# Spacetime Meshing with Adaptive Refinement and Coarsening* 

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#### Abstract

We propose a new algorithm for constructing finite-element meshes suitable for spacetime discontinuous Galerkin solutions of linear hyperbolic PDEs. Given a triangular mesh of some planar domain $\Omega$ and a target time value $T$, our method constructs a tetrahedral mesh of the spacetime domain $\Omega \times[0, T]$ in constant running time per tetrahedron in $\mathbb{R}^{3}$ using an advancing front method. Elements are added to the evolving mesh in small patches by moving a vertex of the front forward in time. Spacetime discontinuous Galerkin methods allow the numerical solution within each patch to be computed as soon as the patch is created. Our algorithm employs new mechanisms for adaptively coarsening and refining the front in response to a posteriori error estimates returned by the numerical code. A change in the front induces a corresponding refinement or coarsening of future elements in the spacetime mesh. Our algorithm adapts the duration of each element to the local quality, feature size, and degree of refinement of the underlying space mesh. We directly exploit the ability of discontinuous Galerkin methods to accommodate discontinuities in the solution fields across element boundaries.


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## 1. INTRODUCTION

Scientists and engineers use conservation laws and hyperbolic partial differential equations to model transient, wavelike behavior in bodies or spatial control volumes. Example applications are numerous, ranging from the Euler equations for compressible gas dynamics, to the Schrödinger equation for time-dependent density functional theory in quantum mechanics, to the equations of elastodynamics in seismic analysis. Closed-form solutions are typically unavailable for these problems, so analysts usually resort to numerical approximations. However, the continuum solutions can exhibit strongly nonlinear behavior as well as shocks and other discontinuities that make this class of numerical problems particularly challenging.

Finite element methods are a good option for solving these problems, especially when the geometry of the analysis domain is complicated. In the standard semi-discrete approach, a finite element mesh discretizes space to generate a system of ordinary differential equations in time that is then solved by a time-marching integration scheme. Most methods enforce a uniform time step size over the entire spatial domain. This approach can be very costly for strongly graded grids, because the allowable time step size is limited by the global minimum element diameter. However, physical causality only implies a local limit on the step size, so algorithms that use a nonuniform time step size can substantially improve computational efficiency.

Dynamic adaptive refinement and coarsening is essential for capturing solution features that move with traveling interfaces and shock fronts. Each discrete remeshing operation requires a projection of the solution fields from the old grid onto the newly adapted mesh. These projections can be costly and inconvenient to compute, and they introduce


Figure 1. An input space mesh and the resulting spacetime mesh computed by Tent Pitcher [8]
significant error. A more continuous approach to adaptive refinement could eliminate these data projections and the error they produce.

Spacetime discontinuous Galerkin finite element methods $[31,6,15,36,35,21]$ are a relatively new alternative to semi-discrete methods. (For further background on discontinuous Galerkin methods in general, we refer the reader to Cockburn, Karniadakis, and Shu [7].) Two features distinguish spacetime discontinuous Galerkin methods from conventional finite element models. First, spacetime discontinuous Galerkin methods work with meshes that cover the entire spacetime analysis domain. For example, simulation of evolving behavior on a three-dimensional spatial domain requires a four-dimensional spacetime mesh. The spacetime discontinuous Galerkin algorithm weakly enforces the governing equations over each spacetime element, eliminating the need for a separate time integration procedure. The second distinguishing feature is the use of discontinuous basis functions with support on individual elements for representing the physical fields. In contrast to conventional finite element methods that use continuous bases, this approach eliminates artificial coupling between adjacent elements when the mesh satisfies certain causality constraints. It also guarantees exact satisfaction of the relevant balance laws on every spacetime element.

Building on ideas from earlier specialized algorithms [15, 28, 32, 34], Üngör and Sheffer [33] and Erickson et al. [8] developed the first algorithm to build graded spacetime meshes over arbitrary simplicially meshed spatial domains, called 'Tent Pitcher'. Unlike most traditional approaches, the Tent Pitcher algorithm does not impose a fixed global time step on the mesh, or even a local time step on small regions of the mesh. Rather, it produces a fully unstructured simplicial spacetime mesh, where the duration of each spacetime element depends on the local feature size and quality of the underlying space mesh. See Figure 1.

Given a triangular mesh of some planar domain $\Omega$, and a target time value $T$, the Tent Pitcher algorithm meshes the spacetime domain $\Omega \times[0, T]$ in $\mathbb{R}^{3}$ using an advancing front method. Elements are added to the evolving mesh in small patches by moving a vertex of the front forward in time. The advance in time is limited by causality, to ensure that the solution in the new patch depends only on boundary data and solution data from previously computed patches. The
use of discontinuous basis functions avoids artificial coupling with subsequent patches. Thus, the spacetime discontinuous Galerkin solution can be computed locally within each new patch as soon as it is created. Provided the initial ground mesh has constant degree, each patch contains only a constant number of elements and can therefore be solved in constant time. The time to generate the spacetime mesh and compute the numerical solution is, therefore, linear in the number of spacetime elements. Moreover, because patches with no causal relationship can be solved independently, the Tent Pitcher algorithm is uniquely well-suited for parallel implementation. In Section 2, we give a new and more intuitive formulation of the Tent Pitcher algorithm and review its theoretical guarantees.

