Composite-grid multigrid for diffusion on the sphere

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Summary
Recently, there has been much interest in the solution of differential equations on surfaces and manifolds, driven by many applications whose dynamics take place on such domains. Although increasingly powerful algorithms have been developed in this field, many straightforward questions remain, particularly in the area of coupling advanced discretizations with efficient linear solvers. In this paper, we develop a structured refinement algorithm for octahedral triangulations of the surface of the sphere. We explain the composite-grid finite-element discretization of the Laplace–Beltrami operator on such triangulations and extend the fast adaptive composite-grid scheme to provide an efficient solution of the resulting linear system. Supporting numerical examples are presented, including the recovery of second-order accuracy in the case of a nonsmooth solution.

KEYWORDS
fast adaptive composite-grid (FAC) algorithm, finite-element discretizations on surfaces, Laplace–Beltrami operator, multigrid

1 INTRODUCTION

Discretization of scalar elliptic partial differential equations (PDEs) over regions in the plane or volumes in \( \mathbb{R}^3 \) is, in general, a well-understood subject, with viable finite-difference, finite-element, and finite-volume approaches seen in standard textbooks and references. Mesh-refinement techniques are also well understood in this setting, with numerous strategies and algorithms appearing in the literature. For standard scalar elliptic problems, on both uniform and refined meshes, the solution of the resulting linear systems is typically accomplished efficiently using multigrid methods, including robust geometric methods and algebraic multigrid (AMG).

A wide body of recent research has been devoted to extending classical discretization approaches to PDEs posed on surfaces and manifolds. Natural techniques that parameterize a surface based on regular coordinates in the plane are one option, but these are limited to simple surfaces and introduce problematic singularities, both in the underlying surface meshes and in the discretized differential operators. Embedding methods are another approach, exemplified by the closest point method (CPM) of Ruuth and coauthors, which inherits attractive numerical properties by appropriately embedding the surface in a surrounding volume and extending the PDE to that volume. Alternatively, one can aim to discretize the surface directly, by defining an approximating triangulated surface and then applying standard finite-element (or finite-volume) approaches to discretize the PDE on the triangulated surface; see the review article by Dziuk et al. and the many references therein.

Within each class of discretization methodology, specific advances have been made, particularly in the development of fast solution algorithms. For example, for the finite-volume discretization of the Laplace–Beltrami operator on the surface of the sphere using latitude–longitude meshes, an optimal multigrid method is known using alternating line relaxation and a transformed restriction operator. Multigrid solvers have also been developed for CPM discretizations. For uniform triangulations...
of surfaces, Bey\textsuperscript{28} and Landsberg et al.\textsuperscript{29} demonstrate the effectiveness of simple multigrid methods, using natural finite-element interpolation operators and simple point smoothing. Motivated by the problem of radiation dose planning (as explained in Section 2), this paper aims to extend these results to locally refined triangulations of the sphere for problems with highly localized sources, using the fast adaptive composite-grid (FAC) methodology.

The FAC methodology was first proposed in the 1980s\textsuperscript{15,16} to extend optimal multigrid performance to meshes generated by structured adaptive mesh refinement algorithms. Many such algorithms generate meshes that are semistructured, which can be characterized by having distinct “levels” of refinement with regular structure within each level. Although such systems can be solved using AMG,\textsuperscript{19} this is naturally inefficient as AMG makes no use of the existing structure in the mesh. FAC, in contrast to AMG, explicitly requires that the composite grid consist of distinct mesh levels, each of which can be considered independently of the others. FAC multigrid cycles consist of independent processing on each level, coupled by intergrid-transfer operators as in standard multigrid, and can be done both multiplicatively\textsuperscript{16} or additively (asynchronously).\textsuperscript{17,18} We note that many other generalizations of the multigrid methodology are possible, including popular subspace and domain decomposition methods\textsuperscript{30,31}; however, the hierarchical structure of FAC maps naturally onto the hierarchical structure of local mesh refinement that will be considered here.

The main goal of this paper is to bring together the aspects of mesh refinement and fast multigrid solution for PDEs dominated by the Laplace–Beltrami operator with localized sources on the surface of the sphere. In particular, we develop and apply the FAC methodology for the Laplace–Beltrami operator on the sphere, discretized on locally refined meshes. Our motivating application, radiation dose planning, is explained in Section 2, including the Fokker–Planck limit in which the Laplace–Beltrami operator is obtained. In Section 3, we propose a structured mesh refinement algorithm, which is later demonstrated to restore second-order convergence when the source function is singular. Finite-element discretization is used on the resulting nonuniform triangulation of the sphere, discussed in Section 4. In Section 5, the FAC methodology is explained and adapted into a solution algorithm for this discretization. Supporting numerical results are presented in Section 6, including for an example with a nonsmooth solution where mesh refinement is expected to be most beneficial. Conclusions and comments on future work are given in Section 7.

## 2 MOTIVATING APPLICATION: RADIATION DOSE PLANNING

As a motivating application, we consider the radiation dose planning problem, which can be posed as an optimal control problem,

$$\min_{(f, g)|\mathcal{L}(f) = g} J(f),$$

where $f$ is the phase-space density of charged particles in tissue. In this model, $J(f)$ is a cost function representing irradiation of cancerous tissue, whose minimum value of 0 is achieved when $f$ is sufficiently large within all identified cancerous tissue in the domain, $\Omega \subset \mathbb{R}^3$, and zero (or sufficiently small) in all healthy tissue. The constraint $\mathcal{L}(f) = g$ represents the solution of the linear Boltzmann transport equation

$$\frac{1}{c} \frac{\partial f}{\partial t} + \omega \cdot \nabla f + \sigma_t(\vec{x}, E)f - \mathcal{K}_s f = 0,$$

subject to initial conditions $f(\vec{x}, \omega, E, t_0) = f_0(\vec{x}, \omega, E)$ and boundary conditions $f(\vec{x}, \omega, E, t) = g(\vec{x}, \omega, E, t)$ for $\vec{x} \in \partial \Omega$. To define the notation, $f(\vec{x}, \omega, E, t)$ is the phase-space density of particles at location $\vec{x} \in \Omega$, moving in direction $\omega \in S$, with energy $E$ at time $t$. Here, $S$ is the unit sphere in $\mathbb{R}^3$. Key parameters in the problem are the particle speed $c$, the probability of interaction (absorption or scattering) per unit distance traveled $\sigma_t$, and the scattering kernel $\mathcal{K}_s$. For charged particles, the mean free path, $\lambda = 1/\sigma_t$, is relatively small, but most interactions result in relatively small changes in angle, direction, and energy. The control variable, $g(\vec{x}, \omega, E, t)$, is chosen from the set of admissible configuration of beams of charged particles, to properly irradiate the cancerous tissue. In real-world applications there are, of course, many practical constraints on the possible values of $g$ imposed by available equipment and medical safety. Efficient solution algorithms for such optimization problems often rely on efficient solution algorithms for the constraining PDEs, and it is this “forward problem” that we concentrate on in this paper.

