

# **Solid Geometry Processing on Deconstructed Domains**

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# Abstract

Many tasks in geometry processing are modelled as variational problems solved numerically using the finite element method. For solid shapes, this requires a volumetric discretization, such as a boundary conforming tetrahedral mesh. Unfortunately, tetrahedral meshing remains an open challenge and existing methods either struggle to conform to complex boundary surfaces or require manual intervention to prevent failure. Rather than create a single volumetric mesh for the entire shape, we advocate for solid geometry processing on deconstructed domains, where a large and complex shape is composed of overlapping solid subdomains. As each smaller and simpler part is now easier to tetrahedralize, the question becomes how to account for overlaps during problem modelling and how to couple solutions on each subdomain together algebraically. We explore how and why previous coupling methods fail, and propose a method that couples solid domains only along their boundary surfaces. We demonstrate the superiority of this method through empirical convergence tests and qualitative applications to solid geometry processing on a variety of popular second-order and fourth-order partial differential equations.

Keywords: numerical analysis, methods and applications

ACM CCS  $\bullet$  Mathematics of computing  $\rightarrow$  Discretization; Partial differential equations; Numerical differentiation

## 1. Introduction

Many tasks in computer graphics and geometry processing can be modelled mathematically as solutions to partial differential equations (PDEs) over a compact spatial domain. For example, shapeaware scattered data interpolation can be modelled as a solution to the Laplace equation ( $\Delta u = 0$ ). Smooth detail-preserving shape deformations can be efficiently parametrized using solutions to a bi-Laplace equation ( $\Delta^2 u = 0$ ). Even computation of geodesic distances (Figure 1) can be captured via iterative solutions to a Poisson equation ( $\Delta u = f$ ). These applications — and many others — rely on *discretization* to realize their solutions on the complex shapes found throughout computer graphics. The most common discretization is via the finite element method (FEM) using piecewiselinear functions defined over a simplicial mesh. For problems over solid regions in  $\mathbb{R}^3$ , this typically requires constructing a tetrahedral mesh that fills the volume bounded by a given surface. Compared to regular grids, unstructured tetrahedral meshes afford spatially varying resolution and complex boundary surfaces — *in theory, at least.* 

In practice, constructing tetrahedral meshes is a fragile process. While the application of linear FEM is often straightforward after posing a problem in the smooth setting, the actual creation of a valid tetrahedral mesh inside a triangle mesh is often left to an *ad hoc* patchwork of heuristics including manual intervention and mesh repair. Existing automatic meshing methods fall short. They either fail too often, create too poor quality elements or approximate too loosely the input domain boundary, as shown in Figure 2.

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**Figure 1:** A complex shape is constructed as the union of 32 solid components. Rather than resolving the mesh union geometrically and struggling to tetrahedralize the result, we propose tetrahedralizing each smaller, simpler model independently. We introduce constraints to join the volume meshes algebraically, enabling partial differential equation-based geometry processing (heat distances [CWW13] shown here).



**Figure 2:** TETGEN fails to tetrahedralize this microscope, even after preprocessing the input using [ZGZJ16]. QUARTET successfully outputs a tet mesh, but thin parts are poorly approximated (inset) and close features are merged (arrow).

We consider an interesting class of shapes that are — at least conceptually if not literally — described as the union of simpler domains (see inset). The traditional



conforming tetrahedralization pipeline would proceed by first computing the result of a surface mesh union operation and then attempt to mesh the interior. However, even if the input triangle meshes are 'clean', the exact mesh boolean result may be host to a number of issues that trip up available tetrahedralization heuristics: fine features, small voids and poorly shaped elements (Figure 3). In general, the *exact* result introduces many new vertices whose coordinates are rational numbers. Naively rounding such vertices to floating-point coordinates may introduce self-intersections, and efficient rounding while preventing intersections in 3D is still an open problem [For97].



**Figure 3:** Even if all input meshes are 'clean' (e.g. four overlapping geodesic spheres on left), their exact mesh-union may have arbitrarily poor quality triangles. These in turn trip up conforming Delaunay tet-meshers: e.g. TetGen [Si03] fails on this example.

In this paper, we propose an alternative to this error-prone pipeline. We consider inputs as *deconstructed domains*, composed as the union of any number of simpler shapes. We tetrahedralize sub-domains independently with no requirement to share vertex positions or combinatorics. Sub-domains are treated democratically, without a priority ordering or hierarchy. We then couple discrete PDEs or variational problems defined on each domain *algebraically*.

This coupling requires care. The given PDE or variational problem must be adapted for overlapping domains to avoid bias or double-counting in twice-covered regions. Via null-space analysis, we show that naive equality constraint leads to *locking*, artificial error that does not vanish under resolution refinement.

We borrow ideas from domain decomposition and immersed boundary methods to derive a general-purpose boundary-only coupling in the smooth setting and then demonstrate its effectiveness for linear FEM discretizations of common problems in 1D, 2D and 3D (e.g. Poisson equations). Domains are coupled with hard constraints resulting in a parameterless method. Further, we extend our results to boundary-only higher order coupling for applications of mixed FEM for fourth-order problems (e.g. bi-Laplace equation).

# 2. Related Work

Our goal is to improve the robustness of geometry processing that requires solving PDEs on solid shapes.

**PDE solvers in geometry processing** Improving the accuracy, robustness and performance of solvers for geometry problems is a core area of interest [BBK05, KFS13, Kaz15, DMZ\*17, HDA17, SPSH\*17]. While we focus on robustness with respect to the input representation, our work relates to these as an *algebraic* pre-process or filter on the eventual linear system or optimization problem. This is in contrast to geometric approaches to robustness such as remeshing [BK04b, SRUL16, GJTP17] or enclosing a shape in a cage [JMD\*07, SVJ15]. The boundary element method avoids volumetric meshing altogether (e.g. [JP99, DHB\*16, SVB17]), however, also limits the class of problems that can be solved. Compared to methods tailored to one application (e.g. character skinning [BTST12]), our method applies to a general class of PDEs.

**Constructive solid geometry** Emerging technologies such as 3D printing and virtual reality have ignited broader interest in geometric modelling. The result is that we have a huge amount of geometric data, but that data are rarely composed of a single, watertight, non-self-intersecting, oriented manifold surface [ZJ16]. Instead, people create using constructive solid geometry (CSG) tools like OPEN-SCAD and TINKERCAD that happily allow overlapping simpler models to create a larger, more complex shape. Early digital CSG complements this modelling paradigm with fast evaluation using implicit functions (e.g. [WMW86]) and GPU-friendly rendering [GHF86]. While the complexity of available meshed surface geometry grows, researchers have devised interesting and interactive ways to create complex models using pre-existing detailed parts [GSP\*06, CK10, CKGK11].

Most volumetric solvers do not consider the upstream modelling process and instead require a single volumetric mesh as input. This puts a heavy burden on modelling tools to maintain a clean surface geometry via mesh 'surgery' operations [SBSCO06, SS10a, SS10b]. Despite recent progress on robust boolean operations for triangle meshes [BF09, BGF15, ZGZJ16], the resulting meshes may have arbitrarily poor aspect ratio (Figure 3) preventing or damaging tetrahedralization. In contrast, we operate directly on the overlapping sub-domain representation common to solid modelling.

