# A fast sweeping method for static convex Hamilton-Jacobi equations<sup>1</sup>

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#### Abstract

We develop a fast sweeping method for static Hamilton-Jacobi equations with convex Hamiltonians. Local solvers and fast sweeping strategies apply to structured and unstructured meshes. With causality correctly enforced during sweepings numerical evidence indicates that the fast sweeping method converges in a finite number of iterations independent of mesh size. Numerical examples validate both the accuracy and the efficiency of the new methods.

# 1 Introduction

We consider a class of static Hamilton-Jacobi equations of the following form,

$$\begin{cases} H(\mathbf{x}, \nabla T(\mathbf{x})) = 1, & \mathbf{x} \in \Omega \setminus \Gamma, \\ T(\mathbf{x}) = g(\mathbf{x}), & \mathbf{x} \in \Gamma \subset \Omega, \end{cases}$$
(1.1)

where  $g(\mathbf{x})$  is a positive, Lipschitz continuous function,  $\Omega$  is an open, bounded domain in  $\mathbb{R}^d$  and  $\Gamma$  is a subset of  $\Omega$ ;  $H(\mathbf{x}, \mathbf{p})$  is Lipschitz continuous in both arguments, and it is convex in the second argument. If  $H(\mathbf{x}, \mathbf{p}) = |\mathbf{p}|H(\mathbf{x}, \mathbf{p}/|\mathbf{p}|) = |\mathbf{p}|F(\mathbf{x})$ , then the eikonal equation for isotropic wave propagation results; in general, the equation is anisotropic in the sense that the wave speed varies along different directions, hence the anisotropic eikonal equation results.

Such equations arise in a multitude of applications, ranging from seismic waves, crystal growth, robotic navigation, and optimal control, to name just a few. Consequently, it is necessary to develop accurate and efficient methods for computing numerical solutions to this nonlinear boundary value problem. In this paper we extend the fast sweeping method [1, 32, 27, 12, 13, 23] to tackle the above static Hamilton-Jacobi equation on triangular meshes.

Mathematical foundation for the well-posedness of the equation traces back to the theory of viscosity solution [5] and computability of such viscosity solution by monotone finite difference methods is established in [6]. There are two crucial tasks in developing an efficient numerical method for such type of equations: one is designing a local

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solver or discretization scheme that can capture causality of the underlying partial differential equation (PDE), and the other is solving the resulting large system of nonlinear equations after discretization. Many finite difference schemes are available for discretizations on rectangular grids, such as Godunov schemes. On structured grids, one-sided or upwind approximation of partial derivatives can be constructed more readily. In most of these schemes causality and consistency are coupled and enforced simultaneously in the local discretization. Due to nonlinearity of the PDE, the corresponding local solver may be quite complicated. For example Godunov solvers for general Hamiltonians can be very difficult.

To tackle such anisotropic eikonal equations in seismics, Dellinger [7] extended an upwind finite difference method [29] to compute the first-arrival based viscosity solutions in an anisotropic medium; Qin and Schuster [24] proposed a wavefront expansion method, and it is based on Huygens' principle and computes the first-arrival traveltimes associated with seismic energy propagating at the group velocity. As pointed out in Dellinger and Symes [8], an anisotropic medium is different from an isotropic medium in that for an isotropic medium the ray velocity vector (ie, the group velocity vector, or the characteristic direction) has the same direction as the (negative) traveltime gradient (ie, the phase velocity vector), which enables us to use the traveltime gradient as a reliable indicator of energy flow (and thus the causality) in propagating the traveltime field [29, 24, 28, 25, 32], while for an anisotropic medium this is no longer true. Therefore one may compute wrong solutions by extending fast marching methods designed for isotropic eikonal equations to anisotropic eikonal equations without taking into account the above essential differences as demonstrated in [26].

Based on the above observation, Qian and Symes [17, 19, 20, 21] proposed a paraxial formulation for the static Hamilton-Jacobi equation by formulating a relation between the characteristic direction and the traveltime gradient direction, so that fast, efficient and accurate methods with linear complexity can be obtained easily; furthermore, Qian, Symes and Dellinger [22] made further improvement by removing the paraxial assumption.

On the other hand, Sethian and Vladimirsky [26] designed ordered upwind methods for the above static Hamilton-Jacobi equation. The spirit of their single-pass method is the following: at a considered node which is to be updated, first one estimates the possible numerical domain of dependency by using the so-called anisotropic coefficient [26], the accepted solution and the mesh size; secondly one uses the so-called controltheoretic update-from-a-single-simplex formula [9] to evaluate a tentative value at the standing node by taking the minimum among all possible values resulting from all the possible virtual simplexes constructed from its numerical domain of dependency; thirdly one accepts the smallest value of all the considered nodes to maintain causality; lastly one maintains the lists of accepted solutions and considered nodes. The resulting ordered upwind methods have the computational complexity of  $O(\eta M \log M)$ , where  $\eta$  is the anisotropic coefficient depending on the Hamilton-Jacobi equation, and M is the total number of mesh points.