To first simplify this model, we consider the time-independent case of monoenergetic transport (constant $c$ and $E$) and assume that all interactions result in scattering, so that

$$\mathcal{K}_s f(\vec{x}, \omega) = (\frac{1}{\lambda})^{-1} \int_S \frac{p(\omega \cdot \omega')}{2\pi} f(\vec{x}, \omega') d\omega',$$
where \( p(\omega \cdot \omega')/2\pi \) is the probability density for scattering from direction \( \omega' \in S \) to direction \( \omega \), depending only on the angle between unit vectors \( \omega \) and \( \omega' \). In this setting, the governing Boltzmann equation simplifies to

\[
\omega \cdot \nabla f(\vec{x}, \omega) = \mathcal{K}_s f(\vec{x}, \omega) - (\overline{\mu})^{-1} f(\vec{x}, \omega) = (\overline{\lambda})^{-1} \left( \int_S \frac{p(\omega \cdot \omega')}{2\pi} f(\vec{x}, \omega') d\omega' - f(\vec{x}, \omega) \right)
\]

(1)

and the domain of \( f \) is now \( \Omega \times S \). The balance in Equation 1 between the advective term on the left and the scattering term on the right depends on the properties of the probability density \( p(\mu) \). Of particular interest is the expected value of \( \mu = \omega \cdot \omega' \), defined as \( \overline{\mu} = \int_S \mu p(\mu) d\mu \), which gives the expected value of the cosine of the angle between an incident direction \( \omega' \) and a scattered direction \( \omega \). In many models of the scattering of charged particles, the scattering occurs very frequently, giving a small mean free path, \( \overline{\lambda} \), but each interaction results in only a small deflection, giving \( \overline{\mu} \approx 1 \). The Fokker–Planck limit considers the case where \( \overline{\lambda} \to 0 \) and \( \overline{\mu} \to 1 \) in a way such that the transport mean-free path, \( \overline{\lambda}_{tr} = \frac{\overline{\lambda}}{1 - \overline{\mu}} = \frac{2}{T} > 0 \), is fixed, defining the linear scattering power, \( T \). In this case, Börgers et al.32 show that the right-hand side of Equation 1 converges weakly to \( \frac{1}{2\overline{\lambda}_{tr}} \Delta_s f \), the Laplace–Beltrami operator on the sphere. The resulting equation,

\[
\omega \cdot \nabla f(\vec{x}, \omega) = \frac{1}{2\overline{\lambda}_{tr}} \Delta_s f,
\]

(2)

is known as the Fokker–Planck equation.

In recent years, several contributions have been made to the literature on both the discretization and efficient solution of Equation 1, primarily drawing on insight from the limiting case in Equation 2. Discretization in space is typically handled using upwind finite differences or DG-type discretizations, yielding a natural downstream ordering of the resulting linear systems. In angle, several discretizations of the unit sphere have been considered, including the \( S_n \) (discrete ordinates) discretization,33 popular in the literature on computational transport, but finite-difference and Galerkin finite-element methods have also been used. Morel et al.34 proposed an angular multigrid method for the one-dimensional analog of Equation 1, with semicoarsening in the angular variable and a “downstream” Gauss–Seidel relaxation used to resolve the spatial variation. This was extended to a two-dimensional problem in the work by Börgers et al.,35 for a finite-difference discretization in “flatland,” where the domain of \( f \) is \([0, 1]^2 \times [0, 2\pi] \); a domain-decomposition preconditioner for this problem was also considered by Börgers.36 Similar work has also been done for three-dimensional scattering, in the case of variable \( E \),37–40 considering the \( S_n \) discretization scheme. These papers consider a much broader class of transport problems than in others,35,36,41,42 including both isotropic (neutron) and anisotropic scattering, as well as the multienergetic case. In studies by Gao et al.,41,42 a uniform-grid finite-element discretization of \( S \) is used in the case of anisotropic scattering, coupled with a finite-difference or DG discretization of the angular terms. A general conclusion from these works is that efficient and accurate simulation of the Boltzmann equation for charged particles is possible, provided that a fast solver is available for an accurate discretization of the angular part of the operator.

Considering the motivating application of radiation dose planning, a natural inefficiency arises in the use of uniform discretizations of the sphere, because the typical boundary conditions impose beams of charged particles that are highly localized in direction. Thus, in this paper, we consider the question of mesh refinement on the sphere, restricted to the Laplace–Beltrami operator (or its reaction–diffusion analog). Although extensions to the full Fokker–Planck and linear Boltzmann equation are necessary for the resulting algorithm to be applied in the context of the radiation dose planning problem, we leave these for future work. In what follows, we propose a structured mesh refinement strategy that addresses (reaction–)diffusion on the sphere with localized sources, a natural finite-element discretization that complements this approach, and we extend the FAC multigrid approach to these meshes and this discretization. The proposed discretization and refinement scheme are similar to those recently proposed for neutral-particle transport by Lau et al.43

3 STRUCTURED REFINEMENT OF THE SPHERE

Let \( S \) be the unit sphere in \( \mathbb{R}^3 \). We henceforth consider the solution of PDEs on the sphere of the form

\[
-\Delta_s u(\omega) + au(\omega) = g(\omega) \quad \forall \omega \in S,
\]

(3)

where \( \Delta_s \) represents the Laplace–Beltrami operator on \( S \) and \( a \geq 0 \) is a constant. We consider specifically the case where \( g(\omega) \) is a function either with localized support or where it is very small except over a small area on \( S \). Our goal is to define a nonuniform triangulation of \( S \), such that the variation of \( g \) (and, consequently, \( u \)) is efficiently resolved by the triangulation. When discretizing Equation 3 using a finite-element approach (discussed in Section 4), this is essentially equivalent to saying that the triangulation yields a good discrete approximation for the solution, \( u(\omega) \).
In this paper, we consider the case where the local structure in \( g(\omega) \) is known in advance and, thus, structured refinement approaches are more natural than adaptive mesh refinement. We start with a “base” triangulation of the sphere, \( \mathcal{T}_0 \). Natural choices are to use an octahedron or icosahedron as the “zeroth” mesh, with all points of the polyhedron lying on \( S \), although refinements of such meshes or alternate base triangulations could also be used. Given an existing mesh, our general (local-then-global) refinement algorithm is considered to have three parts:

1. Mark triangles for local refinement.
2. Refine marked triangles.
3. Refine all triangles (including those created in Step 2).

Whenever a triangle is refined in this algorithm, we consider uniform refinement, bisecting each edge of the triangle to create new nodes, interconnecting these nodes to create four new subtriangles, and then projecting these nodes onto the surface of the sphere. If multiple levels of refinement are to be done simultaneously, either in Step 2 or 3, they are done by sequentially bisecting the edges the appropriate number of times.