**Tetrahedral meshing** Although our work is an effort to subvert tetrahedral meshing and its issues, we still rely heavily on the progress and open source software from this literature. The fundamental challenge at the core of tetrahedral meshing is the balance between ensuring high-quality elements (see, e.g. [She02]) and conforming to a given input surface. While a complete meshing survey is outside the scope of this paper, we identify issues and previous works as they relate to our setting.



Figure 4: This seemingly innocuous union (left) causes the conforming Delaunay method TETGEN [Si03] to fail. Implicit or querybased methods have trouble capturing fine structures (middle) without resorting to high resolutions (right).

Conforming Delaunay tetrahedralization methods maintains the input geometry (and combinatorics) exactly by inserting input vertices and faces into a Delaunay triangulation and then improving element quality via local operations and additional Steiner vertices [CDS12]. In practice, the software TETGEN [Si03] implements many state-of-the-art algorithms and heuristics. The success rate of TET-GEN is not 100% (Figure 4), but it succeeds far more often when inputs are smaller and simpler, without spatially close parts or small triangles. We use TETGEN in most of our examples, but run it on each sub-domain, rather than the complex, complete shape.

Alternatively, other meshing methods work by employing a background grid [LS07] or implicit representation of the input shape [ACSYD05]. These methods ensure good quality elements by construction, but struggle to closely approximate the input shape geometry — especially in the presence of sharp features. Doran *et al.* [DCB13] provide an open source implementation, QUARTET and while robust in the sense of successfully outputting a mesh, this method will join together close features and fail to resolve thin parts (Figure 2). In contrast to Cuilliere *et al.* [CFD12], we avoid computing a unified mesh and do not require matching or correspondence between vertices or combinatorics of overlapping meshes.

**Domain decomposition** The idea of coupling solutions to PDEs across overlapping domains is quite old [Sch70] and well studied. The majority of previous methods for overset and non-matching grids focus on domain decomposition for parallel, offline computation using iterative solvers [SBGG04]. Alternatively, *immersed boundary methods* [Pes73] use similar constraints to couple the simulation of one or many objects embedded in a background simulation. For example, coupling a floating elastic body to a fluid simulation (e.g. [GSLF05]).

In contrast, our interest is in reducing the burden of tetrahedralization while maintaining the complexity of shapes found in graphics and geometry processing. We treat coupling as a hard constraint to single linear system solved using modern, large sparse linear solvers (e.g. [Dav06, AA00]). No sub-domain has preference over another.

English *et al.* [EQYF13] simulate water at varying resolutions by allowing regular finite-difference grids to rigidly overlap. Their method assigns priorities to grids and stitches higher priority grids along their boundaries into lower priority grids to solve a Poisson equation. Similar so-called *Chimera grids* [Ben85] are found in early fluid simulations on comparatively simple domains [BSD83, Hen94, KKRC97, DMYN08]. Henshaw describes how boundary values of one grid are interpolated using ghost points. This method is applied to overlapping regular Cartesian or polar grids. Malgat *et al.* [MGL\*15] couple overlapping discretizations for elasticity simulation via energy minimization. Their method requires a hierarchical ordering.

Overset grid methods (e.g. [Nak99, LSLR01, BS15]) often assume that the domain has been designed with an overlapping solver in mind. High-resolution grids near important areas naturally have well-defined and known priority over coarse background grids. Instead, we consider the case where sub-domain priorities are not known and domains merely serve as an overlapping sub-division. The resolution of a single grid may itself be adaptive.

Schwarz domain decomposition can be interpreted in the context of discontinuous Galerkin finite element method or extended FEM [Kau12], where sub-domains are interpreted as large, high-degree elements and coupling is analogous to interface conditions. Edwards and Bridson [EB15] propose such a solver for Poisson, elasticity and bi-Laplace problems. Their overlapping sub-domains are extracted from a unified grid of the entire domain.

## 3. Smooth Foundations

We first consider a PDE involving a smooth function u defined over a volumetric (i.e. co-dimension zero) domain  $\Omega \subset \mathbb{R}^d$  with appropriate boundary conditions applied to u on the boundary of the domain  $\partial \Omega$ . We focus specifically on elliptic PDEs resulting from energy minimizations common in geometry processing. For example, minimizing the squared gradient (i.e. Dirichlet energy) minus a unit potential, subject to fixing the value of u to a known function g on the boundary of the domain,

$$\min_{u} \int_{\Omega} \left( \frac{1}{2} \| \nabla u \|^2 - u \right) \, dA \tag{1}$$

subject to 
$$u(\mathbf{x}) = g(\mathbf{x}) \quad \forall \mathbf{x} \in \partial \Omega,$$
 (2)

results in the second-order Poisson equation on the interior,

$$\Delta u = 1 \quad \forall \mathbf{x} \in \Omega. \tag{3}$$

Suppose we are incapable of measuring an energy directly over all of the domain  $\Omega$ , but instead are only able to measure energies over two *overlapping* sub-domains  $\Omega_1$ ,  $\Omega_2 \subset \mathbb{R}^d$  whose union comp-



oses the original domain  $\Omega_1 \cup \Omega_2 = \Omega$ . By replacing *u* with new  $u_1$  and  $u_2$  over each respective sub-domain, we can write the original minimization problem in Equation (1), breaking the integral into the non-overlapping parts in each sub-domain  $\Omega_1 \setminus \Omega_2$  and  $\Omega_2 \setminus \Omega_1$  and their intersection ( $\Omega_1 \cap \Omega_2$ ) and adding a *pointwise* equality

coupling constraint,

$$\min_{u_1,u_2} \int_{\Omega_1 \setminus \Omega_2} \left( \frac{1}{2} \| \nabla u_1 \|^2 - u_1 \right) \, dA + \tag{4}$$

$$+\int_{\Omega_2 \setminus \Omega_1} \left(\frac{1}{2} \|\nabla u_2\|^2 - u_2\right) \, dA + \tag{5}$$

$$+\frac{1}{2}\int_{\Omega_1\cap\Omega_2} \left(\frac{1}{2}\|\nabla u_1\|^2 - u_1 + \frac{1}{2}\|\nabla u_2\|^2 - u_2\right) dA \qquad (6)$$

subject to 
$$u_1(\mathbf{x}) = g(\mathbf{x}) \quad \forall \mathbf{x} \in \partial \Omega \cap \partial \Omega_1,$$
 (7)

and 
$$u_2(\mathbf{x}) = g(\mathbf{x}) \quad \forall \mathbf{x} \in \partial \Omega \cap \partial \Omega_2,$$
 (8)

and 
$$u_1(\mathbf{x}) = u_2(\mathbf{x}) \quad \forall \mathbf{x} \in \Omega_1 \cap \Omega_2.$$
 (9)

Advantage of working with energies The appearance of the factor before the integrated energy in the intersection  $\Omega_1 \cap \Omega_2$  region (see Equation (6)) would not be so obvious if we had worked with the Poisson problem directly as a PDE (see Equation (3)). However, viewed as variational problem, the necessity of the 1/2 is clear: we should not double count the energy contributed in this region.

