \dots,$$
 (27)

where G is some given bilinear form in  $V_0 \times V_0$ ,  $\gamma \in \mathbb{R}$  is fixed parameter, and  $\mathbf{u}^k \in V_0$  109 is the k-th approximation to the exact solution of problem (26).

We have proved the next theorem [5, 15] about the convergence of this method.

**Theorem 1.** Suppose that the following conditions hold

$$(\forall \mathbf{u} \in V_0) (\exists R_{\Phi} > 0) (\forall \mathbf{v} \in V_0) \left\{ |\Phi(\mathbf{u}, \mathbf{v})| \le R_{\Phi} \|\mathbf{v}\|_{V_0} \right\}, \tag{28}$$

$$(\exists D_{\Phi} > 0)(\forall \mathbf{u}, \mathbf{v}, \mathbf{w} \in V_0) \left\{ |\Phi(\mathbf{u} + \mathbf{w}, \mathbf{v}) - \Phi(\mathbf{u}, \mathbf{v})| \le D_{\Phi} ||\mathbf{v}||_{V_0} ||\mathbf{w}||_{V_0} \right\}, \quad (29)$$

$$(\exists B_{\Phi} > 0) (\forall \mathbf{u}, \mathbf{v} \in V_0) \left\{ \Phi(\mathbf{u} + \mathbf{v}, \mathbf{v}) - \Phi(\mathbf{u}, \mathbf{v}) \ge B_{\Phi} \|\mathbf{v}\|_{V_0}^2 \right\}, \tag{30}$$

bilinear form G is symmetric, continuous with constant  $M_G>0$  and coercive with 115 constant  $B_G>0$ , and  $\gamma\in(0;2\gamma^*)$ ,  $\gamma^*=B_{\Phi}B_G/D_{\Phi}^2$ .

Then  $\|\mathbf{u}^k - \bar{\mathbf{u}}_*\|_{V_0} \to 0$ , where  $\{\mathbf{u}^k\} \subset V_0$  is obtained by method (27). Moreover,

the convergence rate in norm  $\|\cdot\|_G = \sqrt{G(\cdot,\cdot)}$  is linear, and the highest convergence 118 rate in this norm reaches as  $\gamma = \gamma^*$ .

In addition, we have proposed nonstationary iterative method to solve (26), where bilinear form G and parameter  $\gamma$  are different in each iteration:

$$G^{k}(\mathbf{u}^{k+1}, \mathbf{v}) = G^{k}(\mathbf{u}^{k}, \mathbf{v}) - \gamma^{k} \left[ \Phi(\mathbf{u}^{k}, \mathbf{v}) - L(\mathbf{v}) \right], \ \forall \mathbf{v} \in V_{0}, \ k = 0, 1, \dots$$
 (31)

A convergence theorem for this method is proved in [15].

# 5 Domain Decomposition Schemes for Contact Problems

Now let us apply iterative methods (27) and (31) to the solution of nonlinear penalty variational equation (25) of multibody contact problem. This penalty equation can be written in form (26), where

$$\Phi(\mathbf{u}, \mathbf{v}) = A(\mathbf{u}, \mathbf{v}) - H'(\mathbf{u}, \mathbf{v}) + J'_{\theta}(\mathbf{u}, \mathbf{v}), \ \mathbf{u}, \mathbf{v} \in V_0.$$
(32)

We consider such variants of methods (27) and (31), which lead to the domain 127 decomposition.

Let us take the bilinear form G in iterative method (27) as follows [6, 15]: 129

$$G(\mathbf{u}, \mathbf{v}) = A(\mathbf{u}, \mathbf{v}) + X(\mathbf{u}, \mathbf{v}), \ \mathbf{u}, \mathbf{v} \in V_0,$$
(33)

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$$X(\mathbf{u}, \mathbf{v}) = \frac{1}{\theta} \sum_{\alpha=1}^{N} \left[ \sum_{\beta \in B_{\alpha}} \int_{S_{\alpha\beta}} u_{\alpha n} v_{\alpha n} \, \psi_{\alpha\beta} \, dS + \sum_{\beta' \in I_{\alpha}} \int_{\Gamma_{\alpha\beta'}} \mathbf{u}_{\alpha} \cdot \mathbf{v}_{\alpha} \, \phi_{\alpha\beta'} \, dS \right], \tag{130}$$

where  $\psi_{\alpha\beta}(\mathbf{x}) = \{1, \mathbf{x} \in S^1_{\alpha\beta}\} \vee \{0, \mathbf{x} \in S_{\alpha\beta} \setminus S^1_{\alpha\beta}\}$  and  $\phi_{\alpha\beta'}(\mathbf{x}) = \{1, \mathbf{x} \in \Gamma^1_{\alpha\beta'}\} \vee \{0, \mathbf{x} \in S^1_{\alpha\beta}\}$  $\{0, \mathbf{x} \in \Gamma_{\alpha\beta'} \setminus \Gamma_{\alpha\beta'}^1\}$  are characteristic functions of arbitrary subsets  $S_{\alpha\beta}^1 \subseteq S_{\alpha\beta}$ ,  $\Gamma^1_{\alpha\beta'}\subseteq\Gamma_{\alpha\beta'}$  of possible unilateral and ideal contact areas respectively 134

Introduce a notation  $\tilde{\mathbf{u}}^{k+1} = [\mathbf{u}^{k+1} - \mathbf{u}^k]/\gamma + \mathbf{u}^k \in V_0$ . Then iterative method (27) with bilinear form (33) can be written in such way: 136

$$A\left(\tilde{\mathbf{u}}^{k+1},\mathbf{v}\right) + X\left(\tilde{\mathbf{u}}^{k+1},\mathbf{v}\right) = L(\mathbf{v}) + X\left(\mathbf{u}^{k},\mathbf{v}\right) + H'(\mathbf{u}^{k},\mathbf{v}) - J'_{\theta}(\mathbf{u}^{k},\mathbf{v}), \quad (34)$$

$$\mathbf{u}^{k+1} = \gamma \,\tilde{\mathbf{u}}^{k+1} + (1 - \gamma)\mathbf{u}^k, \ k = 0, 1, \dots$$
 (35)

Bilinear form X is symmetric, continuous with constant  $M_X > 0$ , and nonnegative 138 [15]. Due to these properties, and due to the properties of bilinear form A, it follows that the conditions of Theorem 1 hold. Therefore, we obtain the next proposition:

**Theorem 2.** The sequence  $\{\mathbf{u}^k\}$  of the method (34)–(35) converges strongly to the 141 solution of penalty variational equation (25) for  $\gamma \in (0; 2B_{\Phi}B_G/D_{\Phi}^2)$ , where  $B_G =$  $B_A$ ,  $B_{\Phi} = B$ ,  $D_{\Phi} = M_A + D + \tilde{D}$ . The convergence rate in norm  $\|\cdot\|_G$  is linear. 143

As the common quantities of the subdomains are known from the previous iteration, variational equation (34) splits into N separate equations for each subdomain 145  $\Omega_{\alpha}$ , and method (34)–(35) can be written in the following equivalent form: 146

$$a_{\alpha}(\tilde{\mathbf{u}}_{\alpha}^{k+1}, \mathbf{v}_{\alpha}) + \sum_{\beta \in B_{\alpha}} \int_{S_{\alpha\beta}} \frac{\psi_{\alpha\beta}}{\theta} \, \tilde{u}_{\alpha n}^{k+1} \, v_{\alpha n} \, dS + \sum_{\beta' \in I_{\alpha}} \int_{\Gamma_{\alpha\beta'}} \frac{\phi_{\alpha\beta'}}{\theta} \, \tilde{\mathbf{u}}_{\alpha}^{k+1} \cdot \mathbf{v}_{\alpha} \, dS$$
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$$= l_{\alpha}(\mathbf{v}_{\alpha}) + \frac{1}{\theta} \sum_{\beta \in B_{\alpha}} \int_{S_{\alpha\beta}} \left[ \psi_{\alpha\beta} \, u_{\alpha n}^{k} + \left( d_{\alpha\beta} - u_{\alpha n}^{k} - u_{\beta n}^{k} \right)^{-} \right] v_{\alpha n} \, dS$$
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$$+\frac{1}{\theta} \sum_{\beta' \in I_{\alpha}} \int_{\Gamma_{\alpha\beta'}} \left[ \phi_{\alpha\beta'} \mathbf{u}_{\alpha}^{k} + \left( \mathbf{u}_{\beta'}^{k} - \mathbf{u}_{\alpha}^{k} \right) \right] \cdot \mathbf{v}_{\alpha} \, dS + h_{\alpha}' (\mathbf{u}_{\alpha}^{k}, \mathbf{v}_{\alpha}), \quad \forall \mathbf{v}_{\alpha} \in V_{\alpha}^{0}, \quad (36)$$

$$\mathbf{u}_{\alpha}^{k+1} = \gamma \, \tilde{\mathbf{u}}_{\alpha}^{k+1} + (1 - \gamma) \, \mathbf{u}_{\alpha}^{k}, \ \alpha = 1, 2, \dots, N, \ k = 0, 1, \dots$$
 (37)

In each iteration k of method (36)–(37), we have to solve N linear variational 152 equations in parallel, which correspond to some linear elasticity problems in subdomains with additional volume forces in  $\Omega_{\alpha}$ , and with Robin boundary conditions 154 on contact areas. Therefore, this method refers to parallel Robin–Robin type domain 155 decomposition schemes. 156

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Taking different characteristic functions  $\psi_{\alpha\beta}$  and  $\phi_{\alpha'\beta'}$ , we can obtain different 157 particular cases of penalty domain decomposition method (36)–(37).

Thus, taking  $\psi_{\alpha\beta}(\mathbf{x}) \equiv 0$ ,  $\beta \in B_{\alpha}$ ,  $\phi_{\alpha\beta'}(\mathbf{x}) \equiv 0$ ,  $\beta' \in I_{\alpha}$ ,  $\alpha = 1, 2, ..., N$ , we get 159 parallel Neumann-Neumann domain decomposition scheme.

Other borderline case is when 
$$\psi_{\alpha\beta}(\mathbf{x}) \equiv 1$$
,  $\beta \in B_{\alpha}$ ,  $\phi_{\alpha\beta'}(\mathbf{x}) \equiv 1$ ,  $\beta' \in I_{\alpha}$ ,  $\alpha = 161$  1,2,..., $N$ , i.e.  $S^1_{\alpha\beta} = S_{\alpha\beta}$ ,  $\Gamma^1_{\alpha\beta'} = \Gamma_{\alpha\beta'}$ .

Moreover, we can choose functions  $\psi_{\alpha\beta}$  and  $\phi_{\alpha\beta'}$  differently in each iteration k. 163 Then we obtain nonstationary domain decomposition schemes, which are equivalent to iterative method (31) with bilinear forms

$$G^{k}(\mathbf{u}, \mathbf{v}) = A(\mathbf{u}, \mathbf{v}) + X^{k}(\mathbf{u}, \mathbf{v}), \ \mathbf{u}, \mathbf{v} \in V_{0}, \ k = 0, 1, ...,$$
 (38)

$$X^{k}(\mathbf{u}, \mathbf{v}) = \frac{1}{\theta} \sum_{\alpha=1}^{N} \left[ \sum_{\beta \in B_{\alpha}} \int_{S_{\alpha\beta}} u_{\alpha n} v_{\alpha n} \, \psi_{\alpha\beta}^{k} \, dS + \sum_{\beta' \in I_{\alpha}} \int_{\Gamma_{\alpha\beta'}} \mathbf{u}_{\alpha} \cdot \mathbf{v}_{\alpha} \, \phi_{\alpha\beta'}^{k} \, dS \right]. \tag{167}$$

If we take characteristic functions  $\psi_{\alpha\beta}^k$  and  $\phi_{\alpha\beta'}^k$  as follows [6, 14, 15]:

$$\psi_{\alpha\beta}^{k}(\mathbf{x}) = \chi_{\alpha\beta}^{k}(\mathbf{x}) = \begin{cases} 0, d_{\alpha\beta}(\mathbf{x}) - u_{\alpha n}^{k}(\mathbf{x}) - u_{\beta n}^{k}(\mathbf{x}') \ge 0 \\ 1, d_{\alpha\beta}(\mathbf{x}) - u_{\alpha n}^{k}(\mathbf{x}) - u_{\beta n}^{k}(\mathbf{x}') < 0 \end{cases}, \ \mathbf{x}' = P(\mathbf{x}), \ \mathbf{x} \in S_{\alpha\beta}, \quad \text{169}$$

$$\phi_{\alpha\beta'}^{k}(\mathbf{x}) \equiv 1, \ \mathbf{x} \in \Gamma_{\alpha\beta'}, \ \beta \in B_{\alpha}, \ \beta' \in I_{\alpha}, \ \alpha = 1, 2, \dots, N,$$

then we shall get the method, which can be conventionally named as nonstationary 172 parallel Dirichlet–Dirichlet domain decomposition scheme. 173

In addition to methods (27), (33) and (31), (38), we have proposed another family 174 of DDMs for the solution of (25), where the second derivative of functional  $H(\mathbf{u})$  is used. These domain decomposition methods are obtained from (31), if we choose 176 bilinear forms  $G^k(\mathbf{u}, \mathbf{v})$  as follows

$$G^{k}(\mathbf{u}, \mathbf{v}) = A(\mathbf{u}, \mathbf{v}) - H''(\mathbf{u}^{k}, \mathbf{u}, \mathbf{v}) + X^{k}(\mathbf{u}, \mathbf{v}), \ \mathbf{u}, \mathbf{v} \in V_{0}, \ k = 0, 1, \dots$$
 (39)

Numerical analysis of presented penalty Robin-Robin DDMs has been made 178 for plane unilateral two-body and three-body contact problems of linear elasticity  $(\omega_{\alpha} \equiv 0)$  using finite element approximations [6, 14, 15]. Numerical experiments 180 have confirmed the theoretical results about the convergence of these methods.

Among the positive features of proposed domain decomposition schemes are 182 the simplicity of the algorithms and the regularization of original contact problem 183 because of the use of penalty method. These domain decomposition schemes have only one iteration loop, which deals with domain decomposition, nonlinearity of the stress-strain relationship, and nonlinearity of unilateral contact conditions.