Considering the case where \( g(\omega) \) is a point source (or similar) located at the north pole of the sphere, one instance of this strategy will be to consider the refinement of an octahedron with one vertex located at the north pole. In the algorithm above, we will then mark all triangles adjacent to the north pole for local refinement, in addition to a global refinement. Figure 1 shows the resulting triangulations when each refinement step is a single level, dividing each refined triangle into four subtriangles. Multiple simultaneous levels of refinement are also possible within this algorithm, as is reversing the order of the global and local refinement. Both of these allow for finer grained control of the relative density of points in the refined region and those away from the refinement. Figure 2 shows the resulting meshes from the “reverse” (global-then-local) refinement strategy, where a global refinement step is first applied; then, the resulting elements adjacent to the north pole are refined a second time. In comparison to the meshes in Figure 1, we see a more focused refinement pattern around the north pole, whereas the global base mesh extends further into the northern hemisphere. These are, of course, very naïve approaches to mesh refinement, suitable only to the special case considered here, where \( g(\omega) \) is localized around a known point in the domain. For more broadly applicable mesh adaptation strategies based on local error estimates of the surface finite-element discretization, see, for example, Demlow et al.\(^{44} \) and Camacho et al.\(^{45} \)

![Figure 1](image1.png)  
**FIGURE 1** Three meshes obtained by applying the local-then-global refinement strategy where marked triangles are those adjacent to the north pole

![Figure 2](image2.png)  
**FIGURE 2** Three meshes obtained by applying the global-then-local refinement strategy where marked triangles are those adjacent to the north pole after a global refinement is performed
The leftmost image in Figure 2 highlights an important property of the resulting triangulations, which is that they are naturally nonconforming, in the sense that the local refinement step introduces “hanging nodes.” Due to the projection of such nodes onto the surface of the sphere, these nodes “disconnect” the triangulation; although an additional face could be inserted to fill this area, we choose not to and, instead, simply enforce continuity in the finite-element approximation defined in the following. This avoids the introduction of elements with poor shape regularity that would otherwise negatively impact the discrete problem to be solved.

In what follows, we view such refined meshes as a hierarchy of patches centered at the north pole. If \( \ell \) steps of the refinement algorithm are performed, we can consider a quasi-uniform global mesh \( \mathcal{T}_{\ell,0} \), which is formed by taking the base mesh \( \mathcal{T}_0 \) (an octahedron for the refined meshes shown in Figures 1 and 2) and refining it \( \ell \) times. We then define a sequence of \( \ell \) “patches,” with \( \mathcal{T}_{\ell,1} \) denoting the first refinement patch around the north pole, starting from the elements created by the first local refinement step that have also been uniformly refined \( \ell \) times. Thus, if \( h_0 \) is a representative meshwidth for the global mesh \( \mathcal{T}_{\ell,0} \), then the representative meshwidth for \( \mathcal{T}_{\ell,k} \) is \( h_k \approx h_{k-1}/2 \), indicating that the triangles in \( \mathcal{T}_{\ell,k} \) are at one level of refinement finer than those in \( \mathcal{T}_{\ell,0} \). Note that, due to the projection in both the local and global refinement steps, there will be variations in edge lengths and triangle areas that would not be present for refinement in the plane. To continue, we define \( \mathcal{T}_{\ell,k} \) to be the elements created in the \( k \)th local refinement step, uniformly refined \( \ell \) times. Following the aforementioned argument, the representative meshwidth for \( \mathcal{T}_{\ell,k} \) is \( h_k \approx h_{k-1}/2 \), and the elements in \( \mathcal{T}_{\ell,k} \) can be seen to be uniformly refined \( k \) times relative to the global mesh \( \mathcal{T}_{\ell,0} \).

It is important for what follows to emphasize that each patch, \( \mathcal{T}_{\ell,k} \), contains all elements that are covered by \( \mathcal{T}_{\ell,k+1} \) (and all finer patches), but at the same level of resolution as the nonoverlapped triangles in \( \mathcal{T}_{\ell,k} \). To properly define the triangles active in composite meshes of varying levels of refinement, we also define the “overlap” between \( \mathcal{T}_{\ell,k} \) and \( \mathcal{T}_{\ell,k+1} \) as those elements in \( \mathcal{T}_{\ell,k} \) that are covered by \( \mathcal{T}_{\ell,k+1} \). Formally, one way to express this is as

\[
\mathcal{T}'_{\ell,k} = \{ T \in \mathcal{T}_{\ell,k} \mid \text{all nodes of } T \text{ are also nodes of triangles in } \mathcal{T}_{\ell,k+1} \}.
\]

For a given number of levels of refinement, \( \ell \), the composite mesh can then be defined as

\[
S_{\ell} = \left( \bigcup_{k=0}^{\ell-1} \left( \mathcal{T}_{\ell,k} \setminus \mathcal{T}'_{\ell,k} \right) \right) \cup \mathcal{T}_{\ell,\ell}.
\]

Although the notation for \( S_{\ell} \) would be slightly more transparent with an alternate definition for \( \mathcal{T}_{\ell,k} \setminus \mathcal{T}'_{\ell,k} \), the FAC multigrid method discussed in the following is more natural using this notation. An illustration of this definition of the composite mesh is given in Figure 3; although this figure is restricted to the plane, it matches the connectivity of the composite mesh on the northern hemisphere of the sphere using the global-then-local refinement pattern as shown in Figure 2.

![Figure 3](image-url)
refinement by a single level of refinement in each local/global step, we have the natural relationship that \( T_{\epsilon k} = T_{\epsilon-1,k+1} \), when the latter is well defined (i.e., when \( \epsilon - 1 \geq k + 1 \)), but that this relationship does not hold for general refinement patterns. As such, we cannot directly simplify the notation in Equation 4, unless we make additional assumptions.

## 4 COMPOSITE-GRID FINITE-ELEMENT DISCRETIZATION

To discretize Equation 3, we consider a piecewise linear finite-element discretization on the “composite grid” formed by the refinement process discussed earlier. Discretization then follows by the surface finite-element method (see, e.g., the work by Dziuk et al.\(^ {25} \)), which we adapt to suit the composite-grid setting.

On any given (flat) triangular face \( T \) with normal direction \( \vec{n_T} \), we define the tangential gradient on \( T \) as \( \nabla_T u = \nabla u - (\nabla u \cdot \vec{n_T}) \vec{n_T} \), where \( \nabla u \) is a smooth extension of \( u : T \rightarrow \mathbb{R} \) into a neighborhood of \( T \) in \( \mathbb{R}^3 \). We note that if \( T \) happens to lie parallel to the \( xy \) plane, for example, this reduces simply to the two-dimensional gradient, extended by zero in the \( z \)-direction. This allows us to define the weak form of the Laplace–Beltrami operator in Equation 3 on a triangulation \( S_f \) as finding \( u \in H^1(S_f) \) such that

\[
a_\epsilon(u, v) = \sum_{f \in S_f} \int_T (\nabla_T u \cdot \nabla_T v + a u v) = \sum_{f \in S_f} \int_T g_\epsilon v = \langle g_\epsilon, v \rangle_{\epsilon} \quad \forall v \in H^1(S_f),
\]

where \( H^1(S_f) \) is defined in the natural way. In what follows, we also make use of the space \( C^0(S_f) \) for functions mapping from \( S_f \) into \( \mathbb{R} \). This space is defined in the natural way both within any triangle, \( T \in S_f \), and across common edges between triangles, but extended to the nonconforming situation at boundaries between two levels of refinement in \( S_f \). In this case, continuity is still enforced on the nonconforming edges in \( S_f \), by defining arc-length parameterizations \( e_k : [0, 1] \rightarrow S_f \) and \( e_{k+1} : [0, 1] \rightarrow S_f \) of the single (coarse) edge and multiple (fine) edges, respectively, with \( e_k(0) = e_{k+1}(0) \) and \( e_k(1) = e_{k+1}(1) \) marking the two common nodes, and requiring that \( f(e_k(t)) = f(e_{k+1}(t)) \) for \( 0 \leq t \leq 1 \) for all functions \( f \in C^0(S_f) \). There are, of course, many possible parameterizations of the two sets of edges; we assume both \( e_k \) and \( e_{k+1} \) are arc-length parameterizations in order to impose the natural continuity condition that the function value at the midpoint of a coarse edge matches that at the middle node in a refinement of that edge.