In this paper, we introduce new mechanisms into the Tent Pitcher algorithm that adaptively refine or coarsen the advancing front in response to a posteriori error estimates computed by the numerical code. The front at any stage of the meshing and solution process is a hierarchical subdivision of the original space mesh. A change in the front induces a corresponding refinement or coarsening of future elements in the spacetime mesh. We refine the front using the classical newest vertex bisection method of Sewell [27] and Mitchell [18, 19, 20]; we coarsen the front only by reversing earlier refinements. These modifications generate non-conforming spacetime meshes; two adjacent spacetime elements may not share a common face. However, because discontinuous Galerkin methods directly accommodate discontinuities in the solution field across element boundaries, they do not require conforming meshes or a data projection operation. We describe the remeshing operations in detail in Section 3. In practice, the meshes we compute effectively track shocks and moving interfaces through spacetime; regions of the front are refined in response to an approaching shock, and then coarsened again after the shock passes.

The original Tent Pitcher algorithm of Üngör and Sheffer [33] required every angle in the input mesh to be acute; if the ground mesh contains an obtuse angle, their algorithm can get stuck. Erickson et al. [8] remove this limitation by imposing an additional constraint, beyond those due to causality, on the maximum height of any tent. This so-called progress constraint, which depends on the shape of the underlying space elements, guarantees a lower bound on the minimum height of the tent. Adapting this progress con-
straint to support the changing geometry of the front is the main technical contribution of this paper; see Section 4.

In Section 5 we describe a $2 \mathrm{D} \times$ time linear elastodynamics simulation computed with the help of our meshing algorithm. Finally, we conclude the paper by describing our ongoing work and open problems in spacetime meshing.

## 2. SPACETIME MESHING

The formulation of our spacetime meshing problem relies on the notions of domain of influence and domain of dependence for hyperbolic boundary value problems. We say that a point $\hat{p}$ in spacetime depends on another point $\hat{q}$ if the values of the salient physical fields (e.g., temperature, velocity, pressure, stress, momentum) at $\hat{p}$ depend on the field values at $\hat{q}$; that is, if changing the conditions at $\hat{q}$ can affect the conditions at $\hat{p}$. The domain of influence of $\hat{p}$ is the set of points that depend on $\hat{p}$; symmetrically, the domain of dependence is the set of points that $\hat{p}$ depends on.

If the governing equations of the underlying problem are linear and the material properties are homogeneous and isotropic, the domains of influence and dependence are circular cones with a common apex $\hat{p}$. This double cone can be described by a scalar wave speed $c(\hat{p}) \in \mathbb{R}$, which specifies how quickly the radius of the cones grows as a function of time. In this paper, we consider only linear problems, where the wave speed is uniform and isotropic over the entire spacetime domain. In this case, we can choose an appropriate time scale, independent of the spatial scale, so that $c(\hat{p})=1$ everywhere. For more general problems, the wave speed can vary with direction and as a function of inhomogeneous physical parameters, and might even depend nonlinearly on the solution.

We say that one spacetime element $\Delta$ depends on another spacetime element $\triangle^{\prime}$ if any point $\hat{p} \in \triangle$ depends on any point $\hat{q} \in \triangle^{\prime}$. The solution can be computed in each element one at a time, following any linear extension of the dependency partial order. Alternatively, the solutions within any anti-chain of elements can be computed in parallel.

Although discontinuous Galerkin methods impose no $a$ priori restrictions on the primitive shapes of individual elements, it is convenient to work with simplicial elements. We say that a facet $F$ of a tetrahedral element is causal if it separates the cone of influence from the cone of dependence at every point on $F$ (Figure 2), or equivalently, if $\|\nabla F\| \leq 1$. If a facet is causal, physical information flows in only one direction across that facet. To construct an efficient tetrahedral mesh, we group elements into patches, each with a constant number of elements. The Tent Pitcher algorithm ensures that the boundary facets between patches are causal by construction; we refer to this requirement as the causality constraint (called the cone constraint in Erickson et al. [8]).

In general, the internal facets between elements within a single patch are not causal. Since information can flow in both directions across these facets, the physical response in adjacent elements is coupled, and all of the elements within a patch must therefore be solved simultaneously. However, if we assume polynomial basis functions of bounded degree, the resulting linear system still has bounded size and can therefore be solved in constant time. Thus, the total time to compute the numerical solution is still linear in the number of elements.


Figure 2. The causality constraint: The facet separates the cone of influence (above) from the cone of dependence (below).