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<b>Domain Decomposition</b>	Method	for	<b>Stokes</b>	Problen	r
with Tresca Friction					4

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1 Introduction 12

Development of numerical methods for the solution of Stokes system with slip 13 boundary conditions (Tresca friction conditions) is a challenging task whose difficulty lies in the nonlinear conditions. Such boundary conditions have to be taken 15 into account in many situations arising in practice, in flow of polymers (see [10] and 16 references therein).

The paper is devoted to domain decomposition methods (DDM in short) for the 18 Stokes problem with the slip boundary conditions. The original domain is cut into 19 two sub-domains and the augmented Lagrangian formulation for separate resulting 20 Poisson problems in both domains is used for computations. To relate solutions of 21 these two sub-problems to the original solution, one has to introduce additional con- 22 straints "gluing" them together. The domain decomposition formulation is based on 23 the Uzawa block relaxation method for the augmented Lagrangian involving three 24 supplementary conditions. The paper is concluded by preliminary several numerical 25 examples.

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## 2 Setting Stokes Problem with Nonlinear Boundary Conditions

Let us consider a domain  $\Omega \subset \mathbb{R}^2$  with the Lipschitz boundary  $\partial \Omega$  which is split into 28 two non-empty and non-overlapping parts  $\Gamma_0$  and  $\Gamma$ . We denote by n the outward 29 unit normal to  $\partial \Omega$  and  $u_n$ , respectively  $u_t$ , the normal, respectively the tangential, 30 component of u. We also make use of  $\sigma_t$  for the tangential component of the stress 31 vector  $\sigma(u)n$ . The problem consists in finding the velocity field u and the pressure 32 p for the following Stokes problem with nonlinear boundary condition of Tresca 33 friction type: 3/

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$$\begin{cases}
-div(v\varepsilon(u)) + \nabla p = f & \text{in } \Omega \\
div(u) = 0 & \text{in } \Omega \\
u = 0 & \text{on } \Gamma_0 \\
u_n = 0 & \text{on } \Gamma
\end{cases}$$

$$|\sigma_t| \le g & \text{on } \partial \Omega$$

$$|\sigma_t| < g \Rightarrow u_t = 0 & \text{on } \Gamma$$

$$|\sigma_t| = g \Rightarrow \exists k > 0 \text{ a constant such that} \quad u_t = -k\sigma_t \text{ on } \Gamma$$

where f is in  $L^2(\Omega)$ ,  $g \in L^2(\Gamma)$ , g > 0 is the given slip bound on  $\Gamma$  and  $|\cdot|$  is the 35 euclidean norm. 36

One can derive the variational formulation of (1):

$$\begin{cases} \text{Find } u \in \mathbf{V}_{div}(\Omega) \text{ such that } : \forall v \in \mathbf{V}_{div}(\Omega) \\ a(u, v - u) + j(v) - j(u) \ge L(v - u), \end{cases}$$
 (2)

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with

$$\mathbf{V}(\Omega)=\Big\{v\in\mathbf{H}^1(\Omega),\, v_{|\Gamma_0}=0, v_n=0 \ \mathrm{on} \ \Gamma\Big\},$$
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$$\mathbf{V}_{div}(\Omega) = \left\{ v \in \mathbf{V}(\Omega) , div(v) = 0 \text{ in } \Omega \right\},$$

$$a(u,v) = \int_{\Omega} v \varepsilon(u) : \varepsilon(v) d\Omega, \quad L(v) = \int_{\Omega} f v d\Omega, \quad j(v) = \int_{\Gamma} g|v_t| d\Gamma.$$

Problem (2) is an elliptic variational inequality of the second kind which has a unique 44 solution [3]. Moreover, since the bilinear form  $a(\cdot, \cdot)$  is symmetric (2) is equivalent 45 to the following constrained non-differentiable minimization problem:

Find 
$$u \in \mathbf{V}_{div}(\Omega)$$
 such that :  $\mathcal{J}(u) \le \mathcal{J}(v) \quad \forall v \in \mathbf{V}_{div}(\Omega)$ , (3)

where  $\mathcal{J}(v) = \frac{1}{2}a(v,v) + j(v) - L(v)$  is the total potential energy functional. 47

### 3 Uzawa DDM for Stokes Problem with Tresca Friction

We now study the domain decomposition of (3). We first rewrite (3) in the following 49 more useful form. Suppose that  $\varphi = v_t$ , then the minimization problem (3) becomes: 50

$$\begin{cases} \text{Find } (u, \Phi) \in \Pi \text{ such that:} \\ \Sigma(u, \Phi) \leq \Sigma(v, \varphi) \, \forall \, (v, \varphi) \in \Pi, \end{cases} \tag{4}$$

where

$$\Pi = \{(v, \varphi) \in \mathbf{V}_{div}(\Omega) \times H^{\frac{1}{2}}(\Gamma) \text{ such that } \varphi = v_t\},$$
 52

and  $\Sigma$  is the Lagrangian defined on  $\Pi$  by:

$$\forall (\varphi, v) \in \Pi \qquad \Sigma(v, \varphi) = \frac{1}{2}a(v, v) - L(v) + j(\varphi). \tag{5}$$

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Let  $\{\Omega_1, \Omega_2\}$  be a partition of  $\Omega$ , as shown in Fig. 1, and let

$$\begin{split} &\Gamma_{12} = \Gamma_{21} = \partial \Omega_{1} \cap \partial \Omega_{2}, \quad \Gamma_{i} = \Gamma \cup \partial \Omega_{i}, \quad \Gamma_{i}^{0} = \Gamma_{0} \cup \partial \Omega_{i}, \\ &v_{i} = v|_{\Omega_{i}}, \quad p_{i} = p|_{\Omega_{i}}, \\ &\mathbf{V}(\Omega_{i}) = \Big\{v_{i} \in \mathbf{H}^{1}(\Omega_{i}), \ v_{i|\Gamma_{i}^{0}} = 0, v_{i}.n_{i|\Gamma_{i}} = 0\Big\}, \\ &\mathbf{V}_{div}(\Omega_{i}) = \Big\{v_{i} \in \mathbf{V}(\Omega_{i}), \ div(v_{i}) = 0 \ \text{in } \Omega_{i}\Big\}. \end{split}$$

Restrictions of the functionals a and  $\Sigma$  over  $\Omega_i$  are defined by  $a_i$  and  $\Sigma_i$  respectively. 55 Inner products over a given part S of  $\partial \Omega_i$ , i = 1, 2, and  $\Omega_i$  are defined by 56

$$(u,v)_S = \int_S uv d\Gamma$$
 and  $(u,v)_{\Omega_i} = \int_{\Omega_i} uv dx$ .

We treat the pressure as a Lagrange multiplier associated with the constraint

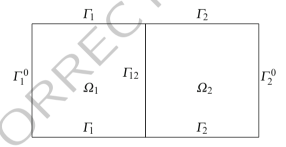


Fig. 1. Decomposition of  $\Omega$  into two subdomains

div(u) = 0. Using the decomposition of Fig. 1, the functional (5) becomes

$$\Sigma(\nu, \varphi) = \Sigma_1(\nu_1, \varphi_1) + \Sigma_2(\nu_2, \varphi_2). \tag{6}$$

It is clear that problem (3) is equivalent to the following constrained minimization 60 problem:

$$\forall (v_{i}, \varphi_{i}) \in \mathbf{V}(\Omega_{i}) \times H^{\frac{1}{2}}(\Gamma_{i}), i = 1, 2$$

$$\Sigma(u_{1}, \Phi_{1}) + \Sigma(u_{2}, \Phi_{2}) \leq \Sigma_{1}(v_{1}, \varphi_{1}) + \Sigma_{2}(v_{2}, \varphi_{2})$$

$$div(u_{i}) = 0 \quad \text{in } \Omega_{i},$$

$$u_{it} - \Phi_{i} = 0 \quad \text{in } \Gamma_{i},$$

$$u_{i} - \Psi = 0 \quad \text{in } \Gamma_{12}.$$

$$(7)$$

The auxiliary interface unknown  $\Psi$  is added to the continuity constraint to avoid 62 coupling between  $u_1$  and  $u_2$  in the penalty term. This so-called *three-field formula-63 tion* has been used in domain decomposition of elliptic problems [9]. To ensure the 64 uniqueness of the pressure, the following constraint can be added 65

$$\int_{\Omega_1} p_1 d\Omega_1 + \int_{\Omega_2} p_2 d\Omega_1 = 0.$$
 (8)

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Then, we introduce the set

$$\mathfrak{P} = \left\{ (q_1, q_2) \in L^2(\Omega_1) \times L^2(\Omega_2) \text{ such that } \int_{\Omega_1} q_1 d\Omega_1 + \int_{\Omega_2} q_2 d\Omega_1 = 0 \right\}$$
 67

We can associate to (7) the augmented Lagrangian functional  $\mathcal{L}_r$  defined by

$$\mathcal{L}_{r} (u, \Phi, \Psi, p, \mu, \lambda) = \Sigma(u_{1}, \Phi_{1}) + \Sigma(u_{2}, \Phi_{2}) 
+ \sum_{i=1}^{2} \left[ (\mu_{i}, \Phi_{i} - u_{it})_{\Gamma_{i}} - (p_{i}, div(u_{i}))_{\Omega_{i}} + (\lambda_{i}, u_{i} - \Psi)_{\Gamma_{12}} \right] 
+ \sum_{i=1}^{2} \left[ \frac{r_{1}}{2} ||div(u_{i})||_{L^{2}(\Omega_{i})}^{2} + \frac{r_{2}}{2} ||\Phi_{i} - u_{it}||_{L^{2}(\Gamma_{i})}^{2} + \frac{r_{3}}{2} ||u_{i} - \Psi||_{L^{2}(\Gamma_{12})}^{2} \right].$$
(9)

where  $r_1$ ,  $r_2$  and  $r_3$  are the penalty parameters which are strictly positive.

Remark 1. The standard  $L^2$  scalar product (not equivalent to the  $H^{1/2}$  scalar product) 70 on the interface  $\Gamma_{12}$  and  $\Gamma_i$  is used in the definition of (9). This approach is easy to 71 implement but it has some negative effects on the convergence of our algorithm.

Then, problem (7) is equivalent to the following saddle-point problem:

$$\begin{cases} \text{Find } (u, \Phi, \Psi, p, \mu, \lambda) \in \mathscr{H} & \text{such that: } \forall (v, \Phi, \Psi, q, \tilde{\mu}, \tilde{\lambda}) \in \mathscr{H} \\ \mathscr{L}_r(u, \Phi, \Psi, q, \tilde{\mu}, \tilde{\lambda}) \leq \mathscr{L}_r(u, \Phi, \Psi, p, \mu, \lambda) \leq \mathscr{L}_r(v, \Phi, \Psi, p, \mu, \lambda). \end{cases}$$
(10)

where  $u=(u_1,u_2)\in \mathbf{V}(\Omega_1)\times \mathbf{V}(\Omega_2)$ ,  $\Phi=(\Phi_1,\Phi_2)\in L^2(\Gamma_1)\times L^2(\Gamma_2)$ ,  $\Psi\in (L^2(\Gamma_{12}))^2$ , 74  $p=(p_1,p_2)\in \mathfrak{P}$ ,  $\mu=(\mu_1,\mu_2)\in L^2(\Gamma_1)\times L^2(\Gamma_2)$  and  $\lambda\in (L^2(\Gamma_{12}))^2$ .  $\mathscr{H}$  is the 75 Cartesian product of all these spaces.

#### 3.1 Uzawa Block Relaxation Method: UBR2

In order to solve (10) we use Uzawa block relaxation algorithm based on ALG2, see 78 [4]. This leads to the following iterations: 79

Initialization:  $\Phi^{-1}$ ,  $\Psi^{-1}$ ,  $p^0$ ,  $\lambda^0$ ,  $\mu^0$  and  $r_i>0$  fixed. 80 Repeat until convergence:

1. Find 
$$u^k \in \mathbf{V}(\Omega_1) \times \mathbf{V}(\Omega_2)$$
 such that:  $\forall v \in \mathbf{V}(\Omega_1) \times \mathbf{V}(\Omega_2)$  82 
$$\mathscr{L}_r(u^k, \Phi^{k-1}, \Psi^{k-1}, p^k, \mu^k, \lambda^k) < \mathscr{L}_r(v, \Phi^{k-1}, \Psi^{k-1}, p^k, \mu^k, \lambda^k).$$
 (11)

Domain Decomposition Method for Stokes Problem with Tresca Friction

2. Find 
$$\Phi^k \in L^2(\Gamma_1) \times L^2(\Gamma_2)$$
 such that:  $\forall \Phi \in L^2(\Gamma_1) \times L^2(\Gamma_2)$ 

$$\mathcal{L}_r(u^k, \boldsymbol{\Phi}^k, \boldsymbol{\Psi}^{k-1}, p^k, \boldsymbol{\mu}^k, \boldsymbol{\lambda}^k) \le \mathcal{L}_r(u^k, \boldsymbol{\Phi}, \boldsymbol{\Psi}^{k-1}, p^k, \boldsymbol{\mu}^k, \boldsymbol{\lambda}^k). \tag{12}$$

3. Find 
$$\Psi^k \in (L^2(\Gamma_{12}))^2$$
 such that:  $\forall \Psi \in (L^2(\Gamma_{12}))^2$ .

$$\mathcal{L}_r(u^k, \Phi^k, \Psi^k, p^k, \mu^k, \lambda^k) \le \mathcal{L}_r(u^k, \Phi^k, \Psi, p^k, \mu^k, \lambda^k). \tag{13}$$

4. Lagrange multipliers update

$$p_i^{k+1} = p_i^k - r_1 div(u_i^k), (14)$$

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$$\lambda_i^{k+1} = \lambda_i^k + r_2(u_{i|\Gamma_{12}}^k - \Psi^k),$$

$$\mu_i^{k+1} = \mu_i^k + r_3(u_{ii}^k - \Phi_i^k).$$
(15)

$$\mu_i^{k+1} = \mu_i^k + r_3(\mu_{it}^k - \Phi_i^k). \tag{16}$$

Subproblem (11) is equivalent to solving, in each subdomain, the following problem: 86

Find  $u_i^k \in \mathbf{V}(\Omega_i)$  such that

$$a(u_{i}^{k}, v) + r_{1}(\nabla . u_{i}^{k}, \nabla . v_{i})_{\Omega_{i}} + r_{2}(u_{i}, v_{i})_{\Gamma_{12}} + r_{3}(u_{t}^{k}, v_{t})_{\Gamma} = (f_{i}, v_{i}) + (p_{i}, \nabla . v_{i})_{\Omega_{i}} + (r_{2}\Psi^{k} - \lambda^{k}, v_{i})_{\Gamma_{12}} + (r_{3}\Phi_{i}^{k-1} - \mu_{i}^{k}, v_{it})_{\Gamma_{i}} \quad \forall v_{i} \in \mathbf{V}(\Omega_{i}).$$
 (17)