Our surface finite-element discretization is then, defined on a triangulation \( S_f \), by taking

\[
\mathcal{V}_\epsilon = \{ u_\epsilon \in C^0(S_f) \mid \forall T \in S_f, u_\epsilon \text{ is linear on } T \},
\]

giving a standard piecewise linear approximation on each triangle. The discrete weak form over \( S_f \) is then defined by restricting Equation 5 to \( \mathcal{V}_\epsilon \), that is, finding \( u_\epsilon \in \mathcal{V}_\epsilon \) such that \( a_\epsilon(u_\epsilon, v_\epsilon) = \langle g_\epsilon, v_\epsilon \rangle_{\epsilon} \) for all \( v_\epsilon \in \mathcal{V}_\epsilon \). Note that in the nonconforming case, the requirement that \( u_\epsilon \in C^0(S_f) \) “slaves” any hanging nodes in the triangulation to take the interpolated values at corresponding points on the unrefined side of the interface. This weak form presumes, however, that the source term \( g_\epsilon \) is defined naturally on \( S_f \), which is not the case because \( g : S \rightarrow \mathbb{R} \) in the Laplace–Beltrami Equation in Equation 3. In order to achieve the best possible order of approximation, we must ensure that \( g_\epsilon \) approximates \( g \) in a suitable way (see, e.g., studies by Dziuk et al.\(^ {25} \) and Dziuk\(^ {46} \)). In what follows, we extend the source function \( g \) to its approximation \( g_\epsilon : S_f \rightarrow \mathbb{R} \) by projecting values from \( S \) onto \( S_f \). In particular, for a point, \( x \in S_f \), we draw the straight-line path from the origin through \( x \) to a point \( \omega \in S \) and fix \( g_\epsilon(x) = g(\omega) \).

When \( \alpha > 0 \), both the weak form in Equation 5 and its restriction to \( \mathcal{V}_\epsilon \) have unique solutions (see, e.g., studies by Dziuk et al.\(^ {25} \) and Dziuk\(^ {46} \)). In the case where \( \alpha = 0 \), we must impose an additional constraint on both \( u_\epsilon \) and \( g_\epsilon \) in order to ensure unique solvability. We extend the natural conditions from the case of planar domains, requiring that \( \langle g_\epsilon, 1 \rangle_{\epsilon} = 0 \), yielding a consistent linear system for the finite-element approximation. To ensure the uniqueness of \( u_\epsilon \), we can either fix the value of \( u_\epsilon \) at a point in \( S_f \) or impose a similar orthogonality condition. In the experiments to follow (only when \( \alpha = 0 \)), we choose the former and fix the value of \( u_\epsilon \) at the south pole to be zero.

In what follows, we make use of two decompositions of \( \mathcal{V}_\epsilon \). A natural overlapping decomposition is to write \( \mathcal{V}_\epsilon = \bigoplus_{k=0}^{\epsilon} \mathcal{V}_{\epsilon,k} \), where

\[
\mathcal{V}_{\epsilon,k} = \{ u_k \in C^0(T_{\epsilon,k}) \mid u_k \text{ is piecewise linear on } T_{\epsilon,k} \}.
\]

To represent \( \mathcal{V}_\epsilon \) without overlap, we introduce the subspaces

\[
\mathcal{W}_{\epsilon,k} = \{ u_k \in C^0(T'_{\epsilon,k}) \mid u_k \text{ is piecewise linear on } T'_{\epsilon,k} \},
\]
noting that \( \mathcal{W}_{\ell,k} \subset \mathcal{V}_{\ell,k} \) (because \( T'_{\ell,k} \subset T_{\ell,k} \)) and that \( \mathcal{W}_{\ell,k} \subset \mathcal{V}_{\ell,k+1} \) (because \( T_{\ell,k+1} \) covers \( T'_{\ell,k} \)). An equivalent, but nonoverlapping, decomposition of \( \mathcal{V}_\ell \) is then given by

\[
\mathcal{V}_\ell = \mathcal{V}_{\ell,0} \oplus_{k=0}^{\ell-1} (\mathcal{V}_{\ell,k} \setminus \mathcal{W}_{\ell,k}),
\]

where boundaries between levels of refinement are implicitly constrained to match the coarser level representation due to the requirement that \( \mathcal{V}_\ell \subset C^0 (S_\ell) \). Considering the decomposition of the composite meshes in Figure 3, we note that \( \mathcal{W}_{\ell,k} \) is the piecewise linear finite-element space over the shaded patches \( T'_{\ell,k} \), whereas \( \mathcal{V}_{\ell,k} \) is the piecewise linear finite-element space over the entire patch \( T_{\ell,k} \), including the shaded patch. In what follows, we will make use of two natural finite-element interpolation operators, \( Q_\ell : \mathcal{W}_{\ell,k} \to \mathcal{V}_{\ell,k} \) and \( P_\ell : \mathcal{W}_{\ell,k} \to \mathcal{V}_{\ell,k+1} \). Although we can consider these both as operators acting on the function spaces and as matrices acting on vectors of coefficients for basis expansions, in what follows we will exclusively consider the matrix representation of these operators.

### 5 FAC METHOD

To solve the discretized linear system on the composite mesh \( S_\ell \),

\[
\text{Find } u_\ell \in \mathcal{V}_\ell \text{ such that } a_\ell (u_\ell, v_\ell) = \langle g_\ell, v_\ell \rangle_\ell \quad \forall v_\ell \in \mathcal{V}_\ell,
\]

we use the FAC methodology.\(^{15-18}\) This is based on the overlapping composite-grid decomposition of \( S_\ell \), where each “patch” \( T_{\ell,k} \) serves as a level in the multigrid hierarchy.

There are many variations on the FAC methodology. The original FAC algorithm of McCormick\(^{15}\) and McCormick et al.\(^{16}\) is based on a decomposition of the mesh into overlapping levels, with the discrete weak form solved sequentially on each level (from coarsest to finest). To break this sequentiality, asynchronous FAC (AFAC) methods were proposed and studied by Lee et al.\(^{17,18}\) and McCormick et al.\(^{47}\) where approximations for each level are computed in parallel and the differences of these approximations are used to correct the composite-grid approximation. AFACx replaces the solves on each level within AFAC with the application of simple relaxation approaches. Here, we consider an “FAC V-cycle,” where each level in the composite grid is treated by a simple relaxation scheme (lexicographical Gauss–Seidel), but the processing is sequential, as in standard multigrid. To enable the V-cycle to be used as a preconditioner for the conjugate gradient (CG) algorithm, we use symmetric relaxation ordering and a V(1,1) cycling scheme.