Richter [22] observed that undesirable numerical dissipation increases in certain discontinuous Galerkin methods as the slopes of boundary facets decrease below the local wave speed. Thus, our goal is to construct an efficient mesh comprised of patches, each containing a small number of simplices, such that the boundary facets of each patch are as close as possible to the cone constraint without violating it.

### 2.1 Advancing-Front Spacetime Meshing

Given a triangulated planar domain $\Omega$ and a target time value $T$, Tent Pitcher constructs a tetrahedral mesh of the spacetime domain $\Omega \times[0, T]$. The algorithm is designed as an advancing front procedure, which alternately constructs a new patch of elements and invokes a spacetime discontinuous Galerkin finite-element method to compute the solution within that patch.

At every iteration, the front is the graph of a continuous time function $t: \Omega \rightarrow \mathbb{R}$, such that within every triangle, $t$ is linear and $\|\nabla t\| \leq 1$, where $\nabla t$ denotes the gradient of $t$. We will assume that the initial time function is constant, but more general initial conditions are also permitted. The front is a terrain whose facets correspond to triangles in the underlying space mesh. To advance the front, the algorithm chooses an arbitrary local minimum vertex $\hat{p}=(p, t(p))$ from the front and moves it forward to a new point $\hat{p}^{\prime}=\left(p, t^{\prime}(p)\right)$ where $t^{\prime}(p)>t(p)$. The volume between the new and old fronts is called a tent. We decompose the tent into simplices sharing the edge $\hat{p} \hat{p}^{\prime}$ and pass these elements, along with the inflow elements just below the tent, to a DG solver. When the algorithm has several choices for the vertex to advance next, we apply one of several heuristics; some heuristics appear to work better in practice than others. The algorithm is free to advance a vertex that is not a local minimum; however, in that case, a finite amount of progress cannot always be guaranteed.

### 2.2 Causality and Progress Constraints

To complete the description of Tent Pitcher, all that remains is to describe how to compute the new time values for each vertex to be pitched, or less formally, how high to pitch each tent. Each triangle in the ground mesh imposes constraints on the time values at each of its vertices. When we pitch a tent over a local minimum vertex $\hat{p}$, the new time value $t^{\prime}(p)$ is simply the largest value that satisfies the constraints for every triangle adjacent to $p$ in the ground mesh.

Thus, it suffices to consider the case where the ground mesh consists of a single triangle $\triangle p q r$. At any stage of the algorithm, the front consists of a single triangle $\triangle \hat{p} \hat{q} \hat{r}$


Figure 3. Pitching tents in spacetime.
whose vertices have time values $t(p), t(q)$, and $t(r)$. Suppose without loss of generality that $t(p) \leq t(q) \leq t(r)$ and we want to advance $\hat{p}$ forward in time. We must choose the new time value to satisfy the causality constraint $\|\nabla t\| \leq 1$. Erickson et al. [8] show that this constraint can be expressed by the following inequality:

$$
\begin{align*}
t^{\prime}(p) \leq t(q) & +\frac{t(r)-t(q)}{|q r|^{2}}(\overrightarrow{q p} \cdot \overrightarrow{q r})  \tag{1}\\
& +\sqrt{1-\left(\frac{t(r)-t(q)}{|q r|}\right)^{2}} d(p, q r)
\end{align*}
$$

Here, $d(p, q r)$ denotes the distance from $p$ to the line $q r$.
Üngör and Sheffer [33] proved that if every angle in $\triangle p q r$ is acute, then exactly meeting the causality constraint guarantees that further advancement is always possible. However, if $\angle p q r$ is obtuse, then lifting $\hat{p}$ as far as possible may prevent us from lifting $\hat{q}$ in the next iteration without violating causality. To avoid this problem, Erickson et al. [8] impose an additional progress constraint, which limits the gradient of the time function restricted to the edges touching the obtuse angle.

The progress constraint can be visualized easily using a circle diagram; see Figure 4. Think of the gradient vector $\nabla t$ as a point in the plane; the causality constraint restricts this point to the unit disk centered at the origin. We can partition this disk into three wedges by the inward normal vectors of the edges of $\triangle p q r$. Each wedge is associated with a vertex of the triangle; for example, the wedge for $p$ is bounded by vectors normal to the edges $p q$ and $p r$. The gradient vector $\nabla t$ lies inside $p$ 's wedge if and only if $p$ has


Figure 4. A triangle and its circle diagram. To guarantee causality, the gradient vector $\nabla t$ must stay inside the circle; to guarantee progress, $\nabla t$ must stay out of the shaded forbidden zones.
the smallest time value, and pitching $p$ moves the point $\nabla t$ along the inward normal vector to the edge $q r$, or in other words, parallel to the vector that does not bound the current wedge.