The subproblems of steps 2 and 3 are uncoupled and consists in the following calcu- 87 lations:

$$\Phi_{i}^{k} = \begin{cases} \frac{||\mu_{i}^{k} + r_{3}u_{it}^{k}||_{0,\Gamma_{i}} - g}{r_{3}||\mu_{i}^{k} + r_{3}u_{it}^{k}||_{0,\Gamma_{i}}} (\mu_{i}^{k} + r_{3}u_{it}^{k}) & \text{if } ||\mu_{i}^{k} + r_{3}u_{it}^{k}||_{0,\Gamma_{i}} \ge g\\ 0 \text{ unless} \end{cases}$$
(18)

and

$$\Psi^k = \frac{1}{2r_2} (\lambda_1^k + \lambda_2^k) + \frac{1}{2} (u_1^k + u_2^k) |_{\Gamma_{12}}.$$
 (19)

Remark 2. For sake of simplicity the given slip bound g is assumed to be non- 91 negative constant in (18). 92

Remark 3. After update (14),  $p^{k+1}$  must be projected onto  $\mathfrak{P}$  to ensure the uniqueness 93 of the pressure. 94

Remark 4. The main advantage of this formulation is that (17) reduces to 2D un- 95 coupled elliptic problems which can be solved in parallel. Moreover, the matrices 96 derived from discret problems are symmetric and positive definite. 97

## 4 Numerical Experiments

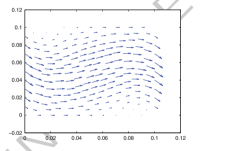
The domain decomposition algorithm **UBR2**, with  $r_1 = r_2 = r_3$ , presented in the previous section was implemented in Matlab V7.9 on a Core2 Duo-1.8 Ghz processor 100 PC. For discrete velocity-pressure-Lagrange multipliers spaces, we use the  $P^1$ -iso- $P^2/P^1$  finite element. These spaces are well known to satisfy the discrete Babuska-Brezzi inf-sup condition [1].

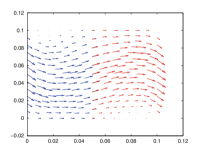
For all the numerical experiments presented, the domain  $\Omega$  is the square  $[0,0.1]^2$ , while  $\Omega_1 = [0, 0.05] \times [0, 0.1]$  and  $\Omega_2 = [0.05, 0.1] \times [0, 0.1]$ . The fluid can slip on 105  $\Gamma_1 \cup \Gamma_2 = [0, 0.1] \times \{0.1\} \cup [0, 0.1] \times \{0\}$ , We set g = 0.015 which is consistent with 106 experimental values, see [5]. The viscosity is taken equal to 0.1 and the stopping 107 tolerance  $\varepsilon$  is  $10^{-6}$ . In addition we enforce parabolic profile on both  $\Gamma_1^0 = \{0\} \times 10^{-6}$ [0,0.1] and  $\Gamma_2^0 = \{0.1\} \times [0,0.1]$ :

$$u|_{\Gamma_1^0} = u|_{\Gamma_2^0} = \begin{bmatrix} y(1-y) \\ -y(1-y) \end{bmatrix}$$

Remark 5. We choose this profile to enforce shear stress near the solid wall to reach 110 the threshold without considering a complicated domain geometry.

In Fig. 2 we report the velocity field for the solution of Stokes problem with Tresca 112 friction (1) in  $\Omega$  and in  $\Omega_1 \cup \Omega_2$ . We can see that we have the same velocity profile. 113 In Table 1 we report the discrete mesh size h, the corresponding number of degree





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Fig. 2. Fluid flow with Tresca BC for one (left) and two domains (right)

of freedom (d.o.f) and number of elements on each subdomain in the follows exper- 115 iments. Table 2 shows the number of iterations IT, the sequential CPU (in seconds) 116 times and the parallel CPU\* times (when subproblems (17) for i = 1, 2 are solved in 117 parallel). For several mesh size and for N<sub>SD</sub> (Number of Sub-Domains) equal to 1 118 or 2. We notice that the **UBR2** algorithm is a h-dependent algorithm and the domain 119 decomposition method to be preferable when dealing with parallel computing using 120 parallel solver.

Table 3 show how the number of iterations and the optimal value of the relax- 122 ation parameter  $r_{opt}$  depend on h. We remark that the speed of convergence is very 123 sensitive to r; this explains the strong increase in the number of iterations for a finer 124 mesh.

Domain Decomposition Method for Stokes Problem with Tresca Friction

$N_{SD}$	h = 0.02	h = 0.01	h = 0.0067	h = 0.005	h = 0,004
1.430	$n/n_{\triangle}$	$n/n_{\triangle}$	$n/n_{\triangle}$	$n/n_{\triangle}$	$n/n_{\triangle}$
1	189/336	665/1284	1577/3032	2829/5496	4393/8548
2	112/188	370/676	806/1516	1396/2668	2220/4284

**Table 1.** h: mesh size; n: number of d.o.f. by domain  $n_{\Lambda}$ : number of elements by domain.

Nen	h = 0.02	h = 0.01	h = 0.0067	h = 0.005	h = 0,004
INSD	IT/CPU/CPU*	IT/CPU/CPU*	IT/CPU/CPU*	IT/CPU/CPU*	IT/ CPU/CPU*
1	199/0.41/-	349/2.8/-	453/10.8/-	509/30.36/-	595/67.3/-
2	486/1/0.81	769/4.8/3.27	993/15.3/7.96	1294/41.14/21.98	1599/99.34/51.59

t2.1 t2.2 t2.3

**Table 2.** Standard speed-up for h: mesh size; IT: number of iterations; CPU & CPU\*: CPU times.

$N_{SD}$			h = 0.0067	h = 0.005	h = 0,004
1.430	r <sub>opt</sub> /IT	$r_{opt}$ /IT	$r_{opt}/\mathrm{IT}$	$r_{opt}$ /IT	r <sub>opt</sub> /IT
1	335/199	590/349	740/453	840/509	1010/595
2	116/486	124/769	175/993	230/1294	290/1599

**Table 3.** Convergence rate with respect  $r_{opt}$ .

5 Conclusion 126

The augmented Lagrangian formulation (9) of domain decomposed Stokes problem with Tresca friction leads to a numerical strategy which solves a classical Poisson problem (17) (in each subdomain  $\Omega_i$ ) and the contribution of Tresca friction (18) in decoupled way. Nevertheless, this algorithm has a mesh dependent convergence and its practical implementation still facing the issue of the optimal choice of the penalties,  $r_i$ , i = 1, 2, 3. To improve this algorithm, different preconditioners will be investigated, especially the Steklov-Poincaré operator on the interface (see e.g. [6–133 8]) and the Cahouet-Chabard preconditioner [2] for the pressure multiplier.

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# A Hybrid Discontinuous Galerkin Method for Darcy-Stokes Problems

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Summary. We propose and analyze a hybrid discontinuous Galerkin method for the solution 10 of incompressible flow problems, which allows to deal with pure Stokes, pure Darcy, and 11 coupled Darcy-Stokes flow in a unified manner. The flexibility of the method is demonstrated 12 in numerical examples.

### 1 Model Problem

Let  $\Omega\subset\mathbb{R}^d$  be a bounded Lipschitz domain in d=2 or 3 dimensions. Given data 15  $\mathbf{f} \in [L^2(\Omega)]^d$  and  $g \in L^2(\Omega)$ , we consider the generalized Stokes problem 16

$$\sigma \mathbf{u} - 2\mu \operatorname{div} \varepsilon(\mathbf{u}) + \nabla p = \mathbf{f}$$
 and  $\operatorname{div} \mathbf{u} = g$  in  $\Omega$ . (1)

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As usual, **u** denotes the velocity, p the pressure, and  $\varepsilon(\mathbf{u}) := \frac{1}{2}(\nabla \mathbf{u} + \nabla \mathbf{u}^T)$  is the symmetric part of the velocity gradient tensor. We require that

$$\sigma \ge 0$$
,  $\mu \ge 0$ , and  $M \ge \sigma + \mu \ge m > 0$  in  $\Omega$ .

For convenience, we assume that  $\sigma$ , the reciprocal of the permeability, and the viscosity  $\mu$  are constant, and consider homogeneous boundary conditions 20

$$\mathbf{u}|_{\partial\Omega} = 0 \quad \text{if} \quad \mu > 0 \quad \text{or} \quad \mathbf{u} \cdot \mathbf{n}|_{\partial\Omega} = 0 \quad \text{if} \quad \mu = 0.$$
 (2)

The unique solvability of the boundary value problem (1)–(2) is guaranteed, if 21 the pressure p and the data g have zero average. For the case  $\mu > 0$ , we then 22 have  $(\mathbf{u},p)\in\mathbf{H}_0^1(\Omega)\times L_0^2(\Omega)$ , where  $\mathbf{H}_0^1(\Omega):=\{\mathbf{v}\in[H^1(\Omega)]^d:\mathbf{v}|_{\partial\Omega}=0\}$  and 23  $L_0^2:=\{q\in L^2(\Omega):\int_\Omega q\ dx=0\}$ . In the Darcy limit  $\mu=0$ , we only have  $\mathbf{u}\in\mathbf{u}\in\mathbf{u}$   $\mathbf{u}\in\mathbf{u}$   $\mathbf{u}\in\mathbf{u}$ 

For the approximation of problem (1)–(2), we consider a hybrid discontinuous 26 Galerkin method, which is capable of treating incompressible flow in the Stokes 27

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and Darcy regimes, as well as coupled problems in a unified manner. Our analysis 28 extends the results of [7] for Stokes flow. Related work on stabilized non-conforming 29 and discontinuous Galerkin methods for Darcy-Stokes flow can be found in [4, 8] and 30 the references given there. We refer to [1, 5] for a unified treatment of discontinuous 31 Galerkin methods for elliptic problems and their hybridization.

### 2 Notation and Preliminaries

Let  $\mathscr{T}_h = \{T\}$  be a shape-regular quasi-uniform partition of  $\Omega$  into affine families 34 of triangles and/or quadrilaterals (tetrahedra and/or hexahedra) of size h. By  $\partial \mathscr{T}_h := 35$  $\{\partial T: T\in \mathcal{T}_h\}$ , we denote the set of element boundaries, and by  $\mathcal{E}_h:=\{E_{ij}=\partial T_i\cap \mathcal{T}_i\}$  $\partial T_i : i > j \} \cup \{E_{i,0} = \partial T_i \cap \partial \Omega\}$  the set of edges (faces) between elements or on the boundary;  $\mathscr{E} = \bigcup_{E \in \mathscr{E}_h} E$  is called the *skeleton*.

For  $s \ge 0$ , let  $H^s(\mathscr{T}_h) := \{ v \in L^2(\Omega) : v|_T \in H^s(T) \text{ for all } T \in \mathscr{T}_h \}$  denote the broken Sobolev space with inner product  $(u,v)_{s,\mathscr{T}_h} := \sum_{T \in \mathscr{T}_h} (u,v)_{s,T}$  and norm  $||u||_{s,\mathscr{T}_h}$ ; 40 the subindex is omitted for s = 0. Piecewise defined derivatives are denoted with 41 the standard symbols. The traces of functions in  $H^1(\mathcal{T}_h)$  lie in  $L^2(\partial \mathcal{T}_h)$ , which is 42 equipped with the scalar product  $\langle u, v \rangle_{\partial \mathscr{T}_h} := \sum_{T \in \mathscr{T}_h} \langle u, v \rangle_{\partial T}$  and norm  $|v|_{\partial \mathscr{T}_h}$ . Any 43 function in  $L^2(\mathscr{E})$  can be identified with a function in  $L^2(\partial \mathscr{T}_h)$  by doubling its values 44 on the element interfaces. Bold symbols are used for vector valued functions. 45

Let  $\mathscr{P}_{p}(T)$  denote the polynomials of degree  $\leq p$  over T, and recall that

$$|v_{\mathbf{p}}|_{\partial T}^2 \le c_T \, \mathbf{p}^2 \mathbf{h}^{-1} ||v_{\mathbf{p}}||_T^2 \qquad \text{for all } v_{\mathbf{p}} \in \mathscr{P}_{\mathbf{p}}(T). \tag{3}$$

Explicit bounds for the constant  $c_T$  in the discrete trace inequality (3) are known for 47 all elements under consideration. The parameter  $c_T$  can be replaced by the shape 48 regularity parameter  $\gamma := \max\{c_T : T \in \mathcal{T}_h\}$ , which is assumed to be independent of h. We then choose a stabilization parameter  $\alpha$  such that

$$4\gamma p^2 h^{-1} \le \alpha \le 4\gamma' p^2 h^{-1},\tag{4}$$

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with  $\gamma'$  independent of p and h, and we define two norms on  $L^2(\partial \mathcal{T}_h)$  by

$$|v|_{\pm 1/2, \partial \mathcal{T}_h} := \left(\sum_{T \in \mathcal{T}_h} |v|_{\pm 1/2, \partial T}^2\right)^{1/2} \text{ with } |v|_{\pm 1/2, \partial T} := \alpha^{\pm 1/2} |v|_{\partial T}.$$

Similar norms are frequently used for the analysis of mixed, non-conforming and 52 discontinuous Galerkin methods; see e.g. [1]. 53

# 3 The Hybrid DG Method

Let us fix  $p \ge 1$ , and choose q = p - 1 or q = p. For the approximation of velocity 55 and pressure in (1)–(2), we will utilize the finite element spaces 56 A Hybrid Discontinuous Galerkin Method for Darcy-Stokes Problems

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$$\mathbf{V}_h := \{ \mathbf{v}_h \in \mathbf{L}^2(\mathscr{T}_h) : \mathbf{v}_h|_T \in [\mathscr{P}_p(T)]^d \quad \text{for all } T \in \mathscr{T}_h \},$$

$$Q_h := \{ q_h \in L_0^2(\Omega) : q_h|_T \in \mathscr{P}_q(T) \quad \text{for all } T \in \mathscr{T}_h \}.$$

We further choose  $\hat{p} = p$  or  $\hat{p} = q$ , and define a space

$$\widehat{\mathbf{V}}_h := \{\widehat{\mathbf{v}}_h \in \mathbf{L}^2(\mathscr{E}) : \widehat{\mathbf{v}}_h|_E \in [\mathscr{P}_{\hat{\mathbf{p}}}(E)]^d \quad \text{for all } E \in \mathscr{E}_h, \ \widehat{\mathbf{v}}_h = 0 \text{ on } \partial\Omega\},$$

of piecewise polynomials for representing velocities on the skeleton. The conditions 58  $p-1 \le q \le p$  and  $q \le \hat{p}$  are explicitly used in the analysis of a Fortin operator; see 59 Proposition 5. In view of Lemma 1, we also require that  $\hat{p} \ge 1$ . Note that the Dirichlet 60 boundary condition has been included explicitly in the definition of the hybrid space 61  $\hat{\mathbf{V}}_h$ . We further denote by  $\pi^p : \mathbf{H}^1(\mathscr{T}_h) \to \mathbf{V}_h$  and  $\hat{\pi}^{\hat{p}} : \mathbf{L}^2(\mathscr{E}) \to \hat{\mathbf{V}}_h$ , the  $L^2$  orthogonal 62 projections onto the discrete spaces. The boundary value problem (1)–(2) is then 63 approximated by the following finite element scheme.