To fully describe the FAC V-cycle, we define two hierarchies of discrete operators corresponding to the constituent pieces of the discretized weak form in Equation 7. Restricting to \( T_{\ell,k} \), we define the system matrix \( A_\ell \) and right-hand side \( g_\ell \) by the weak form

\[
\sum_{T \in T_{\ell,k}} \int_T \nabla u \cdot \nabla v + a uv = \sum_{T \in T_{\ell,k}} \int_T gv,
\]

for \( u, v \in \mathcal{V}_{\ell,k} \), where, again, we project \( g : S \to \mathbb{R} \) onto each triangle, \( T \), in order to define the integrals on the right-hand side. Similarly, restricting to \( T'_{\ell,k} \), we define the system matrix \( \tilde{A}_\ell \) and right-hand side \( g_\ell \) by the weak form

\[
\sum_{T \in T'_{\ell,k}} \int_T \nabla u \cdot \nabla v + a uv = \sum_{T \in T'_{\ell,k}} \int_T gv,
\]

for \( u, v \in \mathcal{W}_{\ell,k} \), with the corresponding projection on the right-hand side.

Although the multigrid interpolation operators were defined earlier, some subtlety exists in the FAC restriction step in order to form the corresponding residual on each coarser level after relaxation on the next finer level of the composite-grid hierarchy. To simplify this, consider restriction from level \( k \) in the hierarchy. After relaxation on level \( k \), the \( \mathcal{V}_{\ell,k} \) residual is naturally defined as \( r_k = g_k - A_k u_k \), where \( u_k \) is the discrete approximation after relaxation. Two residuals must be considered over \( \mathcal{V}_{\ell,k-1} \), that over \( \mathcal{W}_{\ell,k-1} \), \( \tilde{r}_{k-1} = g_{k-1} - \tilde{A}_{k-1} u_{k-1} \), and that over \( \mathcal{V}_{\ell,k-1} \) itself, \( r_{k-1} = g_{k-1} - A_{k-1} u_{k-1} \), where \( u_{k-1} \) and \( u_{k-1} \) are the existing approximations on that level. Again referring to the decomposition of the composite grid pictured in Figure 3, we note that \( \tilde{r}_{k-1} \) is the residual over the shaded patch on level \( k - 1 \), whereas \( r_{k-1} \) is the residual over the entire coarse level. Note that \( r_{k-1} \) and \( \tilde{r}_{k-1} \) should agree over the interior of the patch \( T'_{\ell,k-1} \), but they will differ along the boundary of \( T'_{\ell,k-1} \) (and only \( r_{k-1} \) will be defined on \( T_{\ell,k-1} \setminus T'_{\ell,k-1} \)). The restriction step can, thus, be expressed as

\[
r_{k-1} \leftarrow r_{k-1} + Q_{k-1} (P_{k-1}^T r_k - \tilde{r}_{k-1}).
\]

In this expression, the term \( P_{k-1}^T r_k \) is the natural restriction of the (updated) grid \( k \) residual to \( T'_{\ell,k-1} \) (the shaded patch in Figure 3), while applying \( Q_{k-1} \) to this vector simply extends it (by zero) to all of \( T_{\ell,k-1} \). The difference \( r_{k-1} - Q_{k-1} \tilde{r}_{k-1} \) is, in contrast,
zero within the patch, and equal to the existing residual on $\mathcal{T}_{\ell,k-1} \setminus \mathcal{T}'_{\ell,k-1}$. Thus, the restriction step in Equation 8 appropriately accounts for the combined purpose of mesh $\mathcal{T}_{\ell,k-1}$, to provide both a correction to the refined mesh $\mathcal{T}_{\ell,k}$ and an approximation to the solution on the nonoverlapped mesh, $\mathcal{T}_{\ell,k-1} \setminus \mathcal{T}'_{\ell,k-1}$. Proper treatment at the boundary of the patch is provided by the natural finite-element interpolation operators, $Q_{k-1}$ and $P_k$.

Pseudocode for the FAC V-cycle is given in Algorithm 1. On the “pre-relaxation” side of the cycle, this essentially matches a standard multigrid V-cycle, with a relaxation step and a fine-grid residual calculation; the only difference is in the restriction step, which takes the form in Equation 8, rather than a standard restriction operation. On the global mesh $\mathcal{T}_{\ell,0}$, a standard geometric multigrid V(1,1)-cycle is used as the solver, progressing through coarsened meshes $\mathcal{T}_{\ell-1,0}$, $\mathcal{T}_{\ell-2,0}$, … until the original base mesh is reached, where either an exact solve is performed or a few sweeps of relaxation are used to approximate the coarsest grid solution. The “post-relaxation” side of the cycle is also standard, for both the global mesh and the composite grid. In order to ensure symmetry of the resulting cycle, we use a forward–backward ordering of relaxation, with forward lexicographical sweeps used for pre-relaxation and reverse lexicographical sweeps used for post-relaxation. When used as a preconditioner for the CG algorithm, the right-hand side vectors, $g_k$ and $g_{k+1}$, on all meshes are replaced with the current CG residual on each mesh, and zero initial guesses are used for $u_k$ and $\tilde{u}_k$ on all meshes. We use a single FAC V(1,1)-cycle as the preconditioner when doing so.

Algorithm 1 FAC V-Cycle Pseudocode

```
for $k = \ell', \ell' - 1, \ldots 1$ do
    Relax on interior of $\mathcal{T}_{\ell,k}$
    Compute residual on $\mathcal{T}_{\ell,k}$
    Update residual for $\mathcal{T}_{\ell,k-1}$
end for

Regular V-cycle on $\mathcal{T}_{\ell,0}$

for $k = 1, 2, \ldots, \ell'$ do
    Interpolate and add correction from $\mathcal{T}_{\ell,k-1}$
    Relax on interior of $\mathcal{T}_{\ell,k}$
end for
```

To illustrate Algorithm 1, we consider the cycle on $S_2$, as shown in the bottom line of Figure 3. The algorithm begins with a sweep of relaxation on $\mathcal{T}_{2,2}$, after which a residual is calculated. The restriction step to $\mathcal{T}_{2,1}$, as in Equation 8, can be thought of in two pieces. Outside of the patch $\mathcal{T}_{2,1}'$, we seek to directly approximate the solution and, so, define the current residual there by $r_1 - Q_1 r_1$. Within the patch, we aim to compute a correction to the current approximation and, thus, augment this by the restricted residual over the patch, $Q_1 P_2 r_2$. A similar sequence is then followed on $\mathcal{T}_{2,1}$, with a sweep of relaxation, a residual calculation, and restriction via Equation 8 to the global mesh $\mathcal{T}_{2,0}$. After a V-cycle on the global mesh, a correction over the patch $\mathcal{T}_{2,0}$ is interpolated to $\mathcal{T}_{2,1}$. After relaxation on $\mathcal{T}_{2,1}$, a correction over the patch $\mathcal{T}_{2,1}'$ is interpolated to $\mathcal{T}_{2,2}$, followed by relaxation on this finest level.