## 4. Discrete Locking

The deconstructed energy optimization problem in Equation (4) only involves first derivatives and linear equality constraints. It is tempting to jump to a FEM discretization using piecewise-linear elements for each sub-domain  $\Omega_1$  and  $\Omega_2$ , e.g. hat functions (over polylines for d = 1, triangle meshes for d = 2 and tetrahedral meshes in d = 3)

$$u_i(\mathbf{x}) = \sum_{j=1}^n u_{ij} \varphi_{ij}(\mathbf{x}), \tag{10}$$

with interpolated values at the *n* vertices given as a vector  $\mathbf{u}_i \in \mathbb{R}^n$ .

If the meshes over  $\Omega_1$  and  $\Omega_2$  have only and exactly coincident vertices and compatible combinatorics in the intersection region  $\Omega_1 \cap \Omega_2$ , then we call them *matching*. In this special case, enforcing the pointwise equality constraint in Equation (9) is equivalent to merging the meshes. The solution search space is exactly as rich as linear FEM over the merged mesh.

For meshes with vertices in general position, perfect coincidence never happens. Pointwise equality immediately reduces the search space to piecewise linear functions that exist mutually in both linear FEM function spaces over the intersection  $\Omega_1 \cap \Omega_2$ . In the general *non-matching* case, the constraint reduces the search space dramatically: *only* functions that take on a linear function over  $\Omega_1 \cap \Omega_2$ remain (see Figure 5, left).

This is an extreme case of what is known as *locking* in the FEM literature [ZT00]. Locking is an artificial stiffening of system during discretization. In our case, the constraints are so strict that only rather boring functions remain. These functions can be arbitrarily far from

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**Figure 5:** The constraint space is unrelated to the energy. Consider a simple reconstruction energy  $\min_u \int_{\Omega_1 \cup \Omega_2} (u - g)^2 dA$ . Constraining all points in the overlap locks to a linear function; all vertices look promising for saddle-shaped functions, but cannot reproduce positive curvature functions; boundary vertices avoid locking.

the desired solution and discretization refinement by adding more (general position) vertices will not help (Figure 6).

## 4.1. Constraints at all vertices cause locking

One immediate strategy is to require equality only at mesh vertices. This ties the values at one mesh's vertices to the piecewise linearly interpolated value on the other mesh via a linear equality constraint and *vice-versa*, e.g.:

$$u_{1i} = \sum_{j=1}^{n_2} u_{2j} \varphi_{2j}(\mathbf{v}_{1i}) \quad \forall i \text{ such that } \mathbf{v}_{1i} \in \Omega_1 \cap \Omega_2, \quad (11)$$

$$u_{2j} = \sum_{i=1}^{n} u_{1i} \varphi_{1i}(\mathbf{v}_{2j}) \quad \forall j \text{ such that } \mathbf{v}_{2j} \in \Omega_2 \cap \Omega_1, \quad (12)$$

where  $\mathbf{v}_{ki} \in \mathbb{R}^d$  is the position of the *i*th vertex in the mesh over sub-domain  $\Omega_k$ . The coefficients obtained by evaluating the hat functions  $\varphi_{\ell j}(\mathbf{v}_{ki})$  of the *other* mesh over sub-domain  $\Omega_{\ell}$  are simply the barycentric coordinates of  $\mathbf{v}_{ki}$  in the containing simplex (e.g. tetrahedron for d = 3).

We can collect these constraints in matrix form

$$\mathbf{C} \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} = \mathbf{0},\tag{13}$$

where  $\mathbf{C} \in \mathbb{R}^{(m_1+m_2)\times(n_1+n_2)}$  is a sparse rectangular matrix, where each row corresponds to one of the  $m_1$  vertices of the mesh over  $\Omega_1$  lying in  $\Omega_2$  or *vice-versa*.

These linear equality constraints are easy to implement in practice (e.g. via the null space or Lagrange multiplier method). Unfortunately, these constraints do not alleviate locking.

In  $\mathbb{R}^1$ , constraining all vertices in the overlapping region  $\Omega_1 \cap \Omega_2$ is catastrophic. Intuitively, if a segment of the mesh over  $\Omega_1$  overlaps with a segment of the mesh over  $\Omega_2$ , then both pairs of vertices will have to lie on the same line. In the worst case, an alternating order of vertices from  $\Omega_1$  and  $\Omega_2$  creates a *domino effect*, and the entire intersection region locks to the same linear function (Figure 7).

It is tempting to extrapolate that these constraints will always result in pointwise locking, but in higher dimensions (d > 1), locking from vertex constraints is more nuanced. We observe in Figure 5 that the constraint space created by coupling all vertices struggles to reproduce a round parabolic function and more easily reproduces a saddle-shaped hyperbolic function. Indeed, imposing this constraint when solving a Laplace equation (saddle-shaped solution), we see significantly better convergence with respect to mesh resolution than when solving a parabolic Poisson equation (Figure 8).

This is not a coincidence. The constraint matrix **C** in Equation (13) satisfies many desired properties (constant precision, linear precision, the maximum principle and local support; see [WMKG07]) of a *discrete Laplacian* on the 'joint mesh' over  $\Omega_1 \cap \Omega_2$  created by connecting each vertex of  $\Omega_1$  to the vertices of its containing simplex containing in  $\Omega_2$  and *vice-versa*. Performing eigen analysis on **C** reveals that it responds as a discrete operator strikingly similarly to the FEM discrete (cotangent) Laplacian (Figure 9). Because of this relationship, we call the artificial stiffening due to constraining all overlapping vertices *harmonic locking*.

We will defer our discussion of attempting to soften this equality constraint to Section 7 and instead return to the smooth setting to derive a locking-free solution from first principles.

# 5. Boundary-Only Coupling

The root of the locking troubles is the pointwise equality constraint over the overlapping region  $\Omega_1 \cap \Omega_2$  in Equation (9). Surely coupling is crucial. If we remove this constraint entirely, then  $u_1$  and  $u_2$ will solve independent Poisson equations, subject to emergent natural boundary conditions (in this case,  $\nabla u \cdot \mathbf{n} = 0$ ) on the overlap boundary  $\partial(\Omega_1 \cap \Omega_2)$ . In other words, these zero normal derivative boundary conditions uniquely determine  $u_1$  and  $u_2$ .

The fact that minimizers of our energy in Equation (1) are uniquely determined by boundary conditions can be spun to play in our favour when searching for non-locking coupling constraints.



**Figure 6:** Left-to-right: the Poisson solution in 1D is a simple parabola, easily approximated using linear finite element method. If the discrete domain is given as two overlapping meshes (red and yellow), then enforcing equality at all vertices in the overlap results in the solution locking to a linear function. This problem does not go away with mesh refinement. Coupling the meshes only at the sub-domain boundary (purple and green) alleviates locking, and converges with refinement.