To guarantee that the algorithm never gets stuck, we must ensure that we can always move the point $\nabla t$ out of its current wedge; otherwise, the lowest vertex will never progress past its neighbor. If $\Delta p q r$ has an obtuse angle, this requirement creates "forbidden zones" inside the two obtuse wedges, bounded by lines normal to the edges adjacent to the obtuse angle. The progress constraint is simply to stay out of these forbidden zones. We can express this constraint algebraically as the inequality $t^{\prime}(p) \leq t(r)+(1-\varepsilon) d(p, q r)$, where $0<\varepsilon \leq 1 / 2$ is a tunable parameter.

## 3. HIERARCHICAL MESH REFINEMENT

Our adaptive algorithm uses the newest-vertex bisection refinement method, originally developed by Sewell [27], later adapted by Mitchell [18, 19, 20] in the context of multigrid methods, and still later studied and generalized to three dimensions by Bänsch [3]. This method is similar to the more common longest-edge refinement methods introduced by Rosenberg and Stenger [26] and popularized by Rivara [23, 24, 25] and others [12, 1, 29]. For a general discussion of mesh refinement methods, we refer the the reader to surveys by Jones and Plassmann [9] and by Bern and Plassmann [4].

We call the newest vertex of a triangle its apex and the opposite edge its base. Initially, one vertex of each triangle in the mesh is chosen arbitrarily as its apex. Newest-vertex bisection replaces a triangle with two smaller triangles, each with half the area of the original triangle, obtained by bisecting along the line segment through the apex and the midpoint of the base. The new vertex introduced at the midpoint is the newest vertex of both smaller triangles.

The descendants of any marked triangle under newestvertex bisection fall into only eight homothetic classes-see Figure 5-and there are only four directions along which the triangle or any of its descendants can be further bisected. These four directions are parallel to the three edges of the triangle and the bisecting segment. Any two triangles in the same equivalence class have corresponding newest vertices.

When we refine one triangle in a mesh, we may be forced to refine other nearby triangles in order to maintain a conforming triangulation. Suppose vertex $a$ is the apex of a $\triangle a b c$ that is bisected (Figure 6). If $b c$ is not a boundary edge then some neighboring triangle $\triangle c b e$ shares the edge


Figure 5. Newest vertex refinement.


Figure 6. Refinement propagating to neighboring triangles.
$b c$. To maintain a triangulation, $\triangle c b e$ must be bisected also. If $b c$ is not the base of $\Delta c b e$ then the child of $\triangle c b e$ sharing edge $b c$ must be bisected as well, and the bisection of $\triangle c b e$ will propagate recursively; see Figure 6. It is easy to prove that this propagation terminates, regardless of which vertex is chosen as the apex of each triangle. The propagation path touches every triangle in the worst case, but in practice, the propagation path usually has small constant length; see Suárez et al. [30] for an analysis of a similar refinement algorithm.

The entire refinement sequence can be expressed as a sequence of edge bisection and quad flip operations. The former operation bisects an edge shared by one or two triangles, and the latter replaces one diagonal of a convex quadrilateral with the other diagonal.

Coarsening is the opposite of refinement; we coarsen locally by undoing a single edge bisection. Unlike refinement, coarsening does not require propagation further into the mesh to maintain a conforming triangulation, although one coarsening step may make other coarsenings possible. In particular, if we refine a triangle and then immediately coarsen, we can (but need not) coarsen along the entire refinement path.

Newest-vertex bisection never subdivides any angle of a triangle more than once. Therefore, the degree of any vertex in the mesh at most doubles as a result of refinement. Moreover, given a triangulation with maximum degree $d$ with vertices in general position, if the vertex with largest angle in each triangle is chosen as the apex, then any refinement of the initial triangulation has maximum degree at $\operatorname{most} \max \{d+5,8\}$.

## 4. MAKING TENT PITCHER ADAPTIVE

To incorporate adaptivity into our meshing algorithm, we make a small change to the main loop. At each iteration, just as before, we choose a local minimum vertex $\hat{p}$ of the front,
move it forward in time to create a tent, and pass the tent and its inflow data to the spacetime DG solver. We assume that the solver also computes an a posteriori estimate of its own numerical error. If the error within any element of the tent is above some threshold, the solver rejects the patch, at which point the meshing algorithm throws away the tent and refines the facets of the front whose elements had high error. (Alternately, we could refine the facets until their diameter is smaller than a target length scale computed by the solver.) Note that this refinement may propagate far outside the neighborhood of $\hat{p}$. We accept the numerical solution and update the front only if the error within every element of the patch is acceptable.

On the other hand, the error estimate within an element may fall below some second threshold, indicating that the mesh is finer than necessary to compute the desired result. In this case, the DG solver marks the outflow face of that element as coarsenable. We can coarsen four facets of the front into two only if they are the result of an earlier refinement, they are all marked as coarsenable, and each pair of triangles to be merged is coplanar. To make coarsening possible, our algorithm tries to make coarsenable siblings coplanar, by lowering the top of the tent. However, to avoid very thin elements, we only accept the lower tent if its height is above some threshold. If the lower tent is accepted and its outflow faces are still marked coarsenable, then we coarsen the front.