6 | NUMERICAL EXPERIMENTS

For the numerical results presented here, we use a C++ implementation of the FAC algorithm, using the modular finite-element library MFEM$^{48}$ for managing the discretization, mesh, and interpolation operators. All tests were run in serial on a single node of the Tufts High-Performance Computing Research Cluster, using a 2.2-GHz Intel Xeon CPU and 128 GB available RAM, with code compiled with gcc using full optimizations. For comparison, we present numerical results using the AMG package BoomerAMG.$^{49}$ Unless otherwise stated, we use the preconditioned CG (PCG) implementation in Hypre$^{50}$ to accelerate convergence, as this is found to be more efficient than using the FAC algorithm as a stationary iteration. For all tests, we use a fixed relative stopping tolerance based on the standard estimate of the $\ell^2$-norm of the error within the preconditioned CG algorithm, requiring that

$$\left( z_{\ell}^{(m)} \right)^T r_{\ell}^{(m)} \leq 10^{-8} \left( z_{\ell}^{(0)} \right)^T r_{\ell}^{(0)} $$

where $r_{\ell}^{(m)}$ is the full residual on the finest level (level $\ell$) of the hierarchy after $m$ iterations, and $z_{\ell}^{(m)}$ is the so-called preconditioned residual, obtained by applying the FAC V-cycle to $r_{\ell}^{(m)}$.

We note that for the tests with stationary iterations, this is a nonstandard stopping criterion, because neither of these vectors is typically stored explicitly in a multigrid code. However, we use this here for consistency between tests.
6.1 | Smooth source term

Our first numerical experiments consider the reaction–diffusion analog on the sphere,

\[-\frac{\tau}{2} \Delta \omega u(\omega) + u(\omega) = g(\omega),\]

(9)

where \( g(\omega) \) is calculated so that the solution is given by

\[u(\theta, \phi) = \frac{1}{2} \sum_{i=0}^{\infty} (2i+1)e^{-i(i+1)\tau/2}P_i(\cos \theta),\]

where \( \omega \in S \) is written as \( \omega = (\theta, \phi) \) for polar angle \( \theta \) and azimuthal angle \( \phi \), \( P_i(x) \) is the \( i \)th Legendre polynomial on \([-1, 1]\], and \( \tau = 0.001 \). This solution arises as the fundamental solution of the heat equation on the sphere, with initial data as a (scaled) delta function at the north pole (\( \theta = 0 \)), after time \( \tau \) has elapsed. Thus, Equation 9 models the behavior of the first time-step in an implicit-in-time integration of the heat equation with time-step \( \tau \).

As in the planar case, the fundamental solution to the heat equation is in \( C^\infty(S) \); thus, we expect an optimal order of approximation already on a uniform grid. Because we use a piecewise linear approximation, we expect second-order convergence in the \( L_2 \) error. However, because the solution is highly peaked around \( \theta = 0 \), we expect to see some benefit, in terms of the accuracy per degree of freedom, in the adaptive meshes considered earlier. Figure 4 plots the relative \( L_2 \) error for a uniform mesh and two refined meshes, generated by the local-then-global (“standard”) and global-then-local (“reverse”) refinement patterns discussed earlier and pictured in Figures 1 and 2, respectively. All three methods clearly offer convergence with errors proportional to the inverse of the number of degrees of freedom. For the uniform mesh, this is equivalent to second-order convergence because the surface of the sphere is two dimensional. As expected, the locally refined meshes offer better accuracy per degree of freedom by concentrating elements toward the poles, where the greatest variation in the solution occurs. At 12.5 million degrees of freedom using the local-then-global refinement pattern, the resulting solution has a relative error in this measure of \( 1.9 \times 10^{-4} \), compared with that of \( 3.3 \times 10^{-3} \) when using 16.7 million degrees of freedom on a uniform mesh, showing a clear advantage for the refined meshes.

The advantages in accuracy of the locally refined meshes are, of course, only of benefit if there is no significant additional cost associated with the solution of the resulting linear systems. As a baseline for comparison, Tables 1 and 2 present results for geometric multigrid with rediscretization used for coarse-grid operators on the uniform mesh discretization, both as a standalone iteration (Table 1) and as a preconditioner for CG (Table 2). In these tables, and those that follow, we report grid sizes measured by the number of degrees of freedom in the resulting linear system (i.e., the number of nodes in the mesh), the relative error in the discrete solution, measured in the \( L_2 \) norm (as displayed in Figure 4), the setup time for the geometric multigrid method, and both the solve time and number of iterations for either the stationary multigrid iteration or the multigrid-preconditioned CG algorithm. The setup time includes the total time required to sequentially refine the octahedral base mesh to each level of refinement in the multigrid hierarchy, discretize the equation on each of these meshes, and determine the grid-transfer operators between levels of the hierarchy. Two noticeable conclusions are possible from this data. First, that both setup and solve times scale roughly linearly with the number of degrees of freedom, as is expected. Second, the total time-to-solution is dominated by the setup time, which is consistently a factor of six or more times more costly than the solve time. This illustrates a fundamental difficulty of working with triangulations of surfaces, that mesh refinement and discretization require substantially more effort.
than they do on planar regions, even when using uniform meshes. Comparing the solve times in Tables 1 and 2, we see that there is a clear advantage to using the multigrid method as a preconditioner, although the stationary iteration also offers quite acceptable performance.

Results for the stationary FAC algorithm applied to the locally refined meshes using the local-then-global refinement pattern are shown in Table 3. Corresponding results for multigrid-preconditioned CG are shown in Tables 4 (for the local-then-global refinement pattern) and 5 (for the global-then-local refinement pattern). Most notable from these results is that there is no particular “penalty” to pay in terms of FAC setup and solve times when using these refined meshes. For both refinement patterns, setup time scales roughly linearly with the number of degrees of freedom, as do solve times; again, there is clearly a higher cost to the setup phase of the algorithm, by the same factor of about six or more, except on the finest mesh of the

---

**TABLE 1**  Finite-element accuracy and solver performance for example with smooth solution on uniform meshes, using multigrid (MG) as a stationary iteration

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Relative $L_2$ error</th>
<th>Setup time</th>
<th>Solve time</th>
<th># of MG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>16386</td>
<td>2.84e+00</td>
<td>0.12</td>
<td>0.016</td>
<td>10</td>
</tr>
<tr>
<td>65538</td>
<td>7.89e-01</td>
<td>0.49</td>
<td>0.084</td>
<td>13</td>
</tr>
<tr>
<td>262146</td>
<td>2.07e-01</td>
<td>2.09</td>
<td>0.42</td>
<td>15</td>
</tr>
<tr>
<td>1048578</td>
<td>5.25e-02</td>
<td>8.73</td>
<td>1.78</td>
<td>15</td>
</tr>
<tr>
<td>4194306</td>
<td>1.32e-02</td>
<td>35.55</td>
<td>7.78</td>
<td>15</td>
</tr>
<tr>
<td>16777218</td>
<td>3.31e-03</td>
<td>143.2</td>
<td>33.89</td>
<td>15</td>
</tr>
</tbody>
</table>

---

**TABLE 2**  Finite-element accuracy and solver performance for example with smooth solution on uniform meshes, using multigrid-preconditioned CG