As an iterative method for Hamilton-Jacobi equations, the fast sweeping method was originated in Boue and Dupis [1], and its first PDE formulation was for implicit and non-parametric shape reconstruction from unorganized points using a variational level set method [34]; Zhao [32] proved the O(N) convergence of the method for the eikonal equation based on the Godunov Hamiltonian on Cartesian meshes. Tsai, Cheng, Osher

and Zhao [27] applied the fast sweeping methods to a class of static Hamilton-Jacobi equations based on Godunov numerical Hamiltonians on uniform meshes, and they have derived some explicit update formulae so that the Gauss-Seidel based sweeping strategy can be easily carried out; numerical examples indicate that the sweeping method has linear complexity. Kao, Osher and Tsai [12] proposed a class of fast sweeping methods for the static Hamilton-Jacobi equations based on upwind-discretizing the Bellman formula resulting from the Hamiltonian directly on uniform meshes; numerical examples also indicate that the sweeping method has linear complexity. Kao, Osher and Qian [13] have extended fast sweeping methods to deal with non-convex Hamilton-Jacobi equations based on Lax-Friedrichs numerical Hamiltonians on uniform meshes. Zhang, Zhao and Qian [31] have developed higher order fast sweeping methods based on weighted essentially non-oscillatory schemes [16, 11, 10] on uniform meshes. Zhang, Zhao and Chen [30] proposed fixed-point type sweeping methods on uniform meshes. All of the above cited fast sweeping methods are based upon uniform meshes. In [23] a class of novel fast sweeping methods was developed for isotropic eikonal equations on triangular meshes for the first time. In [3], Cecil, Osher and Qian extended the fast sweeping method to deal with level set equations on adaptive tree-based unstructured meshes. Various parallel implementations of the fast sweeping method are developed in [33]

In this paper we develop a local solver that decouples consistency and causality. This approach allows one to deal with much more general Hamiltonians and applies to both structured and unstructured grids. Since we are solving a nonlinear boundary value problem, a large nonlinear system needs to be solved after discretization. We apply the fast sweeping strategy developed in [23] to solve the system which gives an efficient and unconditionally stable iterative method. In particular the fast sweeping method is an iterative method of Gauss-Seidel type, consisting of correct causality check and alternating sweeping orders. The key point is that all characteristics can be divided into a finite number of groups and each group can be captured simultaneously by one of the orderings. Recently this methodology has been studied extensively and has also been applied successfully to other hyperbolic problems [14].

Here is the outline of this paper. In Section 2 a local solver for general convex Hamiltonian will be described. The full numerical algorithm, the fast sweeping strategy combined with our local solver, is summarized in Section 3. Explicit formulae is derived for a class of anisotropic eikonal equations in Section 4. Finally numerical examples will be shown in Section 5 to demonstrate both the efficiency and the accuracy of our method.

# 2 Local solvers

## 2.1 Some basic facts

Because equation (1.1) arises naturally from geometrical optics for wave propagation [2], without any hesitation we decide to adopt some common terminology from geometrical optics in the following presentation.

For the sake of simplicity in the following derivation we assume that H is strictly convex and homogeneous of degree one; we will comment on general cases later.

Wavefronts of the traveltime are level sets defined by

$$\{\mathbf{x}\in\Omega:T(\mathbf{x})=t,t\in\mathcal{R}\}.$$

The wavefront normal at a point  $\mathbf{x} \in \Omega$  is

$$\mathbf{n}(\mathbf{x}) = \frac{\nabla T(\mathbf{x})}{|\nabla T(\mathbf{x})|},$$

whenever the gradient of traveltime T is well-defined, where  $\mathbf{p} = \nabla T(\mathbf{x})$  is the slowness vector because it has the dimension of the reciprocal of velocity.  $V_p(\mathbf{x}) = \frac{1}{|\nabla T(\mathbf{x})|}$  is the so-called phase speed. Thus we have

$$\mathbf{p}(\mathbf{x}) = \frac{\mathbf{n}(\mathbf{x})}{V_p(\mathbf{x}, \mathbf{n}(\mathbf{x}))}, \qquad (2.1)$$

$$V_p(\mathbf{x}, \mathbf{n}(\mathbf{x})) = H(\mathbf{x}, \mathbf{n}(\mathbf{x})), \qquad (2.2)$$