**Method 1.** Find  $\mathbf{u}_h \in \mathbf{V}_h$ ,  $\widehat{\mathbf{u}}_h \in \widehat{\mathbf{V}}_h$ , and  $p_h \in Q_h$ , such that

$$\begin{cases} \mathbf{a}_h(\mathbf{u}_h, \widehat{\mathbf{u}}_h; \mathbf{v}_h, \widehat{\mathbf{v}}_h) + \mathbf{b}_h(\mathbf{v}_h, \widehat{\mathbf{v}}_h; p_h) = (\mathbf{f}, \mathbf{v}_h)_{\mathscr{T}_h}, \\ \mathbf{b}_h(\mathbf{u}_h, \widehat{\mathbf{u}}_h; q_h) = (g, q_h)_{\mathscr{T}_h}, \end{cases}$$

for all  $\mathbf{v}_h \in \mathbf{V}_h$ ,  $\widehat{\mathbf{v}}_h \in \widehat{\mathbf{V}}_h$ , and  $q_h \in Q_h$ . The bilinear forms are defined as

$$\mathbf{a}_{h}(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}) := \sigma \mathbf{d}_{h}(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}) + 2\mu \mathbf{s}_{h}(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}), \mathbf{b}_{h}(\mathbf{v},\widehat{\mathbf{v}};q) := -(\operatorname{div}\mathbf{v},q)_{\mathscr{T}_{h}} + \langle \mathbf{v} - \widehat{\mathbf{v}}, q\mathbf{n} \rangle_{\partial \mathscr{T}_{h}},$$

and the bilinear forms  $\mathbf{d}_h$  and  $\mathbf{s}_h$  are given by

$$\begin{split} \mathbf{d}_{\textit{h}}(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}) &:= (\mathbf{u},\mathbf{v})_{\mathscr{T}_{\textit{h}}} + \alpha \langle (\widehat{\pi}^{\hat{\mathbf{p}}}\mathbf{u} - \widehat{\mathbf{u}}) \cdot \mathbf{n}, (\widehat{\pi}^{\hat{\mathbf{p}}}\mathbf{v} - \widehat{\mathbf{v}}) \cdot \mathbf{n} \rangle_{\partial \mathscr{T}_{\textit{h}}}, \\ \mathbf{s}_{\textit{h}}(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}) &:= (\varepsilon(\mathbf{u}),\varepsilon(\mathbf{v}))_{\mathscr{T}_{\textit{h}}} - \langle \varepsilon(\mathbf{u}) \cdot \mathbf{n}, \mathbf{v} - \widehat{\mathbf{v}} \rangle_{\partial \mathscr{T}_{\textit{h}}} \\ &- \langle \mathbf{u} - \widehat{\mathbf{u}},\varepsilon(\mathbf{v}) \cdot \mathbf{n} \rangle_{\partial \mathscr{T}_{\textit{h}}} + \alpha \langle \widehat{\pi}^{\hat{\mathbf{p}}}\mathbf{u} - \widehat{\mathbf{u}},\widehat{\pi}^{\hat{\mathbf{p}}}\mathbf{v} - \widehat{\mathbf{v}} \rangle_{\partial \mathscr{T}_{\textit{h}}}. \end{split}$$

One easily verifies that any regular solution of (1)–(2) also satisfies the discrete variational principle above. 68

**Proposition 1 (Consistency).** Let  $(\mathbf{u}, p)$  denote a solution of (1)–(2), and assume 70 additionally that  $\mathbf{u} \in \mathbf{H}^2(\mathcal{T}_h)$  and  $p \in H^1(\mathcal{T}_h)$ . Then

$$\mathbf{a}_h(\mathbf{u}, \mathbf{u}; \mathbf{v}_h, \widehat{\mathbf{v}}_h) + \mathbf{b}_h(\mathbf{v}_h, \widehat{\mathbf{v}}_h; p) = (\mathbf{f}, \mathbf{v}_h)_{\mathscr{T}_h}$$
 and  $\mathbf{b}_h(\mathbf{u}, \mathbf{u}; q_h) = (g, q_h)_{\mathscr{T}_h}$ 

for all  $\mathbf{v}_h \in \mathbf{V}_h$ ,  $\widehat{\mathbf{v}}_h \in \widehat{\mathbf{V}}_h$ , and  $q_h \in Q_h$ ; thus, Method 1 is consistent.

In the Darcy limit  $\mu = 0$ , it suffices to require  $\mathbf{u} \in \mathbf{H}^1(\mathscr{T}_h)$ .

# 4 Stability and Error Analysis

An important ingredient for our analysis will be the following result.

**Lemma 1** (Discrete Korn inequality). Let  $\hat{p} \ge 1$ . Then there is a  $\kappa > 0$  independent 76 of h, such that for all  $\mathbf{v} \in \mathbf{H}^1(\mathscr{T}_h)$  and  $\widehat{\mathbf{v}} \in \mathbf{L}^2(\mathscr{E})$ , there holds

$$\|\varepsilon(\mathbf{v})\|_{\mathscr{T}_h}^2 + |\widehat{\pi}^{\hat{\mathbf{p}}}(\mathbf{v} - \widehat{\mathbf{v}})|_{1/2, \partial \mathscr{T}_h}^2 \ge \kappa \|\nabla \mathbf{v}\|_{\mathscr{T}_h}^2. \tag{5}$$

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*Proof.* The statement follows via the triangle inequality from Korn's inequality for piecewise  $H^1$  functions [3, Eq. (1.12)] established by Brenner. Æ.

**Proposition 2.** For any  $(\mathbf{v}_h, \widehat{\mathbf{v}}_h) \in \mathbf{V}_h \times \widehat{\mathbf{V}}_h$  there holds

$$\mathbf{s}_{h}(\mathbf{v}_{h},\widehat{\mathbf{v}}_{h};\mathbf{v}_{h},\widehat{\mathbf{v}}_{h}) \geq \min\{\frac{5}{12},\frac{\kappa}{4}\}(\|\nabla\mathbf{u}\|_{\mathscr{T}_{h}}^{2} + |\widehat{\pi}^{\widehat{\mathbf{p}}}(\mathbf{u} - \widehat{\mathbf{u}})|_{1/2,\partial\mathscr{T}_{h}}^{2}).$$
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We young's inequality, Eq. (3) and (4), we obtain

*Proof.* By Young's inequality, Eq. (3) and (4), we obtain

$$-2\langle \boldsymbol{\varepsilon}(\mathbf{v}_h) \cdot \mathbf{n}, \mathbf{v}_h - \widehat{\mathbf{v}}_h \rangle_{\partial T} \ge -\frac{3}{4} \|\boldsymbol{\varepsilon}(\mathbf{v}_h)\|_T^2 - \frac{1}{3} |\widehat{\boldsymbol{\pi}}^{\hat{\mathbf{p}}}(\mathbf{v}_h - \widehat{\mathbf{v}}_h)|_{1/2, \partial T}^2.$$

The result then follows by Lemma 1, and the definition of  $s_h$ .

For appropriately characterizing the coercivity of the bilinear form  $\mathbf{d}_h$ , let us introduce the discrete kernel space for the bilinear form  $\mathbf{b}_h$ , namely 83

$$\mathbf{K}_h := \{ (\mathbf{v}_h, \widehat{\mathbf{v}}_h) \in \mathbf{V}_h \times \widehat{\mathbf{V}}_h : \mathbf{b}_h(\mathbf{v}_h, \widehat{\mathbf{v}}_h; q_h) = 0 \ \forall q_h \in Q_h \}.$$

**Proposition 3.** For any pair of functions  $(\mathbf{v}_h, \widehat{\mathbf{v}}_h) \in \mathbf{K}_h$  there holds

$$\mathbf{d}_h(\mathbf{v}_h, \widehat{\mathbf{v}}_h; \mathbf{v}_h, \widehat{\mathbf{v}}_h) \ge \|\mathbf{v}_h\|_{\mathscr{T}_h}^2 + \|\operatorname{div}\mathbf{v}_h\|_{\mathscr{T}_h}^2 + \frac{3}{4}|\widehat{\pi}^{\widehat{\mathbf{p}}}(\mathbf{v}_h - \widehat{\mathbf{v}}_h) \cdot \mathbf{n}|_{1/2, \partial \mathscr{T}_h}^2.$$

*Proof.* Note that for every  $T \in \mathcal{T}_h$  we have  $\operatorname{div} \mathbf{v}_h|_T \in \mathcal{P}_q(T)$ . Testing with  $q_h = 1$  $\operatorname{div} \mathbf{v}_h$  and using (3) yields 89

$$\|\operatorname{div}\mathbf{v}_h\|_T^2 = \langle (\mathbf{v}_h - \widehat{\mathbf{v}}_h) \cdot \mathbf{n}, \operatorname{div}\mathbf{v}_h \rangle_{\partial T} \leq \frac{1}{2} |(\widehat{\boldsymbol{\pi}}^{\widehat{\mathbf{p}}}\mathbf{v}_h - \widehat{\mathbf{v}}_h) \cdot \mathbf{n}|_{1/2, \partial T} \|\operatorname{div}\mathbf{v}_h\|_T,$$

and hence  $\|\operatorname{div} \mathbf{v}_h\|_{\mathscr{T}_h} \leq \frac{1}{2} |(\widehat{\boldsymbol{\pi}}^{\hat{\mathbf{p}}} \mathbf{v}_h - \widehat{\mathbf{v}}_h) \cdot \mathbf{n}|_{1/2, \partial \mathscr{T}_h}$ . The result then follows by adding and subtracting  $\|\operatorname{div} \mathbf{v}_h\|_{\partial \mathscr{T}_h}^2$  from the bilinear form  $\mathbf{d}_h$ .

The two coercivity estimates suggest to utilize the following energy norms for the 90 stability analysis of Method 1, namely,  $||q||_{0,\mathcal{T}_h}$  and 91

$$\begin{split} \|(\mathbf{v},\widehat{\mathbf{v}})\|_{1,\mathcal{T}_h}^2 &:= \sigma\big(\|\mathbf{v}\|_{\mathcal{T}_h}^2 + \|\operatorname{div}\mathbf{v}\|_{\mathcal{T}_h}^2 + |\widehat{\pi}^{\hat{\mathbf{p}}}(\mathbf{v} - \widehat{\mathbf{v}}) \cdot \mathbf{n}|_{1/2,\partial\mathcal{T}_h}^2\big) \\ &+ \mu\big(\|\nabla\mathbf{v}\|_{\mathcal{T}_h}^2 + |\widehat{\pi}^{\hat{\mathbf{p}}}(\mathbf{v} - \widehat{\mathbf{v}})|_{1/2,\partial\mathcal{T}_h}^2\big). \end{split}$$

Remark 1. If  $\mu = 0$ , then  $\|(\cdot, \cdot)\|_{1, \mathcal{T}_h}$  is only a semi-norm on  $\mathbf{V}_h \times \widehat{\mathbf{V}}_h$ . This deficiency 93 can be overcome by eliminating the tangential velocities in the definition of the hybrid space, or by penalizing also the jump of the tangential velocities in the bilinear 95 form  $\mathbf{d}_h$ . Both remedies do not affect our analysis. 96

A combination of Propositions 2 and 3 now yields the kernel ellipticity for  $\mathbf{a}_h$ .

**Proposition 4 (Coercivity).** For any element  $(\mathbf{v}_h, \widehat{\mathbf{v}}_h) \in \mathbf{K}_h$  there holds

$$\mathbf{a}_h(\mathbf{v}_h, \widehat{\mathbf{v}}_h; \mathbf{v}_h, \widehat{\mathbf{v}}_h) \ge \min\{\frac{3}{4}, \frac{\kappa}{2}\} \|(\mathbf{v}_h, \widehat{\mathbf{v}}_h)\|_{1, \mathcal{T}_h}^2.$$

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The constants  $C_i$  appearing in the following results depend on the bounds m and M, 100 but are else independent of the parameters  $\mu$ ,  $\sigma$ , and of h and p. Let us next consider the operator  $(\pi^p, \widehat{\pi}^{\widehat{p}}) : \mathbf{H}_0^1(\Omega) \to \mathbf{V}_h \times \widehat{\mathbf{V}}_h$ . 102

**Proposition 5 (Fortin operator).** *For any*  $\mathbf{v} \in \mathbf{H}_0^1(\Omega)$  *there holds* 

$$b_h(\pi^p \mathbf{v}, \widehat{\pi}^{\widehat{\mathbf{p}}} \mathbf{v}; q_h) = b(\mathbf{v}, q_h) \quad \forall q_h \in Q_h, \tag{6}$$

and 
$$\|(\pi^{\mathbf{p}}\mathbf{v},\widehat{\pi}^{\hat{\mathbf{p}}}\mathbf{v})\|_{1,\mathcal{T}_{h}} \le C_{\pi}\,\mathbf{p}^{1/2}\|\mathbf{v}\|_{1,\Omega}.$$
 (7)

*Proof.* Equation (6) follows from the properties of the projections, and (7) results from stability estimates for the  $L^2$  projections; cf. [7] for details.

The inf-sup stability of  $\mathbf{b}_h$  now follows directly from the previous result.