Figure 7: Locking occurs at any scale and can be understand as a 'Domino effect' when enforcing constraints one-by-one. In 1D, consecutive constraints force the solution to a line: eventually, the entire overlapping region must be a single line.



**Figure 8:** Constraining all vertices in the overlap between two concentric annuli matches the convergence rate of our boundary-only constraints for  $\Delta z = 0$  (left). However, constraining all does not converge for  $\Delta z = 1$  (right), while boundary only maintains convergence. For reference: a non-overlapping, single mesh converges.

Concretely, we will now show that it is sufficient to restrict the pointwise equality constraints from the entire intersection region  $\Omega_1 \cap \Omega_2$  in Equation (9) to *only* its boundary  $\partial(\Omega_1 \cap \Omega_2)$ :

$$u_1(\mathbf{x}) = u_2(\mathbf{x}) \qquad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2).$$
 (14)

We must show that minimizing the deconstructed energy in Equation (4) over  $u_1$  and  $u_2$  with this constraint instead of Equation (9) remains equivalent to the minimization over u in Equation (1).

Assume that  $u_1$  and  $u_2$  are minimizers of Equation (4) satisfying  $u_1 = u_2|_{\Omega_1 \cap \Omega_2}$ , then by uniqueness of energy minimizers and equivalence with the energy in Equation (1),  $u = u_1|_{\Omega_1}$  and  $u = u_2|_{\Omega_2}$ . We must show that minimizing Equation (4) implies that  $u_1 = u_2|_{\Omega_1 \cap \Omega_2}$ .

Given minimizers  $u_1$  and  $u_2$  of Equation (4), let us define  $u_1 = u_2 := h|_{\partial(\Omega_1 \cap \Omega_2)}$ . It does not matter that we do not explicitly know the value of *h*. It is enough that it is well defined implicitly by solving the problem in Equation (4) subject to Equation (14). Since the minimizers  $u_1$  and  $u_2$  satisfy the Dirichlet conditions on their respective boundaries (Equations (7) and (8)), we can add the following constraints to Equation (4) without changing the minimum:

$$u_1(\mathbf{x}) = u_2(\mathbf{x}) = h(\mathbf{x}) \qquad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2).$$
(15)

Minimizers to our quadratic energy are uniquely determined by the values on the boundary of the domain, so we can isolate the problem



**Figure 9:** Eigen modes of the constraint matrix built from fixing all vertices of two overlapping, non-matching disc meshes resemble those of the discrete Laplacian. More irregular meshing produces less smooth modes.

for the overlapping region  $\Omega_1 \cap \Omega_2$ , for example:

$$\min_{u_1} \frac{1}{2} \int_{\Omega_1 \cap \Omega_2} \frac{1}{2} \|\nabla u_1\|^2 - u_1 \, dA \tag{16}$$

subject to 
$$u_1(\mathbf{x}) = h(\mathbf{x}) \quad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2),$$
 (17)

whose optimal argument is identical to the analogous problem replacing  $u_1$  with  $u_2$ , thus implying that the two functions agree on the overlapping region:  $u_1 = u_2|_{\Omega_1 \cap \Omega_2}$ .

Schwarz noticed this over a century ago [Sch70]. Since then, it has been exploited for domain decomposition for parallelization and memory decoupling for iterative solvers discussed in Section 2.

Analogous to the enforcement of Dirichlet boundary conditions, in the discrete linear FEM setting, we constrain only boundary vertices of  $\Omega_1$  lying inside the other domain  $\Omega_2$  or *vice-versa*:

$$u_{1i} = \sum_{j=1}^{n_2} u_{2j} \varphi_{2j}(\mathbf{v}_{1i}) \qquad \forall i \text{ such that } \mathbf{v}_{1i} \in \partial \Omega_1 \cap \Omega_2, \quad (18)$$

$$u_{2j} = \sum_{i=1}^{n_1} u_{1i} \varphi_{1i}(\mathbf{v}_{2j}) \qquad \forall j \text{ such that } \mathbf{v}_{2j} \in \partial \Omega_2 \cap \Omega_1.$$
(19)

These constraints are a subset of the rows of **C** in Equation (11), and we call this much smaller matrix  $\mathbf{A} \in \mathbb{R}^{(b_1+b_2)\times(n_1+n_2)}$ , where the mesh of  $\Omega_i$  has  $b_i$  overlap-boundary vertices.

Not only does fixing the boundary result in a smaller number of constraints and thus typically a better conditioned system,

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**Figure 10:** Constraining all vertices between two concentric spherical shells ('3D annuli'; cut view in middle) does not appear to converge for the Poisson equation  $\Delta u = 1$  (right), while boundary only exhibits similar convergence to a non-overlapping single mesh.

but also the discrete approximations are free of locking artefacts. We see this immediately in the 1D example in Figure 6. The boundary constraints do not show up visible in the constraint space when reproducing hyperbolic or parabolic functions Figure 5. In Figures 8 and 10, convergence with respect to mesh resolution for second-order problems roughly matches that of using a single unified mesh. Recall that we are purposely avoiding creating such a unified mesh, especially in  $\mathbb{R}^3$ , where mesh surgery and likely manual intervention and parameter tuning would be necessary. Instead, complex shapes can be created by overlapping many solid subdomains and coupling solutions using our proposed boundary-only constraints.

# 5.1. Multiple overlapping sub-domains

In general, a complex shape may be composed of the union of K > 1 sub-domains:

$$\Omega = \bigcup_{i=1}^{K} \Omega_i.$$
(20)

All of our derivations so far for K = 2 extend easily to K > 2. Our deconstructed energy has the form

$$\sum_{i=1} \int_{\Omega} \frac{1}{\sum_{j=1}^{K} \chi_j} \left( \frac{1}{2} \| \nabla u_i \|^2 - u_i \right) dA, \tag{21}$$

where  $\chi_j$  is the characteristic function of  $\Omega_j$  (i.e.  $\chi_j(\mathbf{x}) = 1$  for  $\mathbf{x} \in \Omega_j$  and = 0 otherwise). We defer the implementation details and matrix construction to Appendix A.

Many problems in geometry processing are slight variations on the minimization of this energy. For example, implicit timeintegrations of the wave equation replace the unit potential with acceleration, while the heat equation replaces this with a temperature field [SCV14]. While these changes to the basic Poisson solver here are nominal and left to the reader, increasing the differential order of the energy requires specific attention (see Section 6). Before this, we discuss two important considerations during discretization.



**Figure 11:** *Quadrature for estimating partial volume at boundary elements can increase convergence, with diminishing returns.* 

## 5.2. Quadrature

When discretizing the integral in Equation (21), we must approximate the partial volume of tetrahedra straddling the overlap boundaries. We compared various strategies. We specifically *avoid* computing this analytically or splitting elements as this is tantamount to the mesh boolean problem and would inherit its numerical challenges and robustness issues. Instead, we observe that numerical quadrature or Monte Carlo sampling improves accuracy and indeed helps convergence, albeit with diminishing returns (Figure 11). Approximating this integral is simpler than remeshing. We avoid computing exact intersections or new combinatorics.