$$H(\mathbf{x}, \mathbf{p}) = 1. \tag{2.3}$$

Since equation (1.1) is a nonlinear first-order equation, applying the method of characteristics to the equation in phase space along ray trajectories  $(\mathbf{x}(t), \mathbf{p}(t))$  yields

$$\frac{d\mathbf{x}}{dt} = \nabla_{\mathbf{p}} H, \tag{2.4}$$

$$\frac{d\mathbf{p}}{dt} = -\nabla_{\mathbf{x}}H,\tag{2.5}$$

$$\frac{dT}{dt} = \mathbf{p} \cdot \frac{d\mathbf{x}}{dt} = \mathbf{p} \cdot \nabla_{\mathbf{p}} H = 1.$$
(2.6)

The first equation defines the so-called group velocity vector, which points into the same direction as the characteristic (ray) direction. Its magnitude is

$$v_g(\mathbf{x}, \mathbf{p}) = \left| \frac{d\mathbf{x}}{dt} \right| = \left| \nabla_{\mathbf{p}} H \right|, \tag{2.7}$$

which is the so-called group speed depending on the position  $\mathbf{x}$  and the slowness vector  $\mathbf{p}$ , so that the group speed varies as the traveltime gradient does, implying the so-called directional dependence.

In a homogeneous anisotropic medium,  $H(\mathbf{x}, \mathbf{p}) = H(\mathbf{p})$ , the traveltime  $\Delta T$  between any given two points of distance  $\Delta d$  is computed by the following relation:

$$\Delta T = \frac{\Delta d}{v_g(\mathbf{p})},\tag{2.8}$$

where  $\mathbf{p}$  is determined implicitly by using the condition that the ray direction is known which is the unit vector along the straight line connecting the two given points. In general, we have to use a numerical procedure to compute the above traveltime; for example, see [17].

**Remark:** In an isotropic medium the group velocity vector (the ray direction) and the phase velocity vector (the traveltime gradient) are in the same direction, and the group speed and the phase speed are equal. As a result the computation of traveltime between any two points is straightforward. In an anisotropic medium those are no longer true. **Remark:** If  $\lim_{\lambda\to 0} \nabla_{\mathbf{p}} H(\mathbf{x}, \lambda \mathbf{p}) = 0$ , then the strict convexity of  $H(\mathbf{x}, \mathbf{p})$  implies that

$$(\nabla_{\mathbf{p}} H(\mathbf{x}, \mathbf{p}) - \nabla_{\mathbf{p}} H(\mathbf{x}, \lambda \mathbf{q})) \cdot (\mathbf{p} - \lambda \mathbf{q}) > 0 \text{ for } \forall \mathbf{p}, \mathbf{q};$$
(2.9)

consequently, we have

$$\mathbf{p} \cdot \nabla_{\mathbf{p}} H(\mathbf{x}, \mathbf{p}) \ge 0 \quad \text{for } \forall \mathbf{p}.$$
(2.10)

Therefore without using the homogeneity of H in  $\mathbf{p}$  we conclude that the solution value is nondecreasing along ray trajectories. Furthermore, if  $\mathbf{p} \cdot \nabla_{\mathbf{p}} H(\mathbf{x}, \mathbf{p}) > 0$ , then the solution value can be used as the running parameter along ray trajectories in the above formulation. This is the most essential condition for the fast sweeping method to work for Hamilton-Jacobi equations. Hence all the following algorithmic development applies as long as  $\lim_{\lambda\to 0} \nabla_{\mathbf{p}} H(\mathbf{x}, \lambda \mathbf{p}) = 0$  holds.

## 2.2 A local solver based on Fermat's Principle

We tackle the two-dimensional case first. We consider a triangulation  $\mathcal{T}_h$  of  $\Omega$  into nonoverlapping, nonempty and closed triangles  $\mathcal{T}$ , with diameter  $h_{\mathcal{T}}$ , such that  $\bar{\Omega} = \bigcup_{\mathcal{T} \in \mathcal{T}_h} \mathcal{T}$ . We assume that  $\mathcal{T}_h$  satisfies the following conditions:

- No obtuse triangles;
- No more than  $\mu$  triangles have a common vertex;
- $h = \sup_{\mathcal{T} \in \mathcal{T}_h} h_{\mathcal{T}} < 1;$
- $\mathcal{T}_h$  is regular: there exists a constant  $\omega_0$  independent of h such that if  $\rho_{\mathcal{T}}$  is the diameter of the largest ball  $B \subset \mathcal{T}$ , then for all  $\mathcal{T} \in \mathcal{T}_h$ ,  $h_{\mathcal{T}} \leq \omega_0 \rho_{\mathcal{T}}$ .