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Relative $L_2$ error</th>
<th>Setup time</th>
<th>Solve time</th>
<th># of PCG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>16386</td>
<td>2.84e+00</td>
<td>0.12</td>
<td>0.014</td>
<td>7</td>
</tr>
<tr>
<td>65538</td>
<td>7.89e-01</td>
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<td>8</td>
</tr>
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<td>262146</td>
<td>2.07e-01</td>
<td>2.06</td>
<td>0.27</td>
<td>8</td>
</tr>
<tr>
<td>1048578</td>
<td>5.25e-02</td>
<td>8.68</td>
<td>1.26</td>
<td>9</td>
</tr>
<tr>
<td>4194306</td>
<td>1.32e-02</td>
<td>35.26</td>
<td>5.46</td>
<td>9</td>
</tr>
<tr>
<td>16777218</td>
<td>3.31e-03</td>
<td>143.2</td>
<td>23.46</td>
<td>9</td>
</tr>
</tbody>
</table>

---

**TABLE 3**  Finite-element accuracy and fast adaptive composite-grid solver performance for example with smooth solution on meshes following the local-then-global refinement pattern, using multigrid (MG) as a stationary iteration

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Relative $L_2$ error</th>
<th>Setup time</th>
<th>Solve time</th>
<th># of MG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>48894</td>
<td>5.05e-02</td>
<td>0.40</td>
<td>0.077</td>
<td>15</td>
</tr>
<tr>
<td>196094</td>
<td>1.25e-02</td>
<td>1.63</td>
<td>0.34</td>
<td>15</td>
</tr>
<tr>
<td>785406</td>
<td>3.10e-03</td>
<td>6.84</td>
<td>1.42</td>
<td>15</td>
</tr>
<tr>
<td>3143678</td>
<td>7.72e-04</td>
<td>28.38</td>
<td>6.01</td>
<td>15</td>
</tr>
<tr>
<td>12578814</td>
<td>1.93e-04</td>
<td>114.9</td>
<td>26.02</td>
<td>15</td>
</tr>
<tr>
<td>50323454</td>
<td>4.81e-05</td>
<td>467.7</td>
<td>111.4</td>
<td>15</td>
</tr>
</tbody>
</table>

---

**TABLE 4**  Finite-element accuracy and fast adaptive composite-grid solver performance for example with smooth solution on meshes following the local-then-global refinement pattern, using multigrid-preconditioned CG

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Relative $L_2$ error</th>
<th>Setup time</th>
<th>Solve time</th>
<th># of PCG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>48894</td>
<td>5.05e-02</td>
<td>0.40</td>
<td>0.053</td>
<td>9</td>
</tr>
<tr>
<td>196094</td>
<td>1.25e-02</td>
<td>1.63</td>
<td>0.23</td>
<td>9</td>
</tr>
<tr>
<td>785406</td>
<td>3.10e-03</td>
<td>6.85</td>
<td>0.97</td>
<td>9</td>
</tr>
<tr>
<td>3143678</td>
<td>7.72e-04</td>
<td>28.37</td>
<td>4.11</td>
<td>9</td>
</tr>
<tr>
<td>12578814</td>
<td>1.93e-04</td>
<td>114.6</td>
<td>17.63</td>
<td>9</td>
</tr>
<tr>
<td>50323454</td>
<td>4.81e-05</td>
<td>468.5</td>
<td>74.97</td>
<td>9</td>
</tr>
</tbody>
</table>
local-then-global refinement pattern, with approximately 50 million degrees of freedom, where the factor is only five. Note also that the local-then-global refinement pattern offers slightly better accuracy per degree of freedom than the global-then-local pattern. Comparing Tables 3 and 4, we again see that the stationary iteration performs well, but that there is a notable improvement in iteration counts (from 15 to 9) and solve time (a reduction of about 30% on the finest grids), when using it as a preconditioner for CG. For this reason, we only consider the multigrid-preconditioned CG algorithm in the remaining examples in this paper.

A further comparison for the efficiency of the FAC algorithm is against the performance of AMG, for which we use the BoomerAMG code from the Hypre package \(^{49,50}\) with standard parameters. Tables 6 and 7 detail solver timings and iteration counts for this comparison. For clarity, we break out the mesh-refinement and finite-element discretization times in the column labeled “Matrix/RHS setup” from the actual setup time consumed by the AMG algorithm, computing the AMG coarse meshes and interpolation operators, as well as the Galerkin coarse-grid operators. Comparing the timings for the local-then-global refinement pattern in Table 6 to those for FAC in Table 4, we see that the cost for the construction of the composite mesh and finite-element assembly of the matrix and right-hand side is roughly the same as the setup cost for the full FAC algorithm. Additional to this is the cost of the AMG setup and, then, a higher solve cost for AMG, due both to a small increase in the number of iterations and a larger increase in the cost per iteration. We note also in this case that the lower memory overhead of the FAC algorithm allows us to solve a problem with 50 million degrees of freedom using FAC, which we could not do within the memory available to us with AMG. Similar behavior is also seen for the global-then-local refinement pattern, with results for AMG in Table 7 for comparison to those in Table 5. Here, there is a consistent but anomalous penalty of about 20% in CPU time for the construction of the composite mesh and finite-element assembly within the AMG code compared to that in the FAC code. This is due to an increase in time in the composite mesh generation function within our finite-element package, MFEM, for this particular refinement pattern in the code that calls AMG. Although it should be possible to eliminate this difference, it is small enough that we have not done so in the results reported here. There is also a larger increase in solve time (that cannot be

### TABLE 5

Finite-element accuracy and fast adaptive composite-grid solver performance for example with smooth solution on meshes following the global-then-local refinement pattern, using multigrid-preconditioned CG

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Relative (L_2) error</th>
<th>Setup time</th>
<th>Solve time</th>
<th># of PCG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>24450</td>
<td>1.28e-01</td>
<td>0.19</td>
<td>0.026</td>
<td>9</td>
</tr>
<tr>
<td>98050</td>
<td>3.13e-02</td>
<td>0.77</td>
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</tr>
<tr>
<td>392706</td>
<td>7.76e-03</td>
<td>3.27</td>
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<td>9</td>
</tr>
<tr>
<td>1571842</td>
<td>1.93e-03</td>
<td>13.80</td>
<td>1.95</td>
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</tr>
<tr>
<td>6289410</td>
<td>4.81e-04</td>
<td>55.65</td>
<td>8.28</td>
<td>9</td>
</tr>
<tr>
<td>25161730</td>
<td>1.20e-04</td>
<td>228.4</td>
<td>35.64</td>
<td>9</td>
</tr>
</tbody>
</table>

### TABLE 6

Algebraic multigrid (AMG) solver performance for example with smooth solution on meshes following the local-then-global refinement pattern

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Matrix/RHS setup</th>
<th>AMG setup</th>
<th>Solve</th>
<th># of PCG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>48894</td>
<td>0.70</td>
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<td>11</td>
</tr>
<tr>
<td>196094</td>
<td>2.03</td>
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<td>0.51</td>
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</tr>
<tr>
<td>3143678</td>
<td>28.4</td>
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<td>9.75</td>
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<td>12578814</td>
<td>113.7</td>
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<td>39.8</td>
<td>12</td>
</tr>
</tbody>
</table>

### TABLE 7

Algebraic multigrid (AMG) solver performance for example with smooth solution on meshes following the global-then-local refinement pattern