Unless otherwise noted, we simply treat an element as fractionally inside or outside another mesh by averaging the number of domains each corner position lies within (i.e. first-order quadrature).

## 5.2.1. Constraint thinning

1

The simplest way to extend our boundary coupling constraints for K = 2 in Equation (18) is to consider all possible pairs of the K sub-domains:

$$u_{ai} = \sum_{j=1}^{n_b} u_{bj} \varphi_{bj}(\mathbf{v}_{ai}) \qquad \forall i, a \neq b \text{ s.t. } \mathbf{v}_{ai} \in \partial \Omega_a \cap \Omega_b.$$
(22)

For shapes where many sub-domains overlap on the same region, a boundary vertex of one sub-domain may show up in > 1 other sub-domains, resulting in equality constraints for each. This unnecessarily reduces the search space and tarnishes the solution near the overlap boundary (Figure 12). Much like in [PTSZ11], the transitivity of the equality above makes it so that we only *need* one coupling constraint for each boundary vertex. We cannot be satisfied with finding *any* maximal spanning tree of constraints since that may still concentrate constraints near a single vertex.

We experimented with various heuristics for picking which constraint to keep for each fixed vertex. Removing all but the first constraint creates a slight bias to the arbitrary ordering of the domains. Selecting a random constraint works reasonably well, but still results in many vertices involved in multiple constraints. Averaging or softening constraints also helps, but increases complexity.

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Figure 12: Boundary coupling across all sub-domain pairs leads to messy locking near the overlap boundary (6001 constraints). Instead, our heuristic keeps exactly one constraint per overlap boundary vertex (3893 constraints).



Figure 13: Pairwise constraints on multiple overlapping subdomains lead to redundancy, which we remove via constraint thinning. The highlighted vertex of the blue mesh in this didactic example is interpolated by elements in the green, red and purple meshes, when only one of these would be necessary.

Ideally, we would like to maximize the total number of vertices involved in the constraints (to diffuse the constraints) while still ensuring exactly one constraint per overlap-boundary vertex. Viewing the constraint matrix  $\mathbf{A}$  as graph, this selection is a form of vertex cover problem.

We approximate the maximum cover by *scoring* vertices based on how many constraints they are involved in; similarly, we score each constraint by averaging the scores of the vertices involved. For each vertex involved in more than one constraint, we keep the least saturated (lowest scored) constraint only and remove the rest, as shown in Figure 13. *Thinning* the constraints in this way significantly helps avoid issues near boundaries when multiple shapes overlap (Figure 12). The decrease in the number of constraints also reduces the linear system size, albeit with marginal affect on performance.

## 6. Higher Order Partial Differential Equations

Methods in geometry processing often go beyond second-order PDEs to model problems requiring smoother continuity at constraints [BK04a, SLCO\*04, JTSZ10, SGWJ17] or higher order control [FSH11, JC08] (Figure 14). Returning briefly to the smooth setting, we focus on the squared Laplacian energy to extend our consideration of deconstructed domains to higher order PDEs:

$$\min_{u} \int_{\Omega} (\Delta u)^2 \, dA, \tag{23}$$

resulting in the fourth-order bi-Harmonic equation:

$$\Delta^2 u(\mathbf{x}) = 0 \qquad \forall \mathbf{x} \in \Omega.$$
 (24)

The second derivatives of this energy are not immediately discretizable using linear FEM, so we introduce an auxiliary function z and solve the equivalent constrained minimization problem:

$$\min_{u,z} \int_{\Omega} z^2 \, dA,\tag{25}$$

subject to 
$$\Delta u(\mathbf{x}) = z(\mathbf{x}) \quad \forall \mathbf{x} \in \Omega.$$
 (26)

Applying the Lagrange multiplier method and Green's identity, this transforms into a saddle problem involving only first derivatives:

saddle 
$$\int_{\Omega} \left( z^2 + \nabla \mu \cdot \nabla u + \mu z \right) dA$$
 + boundary terms (27)

where  $\mu$  is the Lagrange multiplier function and we defer discussion of boundary terms to previous works (e.g. [SGWJ17]).

We now have a problem involving only first derivatives which we can discretize using multiple sets of linear finite elements (i.e. mixed FEM). After factoring out  $\mu$ , the resulting system has the symmetric matrix form of a KKT system:

$$\begin{pmatrix} \mathbf{0} & \mathbf{L}^{\mathsf{T}} \\ \mathbf{L} & -\mathbf{M} \end{pmatrix} \begin{pmatrix} \mathbf{u} \\ \mathbf{z} \end{pmatrix} = \begin{pmatrix} \mathbf{0} \\ \mathbf{0} \end{pmatrix}.$$
 (28)

## 6.1. Unsuccessful low-order boundary-only coupling

While the second-order Poisson equation requires one set of boundary conditions (e.g. fixed values or fixed normal derivatives), the fourth-order bi-Laplace equation in Equation (24) requires two sets of boundary conditions to identify a unique solution. For example, we can fix both the value *and* the normal derivative along the boundary  $\partial \Omega$  (i.e. fix low-order quantities). If we explicitly fix *only* the value along the boundary when minimizing the squared Laplacian energy, then *natural boundary conditions* will emerge to ensure uniqueness (cf. [SGWJ17]). This also occurs in the mixed FEM discretization. Fixing only the value along the overlapping region for two sub-domains  $\Omega_1 \cap \Omega_2$  couples the function values together, but produces a noticeable 'kink' (Figure 15). We are witnessing the natural boundary conditions on one sub-domain's function (in this case,  $\Delta u = 0$ ) disagreeing with the derivatives of *other* subdomain's function: i.e. in general,  $\Delta u_1 = 0 \neq \Delta u_2$  on  $\partial \Omega_1 \cap \Omega_2$ .

To take advantage of the same uniqueness properties used in Section 5, we must ensure that each function is sufficiently constrained with boundary conditions. One idea would be to trivially extend our boundary-only coupling by fixing the value *and* normal derivative along the overlapping boundary:

$$u_1(\mathbf{x}) = u_2(\mathbf{x}) \quad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2), \tag{29}$$

$$\nabla u_1(\mathbf{x}) \cdot \mathbf{n}(\mathbf{x}) = \nabla u_2(\mathbf{x}) \cdot \mathbf{n}(\mathbf{x}) \quad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2), \tag{30}$$

where  $\mathbf{n}(\mathbf{x})$  is the normal vector pointing outward from the overlapping region  $\Omega_1 \cap \Omega_2$ . In the smooth setting, we can quickly confirm that this is equivalent to the original energy minimization problem in Equation (23) following the same reasoning in Section 5.

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Figure 14: Updating the shape with another stack of wings does not require updating a unified tet mesh. Instead, new components are tet-meshed independently and added to the system algebraically.



**Figure 15:** Enforcing equality at all vertices of two overlap domains results in locking (red and yellow). For higher order PDEs, coupling by value alone at overlap boundaries avoids extreme locking, but resolution refinement reveals non-smoothnesses (pinks). Attempting to enforce derivative continuity by coupling values and first derivatives results in local locking and non-smoothness persists (blues). We couple primary values and auxiliary values constrained to the Laplacian (u'' in 1D) at sub-domain boundaries: the solution is smooth (purple and green).