**Proposition 6 (Inf-sup condition).** *For any*  $q_h \in Q_h$  *there holds* 

$$\sup_{(\mathbf{y}_h,\widehat{\mathbf{y}}_h)\in\mathbf{V}_h\times\widehat{\mathbf{V}}_h} \frac{\mathbf{b}_h(\mathbf{v}_h,\widehat{\mathbf{v}}_h;q_h)}{\|(\mathbf{v}_h,\widehat{\mathbf{v}}_h)\|_{1,\mathscr{T}_h}} \ge C_\beta \, \mathbf{p}^{-1/2} \|q_h\|_{0,\mathscr{T}_h}. \tag{8}$$

As a consequence of Propositions 4 and 6, we obtain by Brezzi's theorem that 106 Method 1 has a unique solution and thus is well-defined. Next, we show the boundedness of the bilinear forms with respect to a pair of stronger norms defined by 108  $|||q_h||_{0,\mathscr{T}_h}^2 := ||q_h||_{\mathscr{T}_h}^2 + |q_h \cdot \mathbf{n}|_{-1/2,\partial\mathscr{T}_h}^2$  and

$$\||(\mathbf{v}_h,\widehat{\mathbf{v}}_h)||_{1,\mathscr{T}_h}^2 := \|(\mathbf{v}_h,\widehat{\mathbf{v}}_h)||_{1,\mathscr{T}_h}^2 + \mu |\partial_{\mathbf{n}}\mathbf{v}_h|_{-1/2,\partial\mathscr{T}_h}^2,$$

The norms  $\|\cdot\|_{0,\mathscr{T}_h}$ ,  $\|(\cdot,\cdot)\|_{1,\mathscr{T}_h}$  and  $\|\cdot\|_{0,\mathscr{T}_h}$ ,  $\|(\cdot,\cdot)\|_{1,\mathscr{T}_h}$  are equivalent on the finite 110 element spaces with equivalence constants less than two. This yields coercivity and 111 inf-sup stability of  $\mathbf{a}_h$  and  $\mathbf{b}_h$  also with respect to the stronger norms. The following bounds then follow from the Cauchy-Schwarz inequality. 113

**Proposition 7 (Boundedness).** For any  $\widehat{\mathbf{u}}$ ,  $\widehat{\mathbf{v}}\in\widehat{\mathbf{V}}_h\oplus\mathbf{L}^2(\mathscr{E})$  and every function  $\mathbf{u}$ ,  $\mathbf{v} \in \mathbf{V}_h \oplus (\mathbf{H}_0^1(\Omega) \cap \mathbf{H}^2(\mathscr{T}_h))$ , there holds 115

$$\mathbf{a}_h(\mathbf{u},\widehat{\mathbf{u}};\mathbf{v},\widehat{\mathbf{v}}) \leq C_a \| (\mathbf{u},\widehat{\mathbf{u}}) \|_{1,\mathscr{T}_h} \| (\mathbf{v},\widehat{\mathbf{v}}) \|_{1,\mathscr{T}_h},$$

and for all  $p \in Q_h \oplus (L_0^2(\Omega) \cap H^1(\mathcal{T}_h))$ , there holds additionally

$$\mathbf{b}_h(\mathbf{u},\widehat{\mathbf{u}};p) \leq C_b \| (\mathbf{u},\widehat{\mathbf{u}}) \|_{1,\mathscr{T}_h} \| p \|_{0,\mathscr{T}_h}.$$

Standard polynomial approximation results [2] imply the following properties.

**Proposition 8 (Approximation).** Assume  $s \ge 1$ . Then for any function  $\mathbf{u} \in \mathbf{H}_0^1(\Omega) \cap$  $\mathbf{H}^{s+1}(\mathscr{T}_h)$  there exist elements  $\mathbf{v}_h \in \mathbf{V}_h$  and  $\widehat{\mathbf{v}}_h \in \widehat{\mathbf{V}}_h$  such that 119

$$\| (\mathbf{u} - \mathbf{v}_h, \mathbf{u} - \widehat{\mathbf{v}}_h) \|_{1,\mathcal{T}_h} \le C_{ap} \, \mathbf{p}^{1/2-s} h^{\min\{\mathbf{p},s\}} \| \mathbf{u} \|_{s+1,\mathcal{T}_h},$$

and for any  $p \in L_0^2(\Omega) \cap H^s(\mathcal{T}_h)$  there exists a  $q_h \in Q_h$  such that

$$|||p-q_h||_{0,\mathscr{T}_h} \le C_{ap} p^{-s} h^{\min\{s,q+1\}} ||p||_{s,\mathscr{T}_h}.$$

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The a-priori estimates now follow by combination of the previous results.

**Proposition 9 (Error estimate).** Let  $(\mathbf{u}, p)$  be the solution of (1)–(2), and let 122  $(\mathbf{u}_h, \widehat{\mathbf{u}}_h, p_h)$  denote the approximation defined by Method 1. Then 123

$$\begin{aligned} \| (\mathbf{u} - \mathbf{u}_h, \mathbf{u} - \widehat{\mathbf{u}}_h) \|_{1,\mathcal{T}_h} + \mathbf{p}^{-1/2} \| p - p_h \|_{0,\mathcal{T}_h} \\ & \leq C_{err} \, \mathbf{p}^{1/2} h^{\min\{\mathbf{p},s\}} \big( \mathbf{p}^{1/2-s} \| \mathbf{u} \|_{s+1,\mathcal{T}_h} + \mathbf{p}^{-s} \| p \|_{s,\mathcal{T}_h} \big). \end{aligned}$$

Proof. The result follows with the usual arguments from the consistency, discrete 124 stability, and boundedness of the bilinear forms, and the approximation properties of 125 the finite element spaces; for details, see [7] or [9]. 126

5 Remarks 127

The analysis of Sect. 4 applies almost verbatim to spatially varying material param- 128 eters  $\mu$  and  $\sigma$ . In particular, a coupling of Darcy and Stokes equations in different 129 parts of the domain is possible and treated automatically. A numerical example for 130 such a case is presented in the next section.

Our results can be extended to shape regular meshes and varying polynomial 132 degree. Also meshes with a bounded number of hanging nodes on each edge or face, 133 and even more general non-conforming mortar meshes can be treated. We refer to 134 [6, 7] for a detailed discussion of conditions on the mesh and polynomial degree 135 distribution. 136

The coercivity and boundedness estimates also hold for more general finite 137 element spaces, but we explicitly utilized the complete discontinuity of the spaces 138 in the proof of the inf-sup condition. Other constructions of a Fortin-operator, cf. 139 e.g. [9], would allow to relax this assumption.

Our analysis also covers equal order approximations q = p, which are stabilized 141 sufficiently by the jump penalty terms.

All degrees of freedom except the piecewise constant pressure and the hybrid 143 velocities can be eliminated by static condensation on the element level. This leads 144 to small global systems, which for  $\hat{p} = 0$  exhibit the same sparsity pattern as a nonconforming  $P_1 - P_0$  discretization. For  $\hat{p} = 0$ , the discrete Korn inequality (5) is not valid, so this choice had to be excluded here. If  $\varepsilon(\mathbf{u})$  in (1) is replaced by  $\frac{1}{2}\nabla\mathbf{u}$ , we 147 however obtain a stable scheme.

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t1.1 t1.2 t1.3 t1.4 t1.5

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#### 6 Numerical Results

Let us now illustrate the capability of the proposed method to deal with incompressible flow in various regimes. Our numerical results were obtained with an implementation of Method 1 in NGSolve.<sup>3</sup>

As a first example, we consider the generalized Stokes equation (1) on the unit 153 square  $\Omega = (-1,1)^2$  with a known analytic solution given by 154

$$\mathbf{u} = (20xy^3, 5x^4 - 5y^4), \quad p = 60x^2y - 20y^3,.$$

The data f and g can be obtained from Eq. (1). For the numerical solution, we employed Method 1 with  $p = \hat{p} = 2$  and q = 1 on a sequence of uniformly refined 156 meshes for different values of  $\mu$  and  $\sigma$ . The analytic solution was used to compute 157 the errors listed in Table 1. As predicted by the theory, we can observe the optimal 158 quadratic convergence.

**Table 1.** Energy errors obtained by simulation on a sequence of uniformly refined meshes for  $(\sigma,\mu) \in \{(1,0),(\frac{1}{2},\frac{1}{2}),(0,1)\}$ , resembling Darcy, Brinkman, and Stokes flow.

				///		
level	Darcy	rate	Brinkman	rate	Stokes	rate
0	4.3996	- /	3.4058	-	3.8578	_
1	1.1261	1.96	0.8628	1.98	0.9764	1.98
2	0.2799	2.00	0.2146	2.00	0.2428	2.00
3	0.0678	2.04	0.0533	2.00	0.0603	2.00
		/				

As a second test case, we consider a coupled Darcy-Stokes flow on a domain consisting of two subdomains  $\Omega_D$  and  $\Omega_S$ , as depicted in Fig. 1. The flow in the subdomains is governed by

$$\sigma_i \mathbf{u}_i - 2\mu_i \operatorname{div} \varepsilon(\mathbf{u}_i) + \nabla p_i = 0$$
 and  $\operatorname{div} \mathbf{u}_i = 0$  in  $\Omega_i$ ,

with  $\mu_D = 0$  in the Darcy domain  $\Omega_D$ , and  $\sigma_S = 0$  in the Stokes domain  $\Omega_S$ , and the subproblems are coupled across the interface  $\partial \Omega_D \cap \partial \Omega_S$  through 164

$$\mathbf{u}_S \cdot \mathbf{n} = \mathbf{u}_D \cdot \mathbf{n}, \ p_S - 2\mu(\varepsilon(\mathbf{u}_S) \cdot \mathbf{n}) \cdot \mathbf{n} = p_D, \ \mathbf{u}_S \cdot \tau + 2\gamma(\varepsilon(\mathbf{u}_S) \cdot \mathbf{n}) \cdot \tau = 0.$$

For  $\gamma = 0$ , these conditions arise naturally when considering the generalized Stokes 165 problem (1) with discontinuous coefficients. In the case  $\gamma \neq 0$  the third Beaver-Joseph-Saffman condition, which restricts the tangential components of the normal 167 stresses, gives rise to an additional interface term that has to be included in the definition of the bilinear form  $\mathbf{a}_h$ ; for details see [8] and the references given there.

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<sup>&</sup>lt;sup>3</sup> visit: http://sourceforge.net/apps/mediawiki/ngsolve

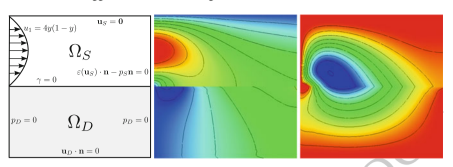


Fig. 1. From left to right: problem setup, and isolines of x- and y-components of the velocity for parameters  $\mu_S = 1$ ,  $\sigma_S = 0$  and  $\mu_D = 0$ ,  $\sigma_D = 1$ ;  $\gamma = 0$ . A part of the flow soaks through the porous medium. The normal component of the velocity is (almost) continuous across the interface, while no continuity is obtained for the tangential component

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# A Parallel Monolithic Domain Decomposition Method for Blood Flow Simulations in 3D

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Summary. We develop a parallel scalable domain decomposition method for the simulation 9 of blood flows in compliant arteries in 3D, by using a fully coupled system of linear elasticity 10 equation and incompressible Navier-Stokes equations. The system is discretized with a finite 11 element method on unstructured moving meshes and solved by a Newton-Krylov algorithm 12 preconditioned with an overlapping additive Schwarz method. We focus on the accuracy and 13 parallel scalability of the algorithm, and report the parallel performance and robustness of the 14 proposed approach by some numerical experiments carried out on a supercomputer with a 15 large number of processors and for problems with millions of unknowns.

1 Introduction 17

Computer modeling of fluid-structure interaction (FSI) is a useful tool for the study 18 of hemodynamics of blood flows in human arteries. Accurate modeling helps the pre- 19 diction and treatment of, for example, vascular diseases. FSI problems are in general 20 difficult to study. One of the main challenges is the effective coupling of the fluid 21 and the structure. Two well-known formulations are iterative and monolithic. In iter- 22 ative approaches, the fluid and the structure equations are solved one after the other 23 repeatedly, until some desired tolerance is reached [7, 10]. The convergence of these 24 approaches is difficult to achieve in some situations [6], since the approaches are very 25 similar to nonlinear Gauss-Seidel with two large blocks. In contrast, we develop a 26 monolithic coupling similar to [2-4], where the fluid and the structure equations are 27 solved simultaneously in a fully coupled fashion and the coupling conditions are en- 28 forced strongly as part of the system. The monolithic approach has been shown to 29 be more robust. Many of the convergence problems encountered within the iterative 30 approaches can be avoided.

With the rapid advancement in high performance computing technologies, high 32 resolution blood flow simulations are expected to provide more details of the physics 33 of blood flows and the artery walls. To obtain highly accurate solutions on a very fine 34 mesh, the parallel performance and scalability of the solution algorithm is becoming 35

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a key issue in the simulation. In [2, 3], a class of parallel scalable Newton-Krylov-Schwarz method was introduced for FSI in 2D. In this paper, we focus on solving 37 the fully coupled FSI system in 3D and also discuss the parallel performance and 38 robustness of the algorithms. The rest of the paper is organized as follows. In Sect. 2, 39 we describe the formulation and the discretization of the fully coupled FSI problem. In Sect. 3, we present the Newton-Krylov-Schwarz method for solving the fully 41 coupled nonlinear system. In Sect. 4, we first validate the method by comparing solutions obtained with the new approach with published results for a straight cylinder 43 problem, then report the parallel performance of the algorithm. Finally, we provide 44 some concluding remarks in Sect. 5.

## 2 Mathematical Formulation and Discretization

Our fully coupled approach can be described by the coupling of three components, 47 the linear elasticity equation for the wall structure in the reference Lagrangian frame, 48 the incompressible Navier-Stokes equations for the fluid in the arbitrary Lagrangian-49 Eulerian (ALE) framework, and the Laplace equation for the displacement of the 50 fluid domain.