### 4.1 New Progress Constraints

The more subtle and important change in our algorithm is the introduction of new progress constraints. In the earlier non-adaptive algorithm [8], the progress constraint was a function of the shape of the underlying space elements. In our new algorithm, the shape of the underlying element is subject to change; each triangle in the current front may be the result of any number of coarsenings or refinements since the last time any ancestor or descendant of that triangle was last pitched. Consequently, our progress constraints must take into account the shapes of all possible descendants of a triangle simultaneously. This requirement motivates our choice of newest-vertex refinement; because the descendants of any triangle fall into only a finite number of homothety classes, we have only a finite number of simultaneous progress constraints.

We can visualize these constraints by referring back to our circle diagram; see Figure 7. A minimal set of progress constraints can be obtained by simply taking the union of the non-adaptive progress constraints of the eight shapes in the hierarchy. These constraints are not necessarily sufficient, however, since it may be impossible for the gradient to leave a wedge for one triangle without violating a progress constraint for a different triangle. Fortunately, we can avoid these conflicts by strengthening some of the constraints.

Omitting the geometric derivation, we formally describe the resulting constraints as follows. Fix two real numbers $0<\delta, \varepsilon<1$. For any triangle $\triangle a b d$ with apex $a$, we define the diminished width of $\triangle a b d$ as follows:

$$
w_{\varepsilon}(\dot{a} b d):=\min \left\{\begin{array}{c}
(1-\varepsilon) d(a, b d), \\
(1-\delta) d(b, a d), \\
(1-\delta) d(d, a b)
\end{array}\right\}
$$

The first distance is measured from the apex to the opposite edge and is scaled differently from the other two altitudes.


Figure 7. Merging eight progress constraints into one.

This definition extends recursively to any descendants of $\triangle a b d$ obtained by newest-vertex subdivision; in the interest of readability, we will always list the vertices of any triangle with the apex first. Now let $c$ be the midpoint of $b d$ and let $e$ be the midpoint of $a b$, as illustrated below. We express our new progress constraints algebraically by limiting the difference in time values along each edge in the subdivided triangle, as follows.


These constraints apply recursively to all descendants of the triangle $\triangle a b d$-for example, we also enforce the constraint $|t(c)-t(e)| \leq w_{\varepsilon}(\dot{c} a b)$-but these recursive constraints are redundant, since they limit the gradient of the time function in exactly the same direction as one of the four constraints above.

Now suppose we are pitching a triangle $\triangle p q r$, where $t(p) \leq$ $t(q) \leq t(r)$. Let $s$ be the midpoint of $q r$; let $u$ be the midpoint of $p r$; and let $v$ be the midpoint of $p q$; see Figure 8. Depending on which of the three vertices is marked as the apex, the new time value $t^{\prime}(p)$ is bounded as a result of these progress constraints in the three different ways enumerated below. Notice that when $p$ is not the apex, lifting $p$ also lifts either $u$ or $v$, so progress constraints along edges $q u$ or $r v$ also indirectly limit $t^{\prime}(p)$.

If $p$ is the apex:

$$
t^{\prime}(p) \leq \min \left\{\begin{array}{l}
t(q)+2 w_{\varepsilon}(\dot{s} p r),  \tag{2}\\
t(s)+w_{\varepsilon}(\dot{p} q r), \\
t(r)+2 w_{\varepsilon}(\dot{s} p q)
\end{array}\right\}
$$


(a)

(b)

(c)

Figure 8. Cases for implementing the combined progress constraints: (a) $p$ is the apex, (b) $q$ is the apex, (c) $r$ is the apex.

If $q$ is the apex:

$$
t^{\prime}(p) \leq \min \left\{\begin{array}{l}
t(q)+2 w_{\varepsilon}(\dot{u} q r),  \tag{3}\\
t(r)+4 w_{\varepsilon}(v q u), \\
2 t(q)-t(r)+2 w_{\varepsilon}(\dot{q} p r)
\end{array}\right\}
$$

If $r$ is the apex:

$$
t^{\prime}(p) \leq \min \left\{\begin{array}{l}
t(q)+4 w_{\varepsilon}(\dot{u} r v)  \tag{4}\\
\left.t(r)+2 w_{\varepsilon} \dot{v} q r\right), \\
2 t(r)-t(q)+2 w_{\varepsilon}(\dot{r} p q)
\end{array}\right\}
$$

### 4.2 Proof of Positive Progress

In this section, we prove that for any triangle $\Delta p q r$ that satisfies the causality and progress constraints, we can lift $\hat{p}$ at least $\Delta$ above $\hat{q}$ without violating either constraint, where $\Delta$ is a positive function of the shape of the triangle and the scaling parameters $\varepsilon$ and $\delta$. The causality constraint and the progress constraints will each imply a different upper bound on the maximum progress $\Delta$ that we can guarantee.