These *low-order* coupling constraints are simple to discretize using linear FEM, but unfortunately do not lead to a convergent system. Fixing directional derivatives across the two functions leads to harmonic locking locally (the one-ring of vertices at the overlapping region boundary). This region shrinks with mesh refinement, but the problem persists: effectively, the solution locks so that natural boundary conditions emerge, albeit one-ring into the overlapping domain (Figure 15).

# 6.2. Higher order boundary-only coupling

Fortunately, the bi-Laplace equation in Equation (24) is also uniquely determined by other combinations of boundary conditions. Such combinations of low- and high-order conditions sometimes appear directly during problem modelling (e.g. [JC08]). The introduction of the auxiliary variable  $z = \nabla u$  in Equation (25) makes the choice of fixing the value and the Laplacian of *u* along the boundary particularly easy to describe:

$$u_1(\mathbf{x}) = u_2(\mathbf{x}) \qquad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2),$$
 (31)

$$z_1(\mathbf{x}) = z_2(\mathbf{x}) \qquad \forall \mathbf{x} \in \partial(\Omega_1 \cap \Omega_2). \tag{32}$$

During discretization using mixed FEM, we add these constraints to the Lagrangian's KKT system in Equation (28) directly, resulting in a larger KKT system:

$$\begin{pmatrix} \mathbf{0} & \mathbf{L}^{\mathsf{T}} & \mathbf{A}^{\mathsf{T}} & \mathbf{0} \\ \mathbf{L} & -\mathbf{M} & \mathbf{0} & \mathbf{A}^{\mathsf{T}} \\ \mathbf{A} & \mathbf{0} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{A} & \mathbf{0} & \mathbf{0} \end{pmatrix} \begin{pmatrix} \mathbf{u} \\ \mathbf{z} \\ \lambda_u \\ \lambda_z \end{pmatrix} = \begin{pmatrix} \mathbf{0} \\ \mathbf{0} \\ \mathbf{0} \\ \mathbf{0} \end{pmatrix}, \quad (33)$$

**Log-log convergence plot for bi-Laplace on 2D annulus** L\_error



**Figure 16:** The behaviour of our different sets of constraints for the bi-Laplace equation  $\Delta^2 u = 0$  mirrors that of the Poisson equation solver shown previously.

where  $\lambda_u, \lambda_z \in \mathbb{R}^{n_1+n_2}$  are vectors of Lagrange multipliers enforcing boundary coupling constraints on **u** and **z**, respectively, and **A** is the linear constraint matrix. While the constraints on the values in **u** are straightforward, the constraints on the auxiliary values **z** may be interpreted as acting orthogonally to the original mixed FEM constraint so that  $\mathbf{Mz} = \mathbf{L}u$ .

This discretization avoids the 'kink' of the low-order boundary coupling in Section 6.1 (Figure 15). We see convergence with respect to mesh refinement (Figure 16).

A remaining issue with our discretization is that mixed FEM results in a saddle problem, rather than a standard convex, linearly constrained quadratic energy minimization. However, this is overcome by rearranging terms algebraically (see Appendix B).

Table 1: Performance timings.

Shape	#Tets	Κ	Build A	Problem	Solve
Android	113 118	33	0.56 s	BBW	9.06 s
Bug	159 533	16	0.47 s	MSBK	21.73 s
Jet	226 548	14	0.48 s	Eigen	1.91 s
Bi-plane	321 237	23	1.05 s	BBW	37.61 s
Microscope	348 099	24	1.09 s	Heat	4.84 s
Pistol	412 798	18	1.08 s	Wave	4.70 s
Alien	682 399	32	2.57 s	Geodesic	13.45 s

Notes: #Tets is the total number of tetrahedra across the *K*overlapping components. We list the runtime for constructing the constraints matrix (*Build* **A**) and then conducting the resulting constrained (example-dependent) optimization (*Solve*).

# 7. Experiments and Results

We have implemented our method using MATLAB using finite element operators from GPTOOLBOX [J\*16] and point location routines from LIBIGL [JP\*18]. We use TETGEN [Si03] to mesh the sub-domains in all examples except the sphere and 3D annulus test cases, for which we use QUARTET [DCB13]. We use TRIANGLE [She96] for 2D meshing. On our MacBook Pro with a 3.5GHz Intel Core i7 with 16 GB of memory, the performance bottleneck is always the linear solve (MATLAB's ldl), eigen decomposition (MATLAB's eigs) or quadratic programming optimization (MOSEK's quadprog). For completeness, we list runtime performance in Table 1.

While our main focus is to improve robustness, we observe systematically predictable trends in the runtime performance. For example, consider solving a Poisson equation on a single mesh of a solid domain with  $O(n^3)$  vertices. For a typical FEM-quality mesh, the performance will be determined by performing a linear system solve on a sparse matrix with  $O(n^3)$  non-zeros. In the absence of other constraints, this matrix will be positive definite, affording Cholesky decomposition. For our deconstruction of the same domain into K overlapping components and  $O(n^3)$  total vertices across all meshes, we build the boundary-only constraints matrix A which (under mild assumptions) will contain  $O(Kn^2)$ non-zeros. In contrast, fixing all vertices in the overlapping region would require  $O(Kn^3)$  non-zeros. Using, e.g. the Lagrange multiplier method to enforce our constraints results in an indefinite sparse system matrix with  $O(n^3 + Kn^2)$  non-zeros (solved, e.g. with LDLT-decomposition). In practice, K is often quite small and the difference in performance between solving on a single mesh and a deconstructed domain boils down to the performance of sparse Cholesky versus sparse LDLT-decomposition - with the important caveat that solving on a *single mesh* is often impossible without user-intervention. In Figure 17, we found an example where meshunion followed by tetrahedralization does create a useable mesh: Cholesky for the single mesh is roughly 3.7× faster than LDLT on our constrained system. As future work, it would be interesting to further exploit our deconstructed domains for performance acceleration and parallelization (see, e.g. [LSLR01]).

An alternative to our boundary-only *hard* constraints would be to enforce *weak* constraints at all vertices in the overlapping regions. In Figure 18, we show that, yes, weak constraints can work, but one



**Figure 17:** Traditional mesh-union then tetrahedralization succeeds on this example, enabling an ad hoc performance comparison. System matrices for solving the Poisson equation require similar memory; however, our MATLAB implementation uses LDLT on our resulting Lagrangian: about  $3.7 \times$  slower than Cholesky, here.



**Figure 18:** Weak constraints are sensitive to the penalty weight. There exists a good value (here,  $\omega = 0.1$ ), but finding this is nontrivial and problem dependent. An incorrect choice can be disastrous. Our boundary coupling is parameterless and achieves low error.

must choose the penalty weight carefully. In this experiment, if the penalty is too weak, the solutions on different sub-domains become decoupled; too strong the solution locks up just as much as the strong constraints. This is not a situation where a different constraint handler will help. For example, the augmented Lagrangian or alternating direction method of multipliers methods are numerical techniques for effectively driving the penalty weight to infinity, but in this limit the solution is simply the locked up solution. Meanwhile, the 'correct' penalty weight will depend on the mesh resolution, constraint constellation and solution. This may vary spatially: a good weight here may cause locking over there.