Let  $\Omega_s \in \mathbb{R}^3$  be the structure domain. The displacement  $\mathbf{x}_s$  of the artery walls is 52 described by

$$\rho_s \frac{\partial^2 \mathbf{x}_s}{\partial t^2} - \nabla \cdot \sigma_s = \mathbf{f}_s \quad \text{in } \Omega_s, \tag{1}$$

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where  $\rho_s$  is the density of the structure, and  $\sigma_s = \lambda_s(\nabla \cdot \mathbf{x}_s)I + \mu_s(\nabla \mathbf{x}_s + \nabla \mathbf{x}_s^T)$  54 is the Cauchy stress tensor. The Lamé parameters  $\lambda_s$  and  $\mu_s$  are related to the 55 Young's modulus E and the Poisson ratio  $v_s$  by  $\lambda_s = v_s E/((1+v_s)(1-2v_s))$  and 56  $\mu_s = E/(2(1+v_s))$ . We fix the structure displacement  $\mathbf{x}_s = 0$  on the inlet and outlet 57 boundary  $\Gamma_s$ , and apply the zero normal traction condition  $\sigma_s \cdot \mathbf{n} = 0$  on the external 58 boundaries.

In order to model the fluid in a moving domain  $\Omega_f(t) \in \mathbb{R}^3$ , the displacement of 60 the fluid domain  $\mathbf{x}_f$  in the reference configuration  $\Omega_0 \in \mathbb{R}^3$  is assumed to satisfy a 61 Laplace equation, 62

$$\Delta \mathbf{x}_f = 0$$
 in  $\Omega_0$ .

We define an ALE mapping  $A_t$  from  $\Omega_0$  to  $\Omega_f(t)$ :

$$A_t: \Omega_0 \to \Omega_f(t), \quad A_t(\mathbf{Y}) = \mathbf{Y} + \mathbf{x_f}(\mathbf{Y}), \quad \forall \mathbf{Y} \in \Omega_0,$$

where **Y** is referred to as the ALE coordinates. The incompressible Navier-Stokes 64 equations defined on the moving domain  $\Omega_f(t)$  are written in the ALE form as 65

$$\rho_f \frac{\partial \mathbf{u}_f}{\partial t} \bigg|_{\mathbf{Y}} + \rho_f [(\mathbf{u}_f - \omega_g) \cdot \nabla] \mathbf{u}_f = \nabla \cdot \sigma_f \qquad \text{in } \Omega_f(t),$$

$$\nabla \cdot \mathbf{u}_f = 0 \qquad \text{in } \Omega_f(t),$$

where  $\rho_f$  is the fluid density,  $\mathbf{u}_f$  is the fluid velocity, and  $\sigma_f = -p_f I + \mu_f (\nabla \mathbf{u_f} + 66$  $\nabla {f u_f}^T$ ) is the Cauchy stress tensor.  $\omega_g = \partial {f x_f}/\partial t$  is the velocity of the moving domain 67 and Y indicates that the time derivative is taken with respect to the ALE coordi- 68 nates. On the inlet boundary  $\Gamma_i$ , a given velocity profile is prescribed. On the outlet 69 boundary  $\Gamma_o$ , the zero traction condition  $\sigma_f \cdot \mathbf{n} = 0$  is considered, where  $\mathbf{n}$  is the unit 70 outward normal. These boundary conditions may be chosen differently, depending 71 on the problem at hand.

More importantly, three coupling conditions are strongly enforced on the fluid-73 structure interface  $\Gamma_w$ 

$$\sigma_s \cdot \mathbf{n}_s = -\sigma_f \cdot \mathbf{n}_f, \ \mathbf{u_f} = \frac{\partial \mathbf{x_s}}{\partial t}, \ \mathbf{x}_f = \mathbf{x}_s,$$
 (2)

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where  $\mathbf{n}_s$ ,  $\mathbf{n}_f$  are unit normal vectors on the fluid-structure interface.

By introducing the structure velocity  $\dot{\mathbf{x}}_{\mathbf{s}}$  as an additional unknown variable, we 76 can rewrite the structure momentum equation (1) as a first-order system of equations. 77 We define the variational space of the structure problem as 78

$$X = \left\{ \mathbf{x_s} \in [H^1(\Omega_s)]^3 : \mathbf{x_s} = 0 \text{ on } \Gamma_s \right\}.$$
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The weak form of the structure problem is stated as follows: Find  $\mathbf{x_s} \in X$  and  $\dot{\mathbf{x}_s} \in X$  80 such that  $\forall \phi_s \in X$  and  $\forall \phi_s \in X$ , 81

$$B_{s}(\{\mathbf{x}_{s},\dot{\mathbf{x}}_{s}\},\{\phi_{s},\phi_{s}\};\sigma_{f}) = \rho_{s}\frac{\partial}{\partial t}\int_{\Omega_{s}}\dot{\mathbf{x}}_{s}\cdot\phi_{s}\,d\Omega + \int_{\Omega_{s}}\nabla\phi_{s}:\sigma_{s}\,d\Omega - \int_{\Gamma_{w}}\phi_{s}\cdot(\sigma_{f}\cdot\mathbf{n}_{s})\,ds - \int_{\Omega_{s}}\mathbf{f}_{s}\cdot\phi_{s}\,d\Omega + \int_{\Omega_{s}}\left(\frac{\partial\mathbf{x}_{s}}{\partial t}-\dot{\mathbf{x}}_{s}\right)\cdot\varphi_{s}\,d\Omega = 0.$$

The variational spaces of the fluid subproblem are time dependent, and the so- 82 lution of the structure subproblem provides an essential boundary condition for the 83 fluid subproblem by (2). We define the trial and weighting function spaces as:

$$V = \left\{ \mathbf{u}_f \in [H^1(\Omega_f(t))]^3 : \mathbf{u}_f = g \text{ on } \Gamma_i, \mathbf{u}_f = \partial \mathbf{x}_s / \partial t \text{ on } \Gamma_w \right\},$$

$$V_0 = \left\{ \mathbf{u}_f \in [H^1(\Omega_f(t))]^3 : \mathbf{u}_f = 0 \text{ on } \Gamma_i \cup \Gamma_w \right\},$$

$$P = L^2(\Omega_f(t)).$$

The weak form of the fluid problem reads: Find  $\mathbf{u}_f \in V$  and  $p_f \in P$  such that  $\forall \phi_f \in V_0$  85 and  $\forall \psi_f \in P$ ,

$$\begin{split} B_f(\{\mathbf{u}_f, p_f\}, \{\phi_f, \psi_f\}; \mathbf{x}_f) &= \rho_f \int_{\Omega_f(t)} \frac{\partial \mathbf{u}_f}{\partial t} \bigg|_{\mathbf{Y}} \cdot \phi_f \, d\Omega - \int_{\Omega_f(t)} p_f(\nabla \cdot \phi_f) \, d\Omega \\ &+ \rho_f \int_{\Omega_f(t)} \left[ (\mathbf{u}_f - \omega_g) \cdot \nabla \right] \mathbf{u}_f \cdot \phi_f \, d\Omega + 2\mu_f \int_{\Omega_f(t)} \varepsilon(\mathbf{u}_f) : \varepsilon(\phi_f) \, d\Omega \\ &+ \int_{\Omega_f(t)} (\nabla \cdot \mathbf{u}_f) \psi_f \, d\Omega = 0, \end{split}$$

where  $\varepsilon(\mathbf{u}_f) = (\nabla \mathbf{u}_f + \nabla \mathbf{u}_f^T)/2$ .

The weak form of the domain movement problem reads: Find  $\mathbf{x}_f \in Z$  such that 88  $\forall \xi \in Z_0$ ,

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$$B_m(\mathbf{x}_f, \xi) = \int_{\Omega_0} \nabla \xi : \nabla \mathbf{x}_f \, d\Omega = 0.$$

And the variational spaces are defined as

$$Z_0 = \{ \mathbf{x}_f \in [H^1(\Omega_0)]^3 : \mathbf{x}_f = 0 \text{ on } \Gamma_i \cup \Gamma_o \cup \Gamma_w \},$$
  

$$Z = \{ \mathbf{x}_f \in [H^1(\Omega_0)]^3 : \mathbf{x}_f = \mathbf{x}_s \text{ on } \Gamma_w, \mathbf{x}_f = 0 \text{ on } \Gamma_i \cup \Gamma_o \}.$$

We discretize the fully coupled problem in space with a finite element method, 91 by using unstructured P1-P1 stabilized elements for the fluid, P1 elements for the 92 structure and P1 elements for the fluid domain motion. We denote the finite element 93 subspaces  $X_h$ ,  $V_h$ ,  $V_{h,0}$ ,  $P_h$ ,  $Z_h$ ,  $Z_{h,0}$  as the counterparts of their infinite dimensional 94 subspaces. Because the fluid problem requires that the pair  $V_h$  and  $P_h$  satisfy the LBB 95 inf-sup condition, additional SUPG stabilization terms are needed in the formulation 96 with equal-order interpolation of the velocity and the pressure as described in [11, 97 12]. The semi-discrete stabilized finite element formulation for the fluid problem 98 reads as follows: Find  $\mathbf{u}_f \in V_h$  and  $p_f \in P_h$ , such that  $\forall \phi_f \in V_{h,0}$  and  $\forall \psi_f \in P_h$ , 99

$$B(\{\mathbf{u}_f, p_f\}, \{\phi_f, \psi_f\}; \mathbf{x}_f) = 0$$

with 100

$$B\left(\left\{\mathbf{u}_{f},p_{f}\right\},\left\{\phi_{f},\psi_{f}\right\};\mathbf{x}_{f}\right)=0,$$

$$B\left(\left\{\mathbf{u}_{f},p_{f}\right\},\left\{\phi_{f},\psi_{f}\right\};\mathbf{x}_{f}\right)$$

$$=B_{f}\left(\left\{\mathbf{u}_{f},p_{f}\right\},\left\{\phi_{f},\psi_{f}\right\};\mathbf{x}_{f}\right)+\sum_{K\in\mathscr{T}_{f}^{h}}\left(\nabla\cdot\mathbf{u}_{f},\tau_{c}\nabla\cdot\phi_{f}\right)_{K}$$

$$+\sum_{K\in\mathscr{T}_{f}^{h}}\left(\frac{\partial\mathbf{u}_{f}}{\partial t}\Big|_{\mathbf{Y}}+\left(\mathbf{u}_{f}-\omega_{g}\right)\cdot\nabla\mathbf{u}_{f}+\nabla p_{f},\tau_{m}\left(\left(\mathbf{u}_{f}-\omega_{g}\right)\cdot\nabla\phi_{f}+\nabla\psi_{f}\right)\right)_{K},$$

where  $\mathscr{T}_f^h = \{K\}$  is the given unstructured tetrahedral fluid mesh, and  $\tau_c$  and  $\tau_m$  are stabilization parameters. 102

We form the finite dimensional fully coupled FSI problem as follows: Find  $x_s \in$  $X_h, \dot{x}_s \in X_h, u_f \in V_h, p_f \in P_h \text{ and } x_f \in Z_h \text{ such that } \forall \phi_s \in X_h, \forall \phi_s \in X_h, \forall \phi_f \in V_{h,0},$ 104  $\forall \psi_f \in P_h$ , and  $\forall \xi \in Z_{h,0}$ , 105

$$B_s(\{x_s, \dot{x}_s\}, \{\phi_s, \varphi_s\}; \sigma_f) + B(\{u_f, p_f\}, \{\phi_f, \psi_f\}; x_f) + B_m(x_f, \xi) = 0.$$
 (3)

The system (3) is further discretized in time with a second-order BDF2 scheme. 106 Since the temporal discretization scheme is fully implicit, at each time step, we obtain the solution  $x^n$  at the nth time step from the previous two time steps by solving 108 a sparse, nonlinear algebraic system

$$\mathcal{F}_n(x^n) = 0, (4)$$

114

133

144

where  $x^n$  corresponds to the nodal values of the fluid velocity  $\mathbf{u}_f$ , the fluid pressure  $p_f$ , the fluid mesh displacement  $\mathbf{x}_f$ , the structure displacement  $\mathbf{x}_s$  and the structure velocity  $\dot{\mathbf{x}}_s$  at the *n*th time step. For simplicity, we ignore the script *n* for the rest of the paper.

### 3 Newton-Krylov-Schwarz Method

In the Newton-Krylov-Schwarz approach, the nonlinear system (4) is solved via the 115 inexact Newton method [8]. At each Newton step the new solution  $x^{(k+1)}$  is obtained 116 from the current solution  $x^{(k)}$  by  $x^{(k+1)} = x^{(k)} + \theta^{(k)} s^{(k)}$ , where the step length  $\theta^{(k)}$  is determined by a cubic line search technique. The Newton correction  $s^{(k)}$  is approximated by solving a preconditioned Jacobian system  $J_k M_k^{-1} M_k s^{(k)} = -\mathscr{F}(x^{(k)})$  with GMRES, where  $M_k^{-1}$  is a one-level restricted additive Schwarz preconditioner [5].

To define the domain decomposition preconditioner, we first partition the finite 121 element mesh (which consists of the meshes for all components of the coupled system) into non-overlapping subdomains  $\Omega^h_\ell,\ \ell=1,\ldots,N,$  where the number of subdomain N is always the same as the number of processors np. Then, each subdomain  $\Omega_\ell^h$  is extended to an overlapping subdomain  $\Omega_\ell^{h,\delta}$ . Note that the decomposition of 125 the mesh is completely independent of which physical variables are defined on a 126 given mesh point. The number of variables at a given mesh point is considered for 127 the purpose of load balancing. The so-called one-level restricted additive Schwarz 128 preconditioner is defined by

$$M_k^{-1} = \sum_{\ell=1}^N (R_\ell^0)^T J_\ell^{-1} R_\ell,$$

where  $R_\ell^0$  and  $R_\ell$  are restrictions to the degrees of freedom in the non-overlapping 130 subdomain  $\Omega_\ell^h$  and the overlapping subdomain  $\Omega_\ell^{h,\delta}$  , respectively.  $J_\ell$  is a restriction of the Jacobian matrix defined by  $J_\ell = R_\ell J_k R_\ell^T$ .

## 4 Numerical Results

Our algorithm is implemented using PETSc [1]. All computations are performed on 134 an IBM BlueGene/L supercomputer.

A benchmark 3D FSI problem is used to study the efficiency and performance 136 of our fully-coupled algorithm and software. The geometry consists of a straight 137 cylinder representing the fluid domain with length 5 cm and radius 0.5 cm, and 138 the surrounding structure with thickness 0.1 cm. A constant traction  $\sigma_f \cdot \mathbf{n} = 1.33 \cdot 139$ 10<sup>4</sup> dynes/cm<sup>2</sup> is imposed on the inlet boundary for 3 ms. A zero traction condition is applied to the fluid at the outlet boundary. The fluid is characterized 141 with viscosity  $\mu_f = 0.03$  poise, and density  $\rho_f = 1.0 \text{ g/cm}^3$ . The Young's modulus  $E = 3 \cdot 10^6 \,\mathrm{g/(cm\,s^2)}$ , the Poisson ratio  $v_s = 0.3$ , and the structure density 143  $\rho_s = 1.2 \text{ g/cm}^3$  are the parameters of the structure model.