Lemma 1. If $\triangle \hat{p} \hat{q} \hat{r}$ satisfies the causality and progress constraints, we can set $t^{\prime}(p)=t(q)+\varepsilon \cdot d(p, q r)$ without violating the causality constraint.

Proof: First, we show that the progress constraints imply $t(r)-t(q)<(1-\varepsilon)|q r|$ no matter which vertex is the apex. If $p$ is the apex, we have the following series of inequalities:

$$
\begin{aligned}
t(r)-t(q) & \leq 4 w_{\varepsilon}(\dot{u} p s) \\
& \leq 4(1-\varepsilon) d(u, p s) \\
& =2(1-\varepsilon) d(r, p s) \\
& \leq 2(1-\varepsilon)|r s|=(1-\varepsilon)|q r|
\end{aligned}
$$

Here, the first inequality is the actual progress constraint, the second follows from the definition of diminished width, and the rest follow from the geometry of the recursively bisected triangle. Similarly, if $q$ is the apex, we have

$$
\begin{aligned}
t(r)-t(q) & \leq 2 w_{\varepsilon}(\dot{u} p q) \\
& \leq 2(1-\varepsilon) d(u, p q) \\
& =(1-\varepsilon) d(r, p q) \leq(1-\varepsilon)|q r|,
\end{aligned}
$$

and if $r$ is the apex, we have

$$
\begin{aligned}
t(r)-t(q) & \leq 2 w_{\varepsilon}(\dot{v} p r) \\
& \leq 2(1-\varepsilon) d(v, p r) \\
& =(1-\varepsilon) d(q, p r) \leq(1-\varepsilon)|q r| .
\end{aligned}
$$

Solving for $\varepsilon$ gives us the inequality

$$
\varepsilon \leq 1-\frac{t(r)-t(q)}{|q r|} \leq \sqrt{1-\left(\frac{t(r)-t(q)}{|q r|}\right)^{2}}
$$

which implies that

$$
\begin{aligned}
t^{\prime}(p) & =t(q)+\varepsilon \cdot d(p, q r) \\
& \leq t(q)+\sqrt{1-\left(\frac{t(r)-t(q)}{|q r|}\right)^{2}} d(p, q r) \\
& \leq t(q)+\frac{t(r)-t(q)}{|q r|^{2}}(\overrightarrow{q p} \cdot \overrightarrow{q r}) \\
& +\sqrt{1-\left(\frac{t(r)-t(q)}{|q r|}\right)^{2}} d(p, q r) .
\end{aligned}
$$

This is exactly the expression of the causality constraint in equation (1).

Lemma 2. If $0<\varepsilon<\delta<(1+\varepsilon) / 2<1$ and $\triangle \hat{p} \hat{q} \hat{r}$ satisfies the causality and progress constraints, we can set $t^{\prime}(p)=t(q)+\Delta$ without violating any progress constraint, where $\Delta>0$ is a function of the triangle $\Delta p q r$ and the parameters $\varepsilon$ and $\delta$.

Proof: As in the previous proof, we have three cases to consider. In each case, we will derive positive upper bounds on the possible value of $\Delta$; setting $\Delta$ to the minimum of these upper bounds will satisfy the conditions of the lemma.

First, suppose $p$ is marked. Since $t(r) \geq t(s) \geq t(q)$, the relevant progress constraint (2) is satisfied if

$$
\Delta \leq \min \left\{2 w_{\varepsilon}(\dot{s} p r), w_{\varepsilon}(\dot{p} q r), 2 w_{\varepsilon}(\dot{s} p q)\right\}
$$

Similarly, suppose $r$ is marked. Since $t(r) \geq t(q)$, we have $2 t(r)-2 t(q) \geq 0$, which implies that the relevant progress constraint (4) is satisfied if

$$
\Delta \leq \min \left\{4 w_{\varepsilon}(\dot{u} r v), 2 w_{\varepsilon}(\dot{v} q r), 2 w_{\varepsilon}(\dot{r} p q)\right\}
$$

Finally, suppose $q$ is marked. By the inductive hypothesis, edge $q r$ already satisfies its individual progress constraint $t(r) \leq t(q)+2 w_{\varepsilon}(\dot{u} p q)$, which implies that $t(q)-t(r) \geq$ $-2 w_{\varepsilon}(\dot{u} p q)$. Since $t(r) \geq t(q)$, the relevant progress constraint (3) is satisfied if

$$
\Delta \leq \min \left\{2 w_{\varepsilon}(\dot{u} q r), 4 w_{\varepsilon}(\dot{v} q u), 2 w_{\varepsilon}(\dot{q} p r)-2 w_{\varepsilon}(\dot{u} p q)\right\} .
$$