<table>
<thead>
<tr>
<th># DOFs</th>
<th>Matrix/RHS setup</th>
<th>AMG setup</th>
<th>Solve</th>
<th># of PCG iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>24450</td>
<td>0.51</td>
<td>0.04</td>
<td>0.05</td>
<td>11</td>
</tr>
<tr>
<td>98050</td>
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<td>0.15</td>
<td>0.25</td>
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</tr>
<tr>
<td>392706</td>
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<td>12</td>
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<tr>
<td>1571842</td>
<td>16.81</td>
<td>2.75</td>
<td>4.83</td>
<td>12</td>
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<tr>
<td>6289410</td>
<td>67.01</td>
<td>10.8</td>
<td>19.98</td>
<td>12</td>
</tr>
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<td>267.5</td>
<td>45.39</td>
<td>90.67</td>
<td>12</td>
</tr>
</tbody>
</table>
readily eliminated) for the AMG approach compared to the local-then-global refinement pattern, about a factor of 2.5 compared to a factor of 2, coming from increased complexity in the AMG hierarchy for this refinement pattern. We note that, due to the use of aggressive coarsening, the AMG grid complexities (computed as the sum of the number of degrees of freedom on all levels of an AMG hierarchy divided by the number on the finest level) are somewhat lower than they are for the FAC algorithm, but that the AMG operator complexities (the sum of the number of nonzero elements in the matrix on all levels divided by the number on the finest level) are slightly higher, due to stencil growth in the coarse-grid operators.

Overall, two main themes emerge from these results. First, there is no substantial penalty in terms of computational cost per degree of freedom for solving the problems on refined meshes using FAC compared to geometric multigrid on uniform meshes. This, then, validates the use of refined meshes for problems with highly localized solutions, such as the example considered here. Furthermore, there is a clear benefit to the use of FAC in this setting over the use of a black-box approach, such as AMG. In particular, the practical costs of the full setup of the FAC algorithm are naturally included in those of assembling the composite-mesh finite-element discretization. This coupled with the greatly improved efficiency of the resulting FAC solve phase, in comparison to either the combined AMG setup and solve phases or just the AMG solve phase, shows the true algorithmic potential of using a semistructured approach with measured speedup factors of two to four times and notably lower memory requirements.

6.2 Impulse source term

In contrast to the example with a smooth solution considered earlier, we now consider the problem

\[-\Delta u = \sum_{i=0}^{\infty} \frac{2i + 1}{4\pi} P_i(\cos \theta) - \frac{1}{4\pi},\]

\[u(\pi, \phi) = 0 \quad \forall \phi,\]

which has an exact solution

\[u = -\frac{1}{2\pi} \log\left( \sin\left( \frac{\theta}{2} \right) \right).\]

We note that the series term on the right-hand side represents the Dirac delta function at the north pole, and should be interpreted in the distributional sense, rather than as a convergent sum. In the distributional sense, its integral over \(S\) is one and, so, the constant shift on the right-hand side is necessary so that the total source term then integrates to zero for consistency. As noted earlier, the condition imposed at the south pole, \(u(\pi, \phi) = 0\), guarantees uniqueness; at the discrete level, this is imposed just as a typical Dirichlet boundary condition would be, albeit at a single point. Figure 5 displays the relative error measured in the \(L_2\) norm between the finite-element solutions and the continuum solution, calculated by omitting those elements adjacent to the north pole to avoid numerical issues evaluating the solutions there. As expected, we now see the clear benefit in rate of convergence given by the adaptive meshes in contrast to the first-order convergence of the uniform mesh (because the mesh size scales as the inverse square root of the number of degrees of freedom in the uniform mesh discretization).

![Figure 5](image-url)
Tables 8 and 9 provide timing statistics for FAC and AMG applied to this problem on the refined mesh using just the local-then-global refinement pattern for brevity. Aside from a slight increase in the number of iterations for each (from 9 to 12 for FAC and 12 to 15 for AMG), the details strongly mirror those for the smooth solution presented earlier. Notably, the setup times between the two approaches are roughly equal, with a small added amount for FAC, noticeable only on the finest meshes (corresponding to the time needed to generate the FAC interpolation operators on top of the composite-mesh finite-element assembly). The solve times for FAC are again about a factor of one half of those for AMG, or about one third if the AMG setup and solve times are counted together. As before, the lower memory footprint of FAC allows the solution of the largest problem size, with over 50 million degrees of freedom, whereas AMG could not solve this problem within the memory available.

### 6.3 Two-spot problem

To demonstrate the flexibility of the FAC algorithm, we next consider an example with a smooth solution that changes rapidly at two points in the domain. To construct this example, we again consider Equation 9, now with source function \( g(\omega) \) chosen so that the solution is given by

\[
  u(\theta, \phi) = \frac{1}{2} \sum_{i=0}^{\infty} (2i + 1)e^{-i(i+1)\tau/2}P_i(\sin \theta \cos \phi) + \frac{1}{2} \sum_{i=0}^{\infty} (2i + 1)e^{-i(i+1)\tau/2}P_i(\cos \theta),
\]

with \( \tau = 0.001 \). This solution arises as a linear combination of fundamental solutions to the heat equation with initial data as the sum of two delta functions, one at the north pole (\( \theta = 0 \)) and one on the equator (\( \theta = \pi/2, \phi = 0 \)). Thus, this example models the behavior of the first time-step in an implicit-in-time integration of the heat equation with time-step \( \tau \) and a “two-spot” initial condition. To match the variation in the solution, we use a corresponding two-spot refinement pattern, with sample meshes pictured in Figure 6. In this example, we focus on the global-then-local refinement pattern, simply because this allows for two disjoint regions of refinement. Note now that each refinement patch consists of two (disconnected) pieces, but that the FAC algorithm as given earlier remains well defined in such a case.

FAC and AMG solver performances are detailed in Tables 10 and 11, respectively, with accuracy proportional to the number of degrees of freedom verified by the relative \( L_2 \) errors reported in Table 10. Similarly to the global-then-local refinement pattern results for the example in Section 6.1, these results show a consistent anomaly in the time required for the mesh construction and finite-element assembly in the AMG results. Aside from this, the same general conclusions can be drawn, with the FAC solve phase being about 2.5 times faster than that of the AMG solve phase, and over four times faster than the combined AMG setup and solve phases for the finest mesh with over 33.5 million degrees of freedom.
CONCLUSIONS AND FUTURE WORK

In this paper, we extend the FAC algorithm to finite-element discretizations on triangulated surfaces, particularly focused on structured-grid refinements of the sphere. The FAC algorithm is shown to be well suited for such problems, with the composite-grid transfer operation naturally expressed in terms of two finite-element interpolation operators. For problems with localized sources, improved finite-element accuracy is seen using refined meshes, including restoration of effectively second-order accuracy for a problem with singular source.

Given the wide body of research currently ongoing into the accurate discretization of PDEs posed on surfaces or manifolds, a natural direction for future work is the comparison of FAC and multigrid algorithms for surface finite-element discretizations with solution algorithms for the CPM or other discretization approaches. Additionally, the work presented here was motivated by the development of fast algorithms for solution of the Boltzmann transport equation in (or close to) the Fokker–Planck limit. A key piece of future work is the coupling of the FAC algorithm presented here with more general scattering kernels and a suitable spatial discretization to address that problem in more detail.

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