The fluid and the structure are initially at rest and the simulation is run on a 145 mesh with  $2.41 \cdot 10^6$  elements and  $3.08 \cdot 10^6$  degrees of freedom, for a total time of 146 10 ms with a time step size  $\Delta t = 0.1$  ms. The simulation proceeds to the next time 147 step when the residual of the nonlinear system is less than  $10^{-6}$ . In Fig. 1, we show 148 the computed fluid pressure and the structure deformation at  $t = 2.5, 5.0, 10.0 \,\mathrm{ms}$ . Our 149 results are similar to the published results in [7, 9]. We observe that the pressure wave 150 propagates along the cylinder and reaches the end of the cylinder at t = 10.0 ms. The wall structure deforms in response to the propagation of the wall pressure, which is 152 a key feature of the fluid-structure interaction.

The strong scalability of the algorithm is presented in Table 1. The results show 154 superlinear scalability for a range of problem sizes and with up to 2,048 processors. It 155 is worth noting that the growth in GMRES iterations for large processor counts may 156 be a problem if we consider to solve the problem on a much larger mesh and with a larger number of processors. In those situations, one possible solution to improve 158 the scalability is the use of a multilevel preconditioner.

Our algorithm is quite robust with respect to physical parameters. In some FSI 160 methods, the convergence becomes difficult to achieve if the density of the fluid and 161 the structure are close to each other. According to Table 2, our solver performs quite 162 well for a wide range of fluid density and structure density.

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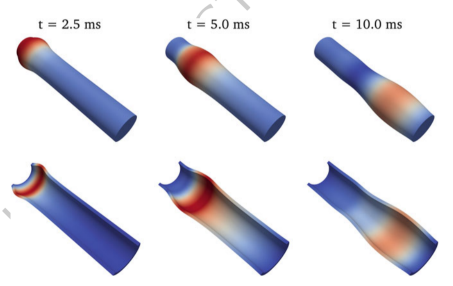


Fig. 1. Pressure wave propagation and structure deformation. The deformation is amplified by a factor of 12 for visualization purpose only

DOF	np	Newton	<b>GMRES</b>	time (s)
	256	2.0	41.60	218.03
$1.24 \cdot 10^6$	512	2.0	49.85	87.53
	1024	2.0	55.65	37.88
	512	2.0	57.60	442.44
$3.07 \cdot 10^6$	1024	2.0	67.15	152.16
	2048	2.0	77.55	65.64

**Table 1.** Performance with respect to the number of processors for two different mesh sizes. "np" denotes the number of processors. "Newton" denotes the average Newton iteration per time step. "GMRES" denotes the average GMRES iterations per Newton step. "time" refers to the average compute time, in seconds, per time step.

$\begin{array}{c ccccccccccccccccccccccccccccccccccc$					
1.0     1.0     2.0     49.85     87.53       1.0     10.0     2.0     53.90     88.07       1.0     100.0     2.0     61.75     88.84       0.01     1.0     2.0     124.60     96.75       0.1     1.0     2.0     60.90     88.77	$\rho_f$	$\rho_s$	Newton	<b>GMRES</b>	time (s)
1.0     10.0     2.0     53.90     88.07       1.0     100.0     2.0     61.75     88.84       0.01     1.0     2.0     124.60     96.75       0.1     1.0     2.0     60.90     88.77	1.0	0.1	2.0	71.65	89.94
1.0         100.0         2.0         61.75         88.84           0.01         1.0         2.0         124.60         96.75           0.1         1.0         2.0         60.90         88.77	1.0	1.0	2.0	49.85	87.53
0.01         1.0         2.0         124.60         96.75           0.1         1.0         2.0         60.90         88.77	1.0	10.0	2.0	53.90	88.07
0.1 1.0 2.0 60.90 88.77	1.0	100.0	2.0	61.75	88.84
	0.01	1.0	2.0	124.60	96.75
10.0 1.0 2.0 60.85 88.79	0.1	1.0	2.0	60.90	88.77
	10.0	1.0	2.0	60.85	88.79

**Table 2.** Different combinations of fluid density  $\rho_f$  and structure density  $\rho_s$ .  $\mu_f$  is kept at 0.03 poise. The tests are run for a problem with 1.25 · 10<sup>6</sup> unknowns and 512 processors.

5 Conclusion 164

In this paper, we developed and studied a parallel scalable overlapping Schwarz domain decomposition method for solving the fully coupled fluid-structure interaction system in 3D. Our algorithm is shown to be scalable on a large scale supercomputer and robust with respect to several important physical parameters. 168

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# A Fully Implicit Compressible Euler Solver for Atmospheric Flows \*

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1 Introduction 9

Numerical methods for global atmospheric modeling have been widely studied in 10 many literatures [5, 7, 9]. It is well-recognized that the global atmospheric flows can 11 be modeled by fully compressible Euler equations with almost no approximations 12 necessary [7]. However, due to the multi-scale nature of the global atmosphere and 13 the high cost of computation, other simplified models have been favorably used in 14 most community codes. 15

There are two main difficulties in using fully compressible Euler equations in 16 atmospheric flow simulations. One is that the fast waves in the equations lead to 17 very restrictive stability conditions for explicit time-stepping methods; see, e.g., [11]. 18 Another difficulty is that the flow is nearly compressible and the low Mach number 19 results in large numerical dissipation errors in many classical numerical schemes. 20

To deal with the fast acoustic and inertio-gravity waves in the fully compressible 21 model, we develop a fully implicit method so that the time step size is no longer 22 constrained by the stability condition. And to treat the low-Mach number flow, an 23 improved version of the Advection Upstream Splitting Method (AUSM<sup>+</sup>-up, [8]) is 24 adapted. This technique has been successfully employed for a shallow water model 25 in [12]. In the fully implicit solver, we use an inexact Newton method to solve the 26 nonlinear system arising at each time step; and the linear Jacobian system for each 27 Newton step is then solved by a Krylov subspace method with an additive Schwarz 28 preconditioner. We show by numerical experiments on a machine with thousands of 29 processors that the parallel Newton-Krylov-Schwarz approach works well for fully 30 compressible atmospheric flows.

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### **2** Governing Equations

Various formulations of the governing equations for mesoscale atmospheric models 33 can be found in, e.g., [6]. In this paper, we focus on the compressible Euler equations 34 by restricting the study on two dimensions (the x-z plane) and omitting the Coriolis 35 terms. The compressible Euler equations for the atmosphere take the following form 36

$$\frac{\partial Q}{\partial t} + \frac{\partial F}{\partial x} + \frac{\partial G}{\partial z} + S = 0,$$

where

$$Q = \begin{pmatrix} \rho \\ \rho u \\ \rho w \\ \rho \theta \end{pmatrix}, F = \begin{pmatrix} \rho u \\ \rho u^2 + p \\ \rho uw \\ \rho u\theta \end{pmatrix}, G = \begin{pmatrix} \rho w \\ \rho wu \\ \rho w^2 + p \\ \rho w\theta \end{pmatrix}, S = \begin{pmatrix} 0 \\ 0 \\ \rho g \\ 0 \end{pmatrix}, \tag{1}$$

where  $g=9.80665\,\mathrm{m/s^2}$  is the effective gravity on the surface of the Earth. In the 38 equation, the prognostic variables are the density  $\rho$ , the velocity (u,w) and the potential temperature  $\theta$  of the atmosphere. The system is closed with the equation of 40 state

 $p = p_{00} \left( \frac{\rho R \theta}{p_{00}} \right)^{\gamma},$ 

where  $p_{00}=1013.25$  hPa is the reference pressure on the surface, R=287.04 J/ 42 (kg·K) is the gas constant for dry air and  $\gamma=1.4$ . For the sake of brevity, we assume 43 the computational domain  $\Omega$  is a rectangle and the boundary conditions are given in 44 Sect. 5. In some cases, a physical dissipation is added to the left-hand-side of the 45 momentum and velocity equations. The dissipation term is  $-\nabla \cdot (v\rho \nabla \phi)$  for  $\phi=u$ , 46 w, and  $\theta$ .

To recover the hydrostatic solution from the equation, instead of using (1) directly, the following shifted system is often preferred [6, 11]:

$$Q = \begin{pmatrix} \rho' \\ \rho u \\ \rho w \\ (\rho \theta)' \end{pmatrix}, F = \begin{pmatrix} \rho u \\ \rho u^2 + p' \\ \rho u w \\ \rho u \theta \end{pmatrix}, G = \begin{pmatrix} \rho w \\ \rho w u \\ \rho w^2 + p' \\ \rho w \theta \end{pmatrix}, S = \begin{pmatrix} 0 \\ 0 \\ \rho' g \\ 0 \end{pmatrix}$$
(2)

where

$$ho'=
ho-ar
ho, \quad p'=p-ar
ho, \quad (
ho\, heta)'=
ho\, heta-ar
ho\,ar heta$$
 51

and the variables with 'bar' satisfy the hydrostatic condition  $\frac{\partial \bar{p}}{\partial z} = -\bar{p}g$  and  $\bar{\theta}$  is 52 obtained from the equation of state. It is clear that the flux Jacobian of the shifted 53 system (2) in each spatial direction is, respectively,

$$\frac{\partial F}{\partial Q} = \begin{pmatrix} 0 & 1 & 0 & 0 \\ -u^2 & 2u & 0 & c^2/\theta \\ -uw & w & u & 0 \\ -u\theta & \theta & 0 & u \end{pmatrix}, \quad \frac{\partial G}{\partial Q} = \begin{pmatrix} 0 & 0 & 1 & 0 \\ -wu & w & u & 0 \\ -w^2 & 0 & 2w & c^2/\theta \\ -w\theta & 0 & \theta & w \end{pmatrix},$$

where  $c = \sqrt{\gamma p/\rho}$  is the sound speed.

55

3 Discretizations 56

Suppose the computational domain is covered by a uniform rectangular  $N_x \times N_z$  57 mesh. Mesh cell  $\mathcal{C}_{ij}$  is centered at  $(x_i, z_j)$ , for  $i = 1, ..., N_x$  and  $j = 1, ..., N_z$ , with 58 mesh size  $\Delta x \times \Delta z$ . The solution in cell  $\mathcal{C}_{ij}$  at time t is approximated as

$$Q_{ij} \approx \frac{1}{\Delta x \Delta z} \int_{z_j - \Delta z/2}^{z_j + \Delta z/2} \int_{x_i - \Delta x/2}^{x_i + \Delta x/2} Q(x, z, t) dx dz.$$

We employ a cell-centered finite volume method for the spatial discretization of 60 the compressible Euler equations (2). Integrating (2) over  $\mathcal{C}_{ij}$  leads to the following semi-discrete system 62

$$\frac{\partial Q_{i,j}}{\partial t} + \frac{F_{i+1/2,j} - F_{i-1/2,j}}{\Delta x} + \frac{G_{i,j+1/2} - G_{i,j-1/2}}{\Delta z} + S(Q_{i,j}) = 0,$$

where the numerical fluxes of F and G are averaged on the edges of each mesh cell. 63

To calculate the numerical fluxes on cell edges, we first employ a piecewise linear 64
formulation to reconstruct constant states in both left and right direction, i.e., 65

$$\begin{split} &Q_{i+\frac{1}{2},j}^{-} = Q_{ij} + \frac{1}{4}(Q_{i+1,j} - Q_{i-1,j}), \quad Q_{i-\frac{1}{2},j}^{+} = Q_{ij} - \frac{1}{4}(Q_{i+1,j} - Q_{i-1,j}), \\ &Q_{i,j+\frac{1}{2}}^{-} = Q_{ij} + \frac{1}{4}(Q_{i,j+1} - Q_{i,j-1}), \quad Q_{i,j-\frac{1}{2}}^{+} = Q_{ij} - \frac{1}{4}(Q_{i,j+1} - Q_{i,j-1}). \end{split}$$

Then we use an improved version of the Advection Upstream Splitting Method 66 (AUSM<sup>+</sup>-up, [8]) to approximate the numerical fluxes based on the reconstructed 67 states. The basic idea of AUSM<sup>+</sup>-up scheme is to split the flux into two parts, e.g., 68

$$F = F^{(c)} + F^{(p)},$$
 69

where the convective flux  $F^{(c)} = \rho u(1, u, w, \theta)^T$  and the pressure flux  $F^{(p)} = 70$   $(0, p', 0, 0)^T$  are estimated separately, both in an upwinded manner. For instance, 71 denote the left and right reconstructed states for the prognostic variables on an edge 72 of a mesh cell as  $(\rho_-, u_-, w_-, \theta_-)$  and  $(\rho_+, u_+, w_+, \theta_+)$ , the pressure flux is approx-73 imated by  $F^{(p)} \approx (0, \widehat{p}', 0, 0)^T$ , where

$$\widetilde{p}' = \mathscr{P}_{5}^{+}(M_{-})p'_{-} + \mathscr{P}_{5}^{-}(M_{+})p'_{+} - (3/2)\mathscr{P}_{5}^{+}(M_{-})\mathscr{P}_{5}^{-}(M_{+})\widetilde{\rho}\,\widetilde{c}(u_{+} - u_{-}),$$

and 75

$$\begin{split} \widetilde{\rho} &= (\rho_{-} + \rho_{+})/2, \quad \widetilde{c} = (\sqrt{\gamma p_{+}/\rho_{+}} + \sqrt{\gamma p_{-}/\rho_{-}})/2, \quad p'_{\pm} = p_{\pm} - \bar{p}, \\ \mathscr{P}_{5}^{\pm}(M) &= \begin{cases} (1 \pm \text{sign}(M))/2, & \text{if } |M| \geq 1, \\ \mathscr{M}_{2}^{\pm}(M) \left[ (\pm 2 - M) \mp 3M \mathscr{M}_{2}^{\mp}(M) \right], & \text{otherwise}, \end{cases} \\ \mathscr{M}_{2}^{\pm}(M) &= (M \pm 1)^{2}/4, \quad M_{\pm} = u_{\pm}/\widetilde{c}. \end{split}$$

More details can be found in [8].