The last term in this expression is the only one that is not obviously positive. To prove that $X=w_{\varepsilon}(\dot{q} p r)-w_{\varepsilon}(\dot{u} p q)$
is in fact bounded away from zero, we expand the definition of diminished width. We have $X=\min \{A, B, C\}$, where

$$
\begin{aligned}
A & =(1-\varepsilon) d(q, p r)-\min \left\{\begin{array}{l}
(1-\varepsilon) d(u, p q), \\
(1-\delta) d(p, q u), \\
(1-\delta) d(q, p u)
\end{array}\right\} \\
& \geq(1-\varepsilon) d(q, p r)-(1-\delta) d(q, p u) \\
& =(\delta-\varepsilon) d(q, p r),
\end{aligned}
$$

$$
\begin{aligned}
B & =(1-\delta) d(r, p q)-\min \left\{\begin{array}{l}
(1-\varepsilon) d(u, p q), \\
(1-\delta) d(p, q u), \\
(1-\delta) d(q, p u)
\end{array}\right\} \\
& \geq(1-\delta) d(r, p q)-(1-\varepsilon) d(u, p q) \\
& =(1-2 \delta+\varepsilon) d(u, p q),
\end{aligned}
$$

$$
\begin{aligned}
C & =(1-\delta) d(p, q r)-\min \left\{\begin{array}{l}
(1-\varepsilon) d(u, p q), \\
(1-\delta) d(p, q u), \\
(1-\delta) d(q, p u)
\end{array}\right\} \\
& \geq(1-\delta)(d(p, q r)-\min \{d(p, q u), d(q, p u)\}) .
\end{aligned}
$$

Since by assumption $\varepsilon<\delta<(1+\varepsilon) / 2$, we clearly have $A>0$ and $B>0$. We prove that $C>0$ as follows:

$$
\begin{aligned}
d(p, q r) & =\frac{2 \operatorname{Area}(\triangle p q r)}{|q r|} \\
& =\frac{2 \operatorname{Area}(\triangle p q u)}{|u v|} \\
& >\frac{2 \operatorname{Area}(\triangle p q u)}{\max \{|q u|,|p u|\}} \\
& =\min \{d(p, q u), d(q, p u)\} .
\end{aligned}
$$

The key observation is that the bisector segment $u v$ must be shorter than at least one of the two edges $p u$ and $q u$.

Theorem 3. Given a triangular mesh $\Omega$ in the plane and a target time value $T$, our algorithm builds a finite tetrahedral mesh of the spacetime domain $\Omega \times[0, T]$, in constant running time per element, provided each triangle is refined only a finite number of times.

## 5. EXPERIMENTAL RESULTS

We have implemented our adaptive spacetime meshing algorithm in $1 \mathrm{D} \times$ time and $2 \mathrm{D} \times$ time. We present here an example to demonstrate its application to a scientific problem of practical complexity: the phenomenon of crack-tip wave scattering within an elastic solid subjected to shock loading.

Figure 9 shows a stationary crack embedded in a rectangular plate subjected to a spatially uniform tensile traction on its top and bottom edges. We prescribe traction-free boundary conditions on the left and right edges and planestrain conditions overall. The load history approximates a step function; as depicted in Figure 9, the load intensity $\sigma$ rises from 0 to a maximum value $\sigma^{*}=10$ over the ramp time $t_{\text {ramp }}=0.002$, so that the ramp covers $3.25 \%$ of the width of the plate. This suddenly-applied loading generates a linear shock wave that moves from the top loaded edge toward the crack surface at the bottom. When the shock reaches the crack, it reflects back up toward the top of the plate and scatters in a circular wave pattern from the crack-tip.

The material properties of the plate are Young's modulus $E=10$, Poisson's ratio $\nu=0.3$, and mass density $\rho=1$.


Figure 9. Crack geometry and loading for the crack-tip wave scattering problem.

Since the domain and initial conditions are symmetric, we explicitly model only the upper right quadrant of the domain, specifying symmetric boundary conditions at the left and bottom edges. The initial space mesh contains only 16 triangles; we depend on the adaptive meshing algorithm to generate all of the necessary refinement. We use a complete quadratic polynomial basis ( $1 x y t x y$ yt $t x x^{2} y^{2} t^{2}$ ) to represent the displacement field within each tetrahedral spacetime element.

A dissipation-based error indicator drives the adaptive meshing scheme. The energy dissipated by each patch is compared to the total energy flux into the patch; zero dissipation indicates exact energy balance. We set a normalized dissipation target, dissipation/energy influx $=1.0 \times 10^{-7}$. Elements with dissipation more than $20 \%$ above the target are rejected and marked for mandatory refinement. Elements with dissipation more than $20 \%$ below the target are accepted and marked as eligible for coarsening. We also specify a minimum dissipation of $6.0 \times 10^{-10}$ and a minimum element size of $5.0 \times 10^{-5}$, below either of which no further refinement is allowed.