We designed 2D and 3D convergence test scenarios (Figures 8 and 10). We compare  $L_{\infty}$  error to an analytic solution. In Figure 11, we use the same setup to test partial area estimation. For 10-point quadrature, we use the symmetric rules of Zhang *et al.* [ZCL09].

For irregular tetrahedral meshes, elements overlapping the boundary of another domain typically contain multiple boundary vertices of that domain and thus participate in multiple constraints (even in the simple overlapping 3D annuli in Figure 10, *involved* tetrahedra contain on average 3.18 boundary vertices). For more complex shapes, the interior boundaries inherit the irregularity of the overlapping parts. Our method does not smooth or alter these potentially irregular boundaries (Figure 1).



Figure 19: Our general method can be applied to a variety of problems involving discrete differential geometry including eigen analysis.



**Figure 20:** Our method is accurate enough to repetitive solve for dynamics, as seen in this simulation of a shockwave propagating from one solid component into the rest.

We demonstrate the versatility of our constraints by expanding beyond the Laplace ( $\Delta u = 0$ ) and Poisson equations ( $\Delta u = f$ ) to other equations found in solid geometry processing. In Figure 2, we demonstrate our boundary-only constraints for solving an implicit time step of the heat equation ( $u - \delta t \Delta u = u_0$ ). In Figure 1, we solve the same heat equation for u and then the Poisson equation  $\Delta \varphi = -\Delta u/|\nabla u|$  to approximate interior distances  $\varphi$  using the method of Crane *et al.* [CWW13]. In Figure 20, we visualize shock wave through a pistol composed of many overlapping components  $(u - \delta t^2 \Delta u = u_0 + \delta t \dot{u}_0)$ .

In Figure 19, we use our boundary-only constraints to conduct a Laplacian modal analysis on a deconstructed domain. We enforce constraints during eigen decomposition via the null space method [Gol73], but replace the QR decomposition with the sparser LUQ decomposition. In Figure 21, we quantitatively validate our method using the Laplacian spectrum. The smallest 100 eigenvalues using our method match the theoretical groundtruth for a sphere domain (and those computed using standard linear FEM on a single mesh). To extend this comparison to a more complex example where theoretical values are not known, we found a shape where mesh-union followed by tetrahedralization succeeds. Compared to second-order finite differences over a high-resolution voxelization, our spectrum better matches the spectrum found using a single unified mesh. Higher order elements — known to improve spectral convergence [RBG\*09] — could be used in either method, but do not affect our main contribution of setting up constraints.

By rearranging our higher order coupling for bi-Laplacian problems in Section 6.2 into a convex energy minimization (see Appendix B), we can immediately implement advanced methods involving  $L_1$  sparsity inducing norms for shape descriptors, such as the multiscale pre-biharmonic kernels [Rus11] in Figure 22 and inequality constraints such as the bounded biharmonic weights [JBPS11], used for real-time skinning deformations in Figure 23. In Figure 14, we demonstrate the robustness of our method to large-scale geometry changes. Wings are added to the plane simply by overlapping



**Figure 21:** The spectral behaviour of our Laplacian operator constructed using solely the information from the primitive's meshes (green lines) approaches the analytical spectra (left) in the same way as that obtained from traditional FEM on a unified mesh of the domain (left and right, blue line).

16 overlapping solids Volumetric Multiscale Biharmonic Kernels



**Figure 22:** Our high-order boundary-coupling constraints complement advanced biharmonic energy-minimization methods and additional constraints such as  $L_1$  sparsity.

new solid components: we only need to tet-mesh the new components and add their linear constraints to the system. In the classic geometry processing pipeline, we would need to invoke mesh union and fragile global tet-meshing algorithms. Our method avoids this.

## 8. Limitations and Future Work

We make a heavy assumption that the input domain is or can be deconstructed into simple tetrahedralizable sub-domains. While many models are originally created using CSG operations, often only the (typically poor triangle-quality) mesh-boolean result is available when it comes time to solve a volumetric PDE. Therefore, we advocate to *retain* these simpler domains and the construction tree rather than preemptively resolving the mesh-boolean. Recent developments on obtaining the CSG tree from a given shape via reverse



Figure 23: Minimizing the squared Laplacian subject to bound constraints produces automatic deformation bases: our method enables this over deconstructed domains.



**Figure 24:** Our method can be generalized to deal with other 2D set operations, such as intersections and substractions.

engineering [DIP\*18] are promising towards reducing the severity of this limitation.

Nonetheless, our tetrahedralizers, TETGEN [Si03] and QUARTET [DCB13], still occasionally fail even on simpler sub-domains. To mitigate this, we can pre-process problematic sub-domains on a case-by-case basis using MESHFIX [Att10] and generalized winding numbers [BDS\*18]. Alternatively, one could use the recently published TETWILD [HZG\*18] directly on the primitive shapes.

In this paper, we consider volumetric unions of polyhedral subdomains. Other domains such as those modelled using metaball implicits [WMW86] or reconstructed from unstructured point clouds (e.g. [KBH06]) are not immediately suitable for our method. It is exciting to consider automatic methods for converting such domains into unions of simpler primitives, perhaps with inspiration from advances in approximate convex decomposition [AGCO13].

While all examples presented in this paper deal exclusively with unions of different shapes, one can conceive of certain variations that would make our method valid for all CSG operations, such as intersections or differences. We have promising initial results for intersecting two-domains in 2D (Figure 24) and are working an extension to full 3D CSG trees.

#### Acknowledgements

The first four authors were supported by the 2017 Fields Undergraduate Summer Research Program. Silvia Sellán is also funded by the Maria Cristina Masaveu Peterson Scholarship for Academic Excellence. We thank: Eitan Grinspun, David I.W. Levin, Oded Stein and Héctor Jardón Sánchez for insightful and helpful conversations; Sarah Kushner for presenting our poster at SGP 2018; Darren Moore, Gavin Barill, Michael Tao and Sarah Kushner for proofreading; and the TOMOGRAPH 2017 workshop for hosting an early presentation. This research is funded in part by NSERC Discovery (RGPIN2017–05235, RGPAS–2017–507938), Connaught Funds (NR2016–17), the Canada Research Chairs Program, the Fields Centre for Quantitative Analysis and Modelling and gifts by Adobe Systems, Autodesk and MESH Inc.

#### **Appendix A: Deconstructed Domains Solver**

This appendix provides a step-by-step construction of the discrete solver for deconstructed domains. Without loss of generality, let us assume 3D domains (d = 3). The input to our method is a set of K overlapping, embedded, manifold tetrahedral meshes with vertices  $\{\mathbf{V}_1, \ldots, \mathbf{V}_K\}$  so that  $\mathbf{V}_i \in \mathbb{R}^{n_i \times 3}$  contains the positions of the *i*th sub-domain's  $n_i$  vertices in its rows and list of tetrahedral indices  $\{\mathcal{T}_1, \ldots, \mathcal{T}_K\}$ , where the row-indices into  $\mathbf{V}_i$  of the *i*th sub-domain's  $t_i$  tetrahedra appear as rows  $\mathcal{T}_i \in [1, \ldots, n_i]^{t_i \times 4}$ .