For the temporal integration, instead of using explicit methods that suffer from 77 severe stability restriction on the time step size, we employ a fully implicit method. 78 Given a semi-discrete system 79

$$\frac{\partial Q}{\partial t} + \mathcal{L}(Q) = 0,$$
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we use the following second-order backward differentiation formula (BDF-2):

$$\frac{1}{2\Delta t} \left( 3Q^{(k)} - 4Q^{(k-1)} + Q^{(k-2)} \right) + \mathcal{L}(Q^{(k)}) = 0.$$

Here  $Q^{(k)}$  denotes the solution vector Q evaluated at the k-th time step with a fixed 82 time step size  $\Delta t$ . Only at the first time step, a first-order backward Euler method is 83 used.

## 4 Newton-Krylov-Schwarz Solver

The fully implicit method leads to a large sparse nonlinear algebraic system at each 86 time step. In this study, we use the Newton-Krylov-Schwarz (NKS) algorithm as the 87 nonlinear solver. Given a nonlinear system  $\mathscr{F}(X) = 0$ , an inexact Newton method 88 is used to solve the system in the outer loop of the NKS approach. Let  $X_n$  be the 89 approximate solution for the *n*-th Newton iterate, we find the next solution  $X_{n+1}$  as

$$X_{n+1} = X_n + \lambda_n s_n, \quad n = 0, 1, \dots$$

where  $\lambda_n$  is the steplength decided by a linesearch procedure and  $s_n$  is the Newton 91 correction. We then use the right-preconditioned GMRES (restarted every 30 itera- 92 tions) method to solve the Jacobian system 93

$$J_n M^{-1}(Ms_n) = -\mathscr{F}(X_n), \quad J_n = \mathscr{F}'(X_n)$$
 94

until the linear residual  $r_n = J_n s_n + \mathcal{F}(X_n)$  satisfies

$$||r_n|| \leq \eta ||\mathscr{F}(X_n)||,$$
 96

where  $\eta > 0$  is the nonlinear forcing term that has been set to be a fixed value  $\eta = 97$  $1.0 \times 10^{-6}$  in our test. A multi-coloring finite difference method [4] is used to form 98 the Jacobian  $J_n$  in the calculation. To achieve uniform residual error at each time 99 step, we use the same adaptive stopping conditions as in [13]. 100

Given the computational domain  $\Omega$ , we first decompose it into non-overlapping 101 subdomains  $\Omega_k$ ,  $k=1,\ldots,np$ , where np is the number of subdomains and also the 102 number of processor cores. Then each subdomain  $\Omega_k$  is extended to  $\Omega_k^{\delta}$  within  $\Omega$ and the number of overlapping mesh layers between subdomains is  $\delta$ . For the overlapping domain decomposition, a preconditioner  $M^{-1}$  is then constructed using the 105 one-level restricted additive Schwarz (RAS, [2]) method defined as follows

$$M^{-1} = \sum_{k=1}^{np} (R_k^0)^T (J_n)_k^{-1} R_k^{\delta}.$$

Here  $(J_n)_k$  is the Jacobian matrix defined on subdomain  $\Omega_k^{\delta}$  and  $R_k^{\delta}$  and  $(R_k^0)^T$  are restriction and prolongation operators respectively. Given a solution vector defined on 108  $\Omega$ ,  $R_k^{\delta}$  restricts the vector to the overlapping subdomain  $\Omega_k^{\delta}$  while  $(R_k^0)^T$  prolongates 109 the restricted vector back to the whole domain  $\Omega$  by putting zeros not only outside 110  $\Omega_k^{\delta}$  but also within  $\Omega_k^{\delta} \setminus \Omega_k$ . In the implementation of the NKS solver, we use a 111 point-block ordering for both the unknowns and the nonlinear equations, resulting 112 in Jacobian matrices with 4 × 4-block entries. A point-block version of sparse LU 113 factorization is then used to solve the subdomain problems.

### **5 Numerical Results**

An IBM BlueGene/L supercomputer with 4,096 nodes is used to conduct our numerical tests. Each node of the computer has a dual-core IBM PowerPC 440 processor 117 running at 700 MHz and 512 MB local memory. We implement the NKS algorithm 118 based on the Portable, Extensible Toolkits for Scientific computations (PETSc, [1]) 119 library. In the numerical tests, the overlapping factor in the NKS solver is fixed at 120  $\delta = 2$ .

We study a test case describing a rising thermal bubble that is similar to those 122 studied in [3] and [10]. The computational domain is 123

$$\Omega = \{(x,z) | x \in [-10.0 \text{ km}, 10.0 \text{ km}], z \in [0,10.0 \text{ km}] \},$$

which is assumed to be horizontally periodic with rigid walls (zero normal velocity, 125 i.e., w = 0 here) at the bottom and top boundaries. The initial condition for the problem is obtained from a hydrostatic state with u = w = 0 and  $\bar{\theta} = 300$  K by adding a 127 perturbation

$$\Delta\theta = \begin{cases} 2.0\cos(0.5\pi L) \, \text{K} & \text{if } L \le 1.0, \\ 0.0 \, \text{K} & \text{otherwise,} \end{cases}$$

114

115

128

130

where

$$L = \sqrt{\left(\frac{x - 0.0 \,\mathrm{km}}{2.0 \,\mathrm{km}}\right)^2 + \left(\frac{z - 2.0 \,\mathrm{km}}{2.0 \,\mathrm{km}}\right)^2}.$$

A physical dissipation  $v = 15.0 \text{ m}^2/\text{s}$  is employed in the calculation. The results on a  $1,000 \times 500$  mesh using the fully implicit method with  $\Delta t = 2.0$  s are provided 133 in Fig. 1. We find that the results are in agreement with those provided in several 134 publications; see, e.g., [3, 10] and [6]. 135

To investigate the performance of the preconditioner, we run a fixed size problem 136 on a 1,920  $\times$  960 mesh for 50 time steps with  $\Delta t = 2.0$  s by using gradually doubled numbers of processor cores (np). The results on the averaged number of Newton and 138 GMRES iterations per time step are provided in Fig. 2, from where we observe that 139

### Potential temperature perturbation

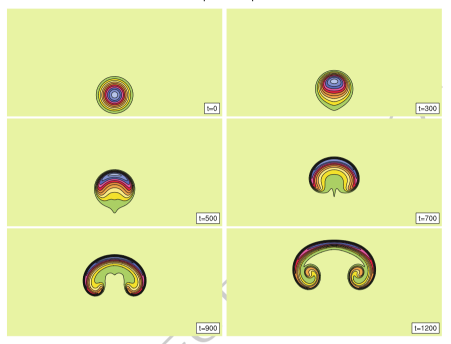


Fig. 1. Contour plots of the potential temperature perturbation (contour interval: 0.2 K)

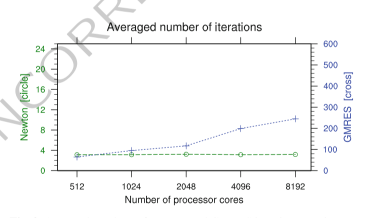


Fig. 2. Averaged numbers of Newton and GMRES iterations per time step

the number of Newton iterations is not sensitive to np but the number of GMRES iterations needed for each time step increases as np increases. The total compute time 141
and the parallel scalability are provided in Fig. 3, which clearly shows that as more 142
processors are used for the fixed size problem, the total compute time is reduced 143
accordingly and the parallel scalability from 512 to 8, 192 processor cores is nearly 144

#### A Fully Implicit Compressible Euler Solver for Atmospheric Flows

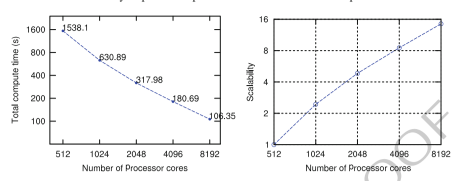


Fig. 3. Total compute time (*left*) and parallel scalability (*right*) results

optimal, with the parallel efficiency reaching 90.38%. Because of the page limit, 145 we only present a one-level restricted additive Schwarz method for the compressible 146 Euler problem and only provide some preliminary results in this paper. More ad- 147 vanced algorithms such as multilevel hybrid Schwarz methods will be investigated 148 in a forthcoming paper and more numerical experiments will be carried out in it.

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Preface

Domain decomposition, a form of divide and conquer for mathematical problems 3 posed over a physical domain, as in partial differential equations, is the most common paradigm for large-scale simulation on massively parallel distributed, hierarchi- 5 cal memory computers. In domain decomposition, a large problem is reduced to a 6 collection of (typically many) smaller problems, each of which is easier to solve computationally than the undecomposed problem and most or all of which can be solved 8 independently and concurrently. Typically, it is necessary to iterate over the collec- 9 tion of smaller problems, and much of the theoretical interest in domain decomposi- 10 tion algorithms lies in ensuring that the number of iterations required is very small. 11 Indeed, the best domain decomposition methods share with their cousins, multigrid 12 methods, the property that the total computational work is linearly proportional to the size of the input data or that the number of iterations required is at most logarithmic 14 in the number of degrees of freedom of individual subdomains. Algorithms whose 15 work requirements are linear in the size of the input data in this context are said to 16 be "optimal." Optimal domain decomposition algorithms are now known for many, 17 but certainly not all, important classes of problems that arise from science and engi- 18 neering. Much of the practical interest in domain decomposition algorithms lies in 19 extending the classes of problems for which optimal algorithms are known. Domain 20 decomposition algorithms can be tailored to the properties of the physical system 21 as reflected in the mathematical operators, the number of processors available, and 22 even to specific architectural parameters, such as cache size and the ratio of memory 23 bandwidth to floating-point processing rate.

Since the first meeting was held in Paris in 1987, the International Conference 25 on Domain Decomposition Methods is the only regularly occurring international forum dedicated to interdisciplinary technical interactions between theoreticians and 27 practitioners working in the creation, analysis, software implementation, and application of domain decomposition methods. The conferences have now been held in 29 12 countries in the Far East, Europe, the Middle East, and North America. To date, 30 there are essentially no real alternatives to domain decomposition as a strategy for 31 parallelization on petascale computers and beyond, with hundreds of thousands or 32 even millions of processor cores. Domain decomposition has proved to be an ideal 33

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paradigm not only for execution on advanced architecture computers but also for the 34 development of reusable, portable software. The most complex operation in a typical 35 domain decomposition method is the application of a preconditioner that carries out 36 in each subdomain step nearly identical to those required to apply a conventional pre- 37 conditioner to the global domain. Hence, software developed for the global problem 38 can readily be adapted to the local problem, instantly presenting wealth of "legacy" scientific code to be harvested for parallel implementations. Furthermore, since the 40 majority of data sharing between subdomains in domain decomposition codes oc- 41 curs in two archetypal communication operations – ghost point updates in overlap- 42 ping zones between neighboring subdomains and global reduction operations, as in 43 forming an inner product – domain decomposition methods map readily onto opti- 44 mized, standardized message-passing environments, such as MPL Finally, it should 45 be noted that domain decomposition is often a natural paradigm for the modeling 46 community. Physical systems are often decomposed into two or more contiguous 47 subdomains based on phenomenological considerations, such as the importance or 48 negligibility of viscosity or reactivity, or any other feature, and the subdomains are 49 discretized accordingly, as independent tasks. This physically based domain decomposition may be mirrored in the software engineering of the corresponding code, and 51 leads to threads of execution that operate on contiguous subdomain blocks, which 52 can either be further subdivided or aggregated to the granularity of an available par- 53 allel computer, and have the correct topological and mathematical characteristics for 54 scalability. Much of the reputation of this conference series results from the close 55 interaction between experts in mathematics, computer science, and large-scale computational science in various application areas.

This volume contains a selection of 83 papers presented at the 20th International 58 Conference on Domain Decomposition, DD20, hosted by the Center for Compu- 59 tational Mathematics at the University of California at San Diego, held at the San 60 Diego Supercomputer Center on the UCSD campus during the week of February 61 9–13, 2011. The conference featured 16 plenary lectures delivered by leaders in the 62 field, 18 minisymposiums, as well as contributed talks and a poster session. In ad- 63 dition, Olof Widlund gave an introductory short course on domain decomposition 64 on Sunday February 8 to a packed room of more than 40 participants in the Cen- 65 ter for Computational Mathematics, a short walk from the San Diego Supercom- 66 puter Center. Attending the regular conference during the week were 199 scientists 67 from 21 countries, giving a total of 173 presentations, which accentuates the international scope and relevance of this meeting. To add a unique local flavor to the 69 UCSD meeting, three special plenary talks were scheduled for Tuesday, given by 70 world-renowned local UCSD computational scientists in fields spanning computa-71 tional chemistry to galaxy collision simulation. In addition to the scientific talks dur- 72 ing the day throughout the week, participants gathered for a poster session with wine 73 and cheese in the early evening on Monday, and the plenary speakers gathered for a 74 small dinner in Del Mar on Tuesday evening. The Scientific Committee met with the 75 local organizing committee and discussed plans for the next conference in the series 76 on Wednesday evening, aided by samplings from local San Diego microbreweries. 77

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The large conference banquet for all the participants was held in the UCSD Faculty 78 Club on Thursday evening, and the conference came to a close at noon on Friday.

For further information, we recommend the homepage of International Domain 80 Decomposition Conferences, www.ddm.org, maintained by Martin Gander. This 81 site features free online access to the proceedings of all previous DD conferences, 82 information about past and future meetings, as well as bibliographic and personal 83 information pertaining to domain decomposition. A bibliography with all previous 84 proceedings is provided below, along with some major review articles and mono- 85 graphs. (We apologize for unintentional omissions to our necessarily incomplete 86 list.) No attempts have been made to supplement this list with the larger and closely 87 related literature of multigrid and general iterative methods, except for the books by 88 Hackbusch and Saad, which have significant domain decomposition components.

The editors wish to thank all members of the International Scientific Committee 90 for Domain Decomposition Conferences, chaired by Ralf Kornhuber, for their help 91 in setting the scientific direction of this conference. We are also grateful to the organizers of the minisymposiums for shaping the profile of the scientific program and 93 attracting high-quality presentations. The local organizers were Randolph Bank and 94 Michael Holst, aided by Rob Falgout, David Keyes, Rich Lehoucq, and Jinchao Xu. 95 We gratefully acknowledge administrative assistance from the San Diego Computer 96 Center (SDSC) and the California Institute for Telecommunications and Information 97 Technology (CalIT2).

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