The final spacetime mesh contains $17,052,587$ tetrahedra, clustered into $2,964,477$ patches. During the solution process, $3,679,040$ patches containing 21, 956,046 tetrahedra were computed and solved; approximately $20 \%$ of these patches were rejected, causing the front to be refined. The ratio of the largest to smallest element diameters in the spacetime mesh is 1024 , reflecting 20 levels of refinement. The normalized dissipation for the entire spacetime analysis domain is $0.079 \%$.

Figure 10 shows two pairs of visualizations of the computed solution within the upper right quadrant of the plate. Each image illustrates the intersection of the spacetime data set with a constant-time plane. The images on the left display two axes of the solution data, by mapping strain-energy-density to a color field on a logarithmic scale, and mapping the magnitude of the velocity field to a height field. The images on the right show the intersection of the unstructured tetrahedral spacetime mesh with the constant-time plane, together with the strain-energy-density color field.

The first pair of images in Figure 10 show the shock front as it approaches the crack surface for the first time. The pattern of mesh refinement accurately tracks the fine details of the traveling waves, including the primary shock front as well as dilitation and shear waves generated along the traction-free top edge.

The second pair of images show the initial crack-tip wave scattering pattern some time after the shock first arrives at the crack. The circular scattering pattern is reflected in the mesh refinement; the outer annulus of less intense refinement indicates the extent of the faster dilitational wave component, while the circle of more intense refinement indicates the slower shear wave. A singular strain-energy-density field is visible at the crack tip, a stationary feature that generates a persistent region of intense mesh refinement.

Figure 11 shows two views of the entire spacetime mesh at roughly the same time depicted in the second row of Figures 10. The vertical axis is time, and the pattern of refinement on the vertical faces follows closely the spacetime trajectories of the shocks; see Figure 11(a). The cone-shaped region of intense mesh refinement in Figure 11(b) covers the domain of influence of the initial crack-tip scattering event. The asymmetry of the cone is consistent with the form of the singular crack-tip field, which vanishes in the plane of the crack ahead (to to the right) of the tip.

As expected, our experimental results show a significant improvement in the quality of the solution as a result of adaptivity, especially near discontinuities in the solution or its derivative. Also, we were able to achieve a better solution without using a fine mesh everywhere, which would have resulted in a massive increase in computation time.

## 6. CURRENT AND FUTURE WORK

We implemented a sequential version of our adaptive Tent Pitcher algorithm for one- and two-dimensional spatial domains which we used to solve linear PDEs with constant wavespeed and nonlinear problems where the maximum wavespeed is bounded. We also prototyped a parallel version for $1 \mathrm{D} \times$ time that leverages the mutual independence of any two patches over the same front. Since the patches can be solved independently of each other, we were able to solve them simultaneously on different processors. We used the CHARM + + parallel language and runtime system [11] developed by the Parallel Programming Group at the University of Illinois [10] to manage interprocessor communication, load balancing, and the automatic dispatch and scheduling of patches to be solved. In our preliminary experiments, we observed a parallel speedup of about 20 . We observed that the speedup was limited mainly by the mesh generation, which was not running in parallel.

We are also investigating spacetime discontinuous Galerkin methods for nonlinear conservation laws. In such problems, the wavespeed is not constant throughout the spacetime domain. Instead, the wavespeed is one of the physical parameters governed by the underlying PDE, and can therefore change in unpredictable ways. We have extended Tent Pitcher to adapt the number and duration of spacetime elements to changing wavespeeds, for problems defined over one- and two-dimensional spatial domains. As in the current paper, the major theoretical bottleneck lies in developing appropriate progress constraints. We have implemented our algorithm for nonlinear problems in 1D $\times$ time, and we are currently implementing the algorithm in $2 \mathrm{D} \times$ time. We expect to publish these results in a subsequent paper.

We are also working on combining the mesh refinement with adaptivity to local wavespeeds. We plan to implement this combined, completely adaptive algorithm in parallel.


Figure 10. Adaptive solution of the crack-tip wave scattering problem. (a)-(b) The shock front approaching the crack surface from above. (c)-(d) The initial scattering pattern after reflection off the crack surface.


Figure 11. Spacetime mesh for the crack-tip wave scattering problem: (a) overview; (b) detail of the initial crack-tip wave scattering pattern.

Finally, we would like to extend all the current results to higher dimensions, specifically to $3 \mathrm{D} \times$ time. For the actual mesh refinement, we can directly use the generalization of newest-vertex bisection to three dimensions by Bänsch [3] and others $[2,5,13,14,16,17]$. Again, the main theoretical hurdle in higher dimensions is deriving minimal progress constraints that guarantee that the front can always advance. Another concern is to be able to describe the constraints in such a fashion that they are easy to implement. Up to our current work in $2 \mathrm{D} \times$ time, we have encountered only a constant number of linear constraints per element. It seems likely that such constraints that can be described analytically should suffice in higher dimensions, but it is not clear that they are the best possible.

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