We first build the constraint matrix  $\mathbf{A} \in \mathbb{R}^{(b_1 + \dots + b_K) \times (n_1 + \dots + n_K)}$ , where  $b_i$  are the number of boundary vertices of the *i*th mesh lying inside a tetrahedron of *any* other mesh.

Next, we build the sparse discrete gradient matrix  $\mathbf{G}_i \in \mathbb{R}^{3t_i \times n_i}$  for each domain and compute *adjusted* volumes for each tetrahedron  $\mathbf{a}_i \in \mathbb{R}^{t_i}$  (accounting for the  $1/\sum_{j=1}^{K} \chi_j$  term in Equation (21), see Section 5.2).

From these, we can construct a quadratic coefficients matrix (i.e. discrete Laplacian)  $\mathbf{L}_i \in \mathbb{R}^{n_i \times n_i}$  for each domain:

$$\mathbf{L}_i = \mathbf{G}_i^{\mathsf{T}} \operatorname{diag}\left(\mathbf{a}_i\right) \mathbf{G}_i, \qquad (A.1)$$

where diag(**x**) for a vector  $\mathbf{x} \in \mathbb{R}^n$  creates  $n \times n$  matrix with **x** along the diagonal. We concatenate the contributions from each subdomain into a monolithic Laplacian  $\mathbf{L}_i \in \mathbb{R}^{(n_1+\dots+n_K)\times(n_1+\dots+n_K)}$ ,

$$\mathbf{L} = \text{blkdiag}\left(\mathbf{L}_{1}, \dots, \mathbf{L}_{K}\right), \qquad (A.2)$$

where  $blkdiag(\mathbf{A}, \mathbf{B}, ...)$  creates a block diagonal matrix from matrices  $\mathbf{A}, \mathbf{B}, ...$ 

Using the adjusted tetrahedral volumes in  $\mathbf{a}_i$ , we build a 'barycentric' lumped diagonal mass matrix for each mesh  $\mathbf{M}_i \in \mathbb{R}^{n_i \times n_i}$  and stack these as well to create the mass matrix of the entire system  $\mathbf{M} \in \mathbb{R}^{(n_1+\dots+n_K)\times(n_1+\dots+n_K)}$ ,

$$(\mathbf{M}_i)_{jj} = \sum_{k \in N(j)} \frac{1}{4} (\mathbf{a}_i)_k \tag{A.3}$$

$$\mathbf{M} = \text{blkdiag}\left(\mathbf{M}_{1}, \dots, \mathbf{M}_{K}\right), \qquad (A.4)$$

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where N(j) are the tetrahedra incident on vertex j. Further accuracy could possibly be achieved by using a hybrid 'Voronoi' mass matrix [MDSB03, JTSZ10].

Finally, we define  $\mathbf{u} \in \mathbb{R}^{(n_1+\dots+n_K)}$  as the vertically stacked vectors of unknown per-vertex values across the *K* sub-domain meshes.

We may now pose the discretization of the energy minimization problem in Equations (21) and (22) using a standard matrix form:

$$\min \mathbf{u}^{\mathsf{T}} \mathbf{L} \mathbf{u} - \mathbf{u}^{\mathsf{T}} \mathbf{M} \mathbf{1}$$
 (A.5)

subject to 
$$u_{ii} = g(\mathbf{v}_{ii}) \quad \forall \mathbf{v}_{ii} \in \partial \Omega \cap \Omega_i$$
 (A.6)

and 
$$\mathbf{A}\mathbf{u} = \mathbf{0}$$
, (A.7)

where **1** and **0** are vectors ones and zeros, respectively. Vertices receiving boundary conditions or constraints are identified combinatorially and located inside other meshes efficiently using a spatial acceleration data structure (e.g. we use LIBIGL'S AABB tree [JP\*18]), and then *thinned* by removing rows according to our approximate max-cover criteria (see Section 5.2.1). We use the MATLAB or MOSEK quadratic programming solvers to find an optimal **u**.

#### **Appendix B: Rearrangement into Quadratic Minimization**

A remaining issue with our discretization is that mixed FEM results in a saddle problem, rather than a standard convex, linearly constrained quadratic energy minimization. This means in practice we cannot send the system in Equation (33) to a standard quadratic programming solvers because the top-left sub-block

$$\begin{pmatrix} \mathbf{0} & \mathbf{L}^{\mathsf{T}} \\ \mathbf{L} & -\mathbf{M} \end{pmatrix} \tag{B.1}$$

is not positive semi-definite. However, we can resolve this by factoring out  $\mathbf{z} = \mathbf{M}^{-1}(\mathbf{L}u + \mathbf{A}^{\mathsf{T}}\lambda_z)$  resulting in the smaller KKT system:

$$\begin{pmatrix} \mathbf{L}^{\mathsf{T}}\mathbf{M}^{-1}\mathbf{L} & \mathbf{L}\mathbf{M}^{-1}\mathbf{A}^{\mathsf{T}} & \mathbf{A}^{\mathsf{T}} \\ \mathbf{A}\mathbf{M}^{-1}\mathbf{L}^{\mathsf{T}} & \mathbf{A}\mathbf{M}^{-1}\mathbf{A}^{\mathsf{T}} & \mathbf{0} \\ \mathbf{A} & \mathbf{0} & \mathbf{0} \end{pmatrix} \begin{pmatrix} \mathbf{u} \\ \lambda_z \\ \lambda_u \end{pmatrix} = \begin{pmatrix} \mathbf{0} \\ \mathbf{0} \\ \mathbf{0} \end{pmatrix}, \quad (B.2)$$

where the top-left  $2 \times 2$  sub-block is now positive semi-definite. This system arrives as the Euler–Lagrange equation for the constrained convex quadratic minimization problem:

$$\min_{\mathbf{u},\lambda_z} \|\mathbf{M}^{-1/2}(\mathbf{L}\mathbf{u} + \mathbf{A}^{\mathsf{T}}\lambda_z)\|^2$$
(B.3)

subject to 
$$\mathbf{A}\mathbf{u} = \mathbf{0}$$
. (B.4)

To avoid inverting the mass matrix and improve the conditioning of the objective term, we introduce another auxiliary variable  $\mathbf{y} \in \mathbb{R}^{n_1+n_2}$ , arriving at the final constrained problem in standard form:

$$\min_{\mathbf{y}, \lambda \in \mathbf{y}} \|\mathbf{y}\|^2 \tag{B.5}$$

subject to 
$$Au = 0$$
, (B.6)

and 
$$\mathbf{L}\mathbf{u} + \mathbf{A}^{\mathsf{T}}\lambda_z = \mathbf{M}^{\mathsf{T}/2}\mathbf{y}.$$
 (B.7)

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