Therefore, equation (1.1) is solved in the domain  $\Omega$ , which has a triangulation  $\mathcal{T}_h$  consisting of triangles. We consider every vertex and all triangles which are associated with this vertex. See Figure 2.1 for a node C and its n triangles  $\mathcal{T}_1, \mathcal{T}_2, \dots, \mathcal{T}_n$ . For a typical triangle  $\triangle ABC$  we denote  $A : (x_A, y_A), B : (x_B, y_B)$  and  $C : (x_C, y_C); \ \angle A = \beta$ ,  $\ \angle B = \alpha$ , and  $\ \angle C = \gamma; \ \overline{AB} = c, \ \overline{AC} = b$ , and  $\overline{BC} = a$  are the lengths of the edges AB, AC and BC, respectively.

During the solution process we need a local solver at vertex C for each triangle; see Figure 2.2. Given the values  $T_A$  and  $T_B$  at A and B of triangle  $\triangle ABC$ , we want to calculate the value  $T_C$  at C.

If  $T_A$  and  $T_B$  are used to update  $T_C$ , then there must be a ray emanating from the segment AB and hitting point C; namely there is an F(s) located in between A and B, where s parametrizes the segment  $\overline{AB}$ : F(0) = A and F(1) = B.

According to Fermat's principle the traveltime at C is given by minimizing the functional

$$T_C(s) = sT_B + (1-s)T_A + \frac{d(s)}{v_g(C)}$$
(2.11)

with respect to s, where

$$d(s) = \overline{CF(s)} = \sqrt{b^2 + c^2 s^2 - 2bcs \cos\beta}, \qquad (2.12)$$

$$v_g(C;s) = v_g(C;T_C(s),T_A,T_B);$$
 (2.13)



Figure 2.1: Vertex  ${\cal C}$  and the local mesh.



Figure 2.2: Update the value at C in a triangle.

namely,

$$T_C = \min_{s \in [0,1]} \left\{ sT_B + (1-s)T_A + \frac{d(s)}{v_g(C;s)} \right\}.$$
 (2.14)

This is the so-called control-theoretic update-from-a-single-simplex formula as used in [9, 26].

The main difficulty in implementing this formula is that we have to compute the group speed  $v_g(C; s)$  by using the current ray direction defined by C and F(s). By the above formula we can immediately conclude that

$$T_C \ge \min\{T_A, T_B\} \tag{2.15}$$

if there exists a characteristic emanating from the segment AB to hit point C. However, since the wave front normal  $\nabla T$  does not coincide with the characteristic direction in general,  $\nabla T$  may not fall into the triangle; therefore, it is not necessarily true that

$$T_C \ge \max\{T_A, T_B\}.$$
(2.16)

These two facts are observed in our numerical examples.

**Remark:** In the special case of the eikonal equation, the wave front normal coincides with the direction of characteristics and hence either one can be used to check causality condition; since  $\nabla T$  points away from C and is in between the two sides CA and CB a causality-satisfying  $T_C$  must be larger than max{ $T_A, T_B$ }.

**Remark:** The above formulation involves optimization which is avoidable by adopting a fully Eulerian viewpoint.

# 2.3 A local solver based on an Eulerian discretization

By definition we have

$$\frac{T_C - T_A}{b} = \nabla T(C) \cdot \left(\frac{x_C - x_A}{b}, \frac{y_C - y_A}{b}\right)^t + o(h^2),$$
(2.17)

$$\frac{T_C - T_B}{a} = \nabla T(C) \cdot \left(\frac{x_C - x_B}{a}, \frac{y_C - y_B}{a}\right)^t + o(h^2), \qquad (2.18)$$

where t denotes the transpose of vectors. Furthermore we have

$$\begin{pmatrix} \frac{T_C - T_A}{b} \\ \frac{T_C - T_B}{a} \end{pmatrix} = \mathbf{P} \nabla T(C) + o(h^2), \qquad (2.19)$$

where

$$\mathbf{P} = \begin{pmatrix} \frac{x_C - x_A}{b} & \frac{y_C - y_A}{b} \\ \frac{x_C - x_B}{a} & \frac{y_C - y_B}{a} \end{pmatrix} \equiv \begin{pmatrix} \mathbf{r}_1^t \\ \mathbf{r}_2^t \end{pmatrix}.$$
 (2.20)

Assuming a linear approximation of T locally near C to ignore higher order terms and solve for  $\nabla T_C$ , we have

$$\nabla T(C) \approx \mathbf{P}^{-1} \left( \begin{array}{c} \frac{T_C - T_A}{b} \\ \frac{T_C - T_B}{a} \end{array} \right), \tag{2.21}$$

where  $\mathbf{P}^{-1} = \mathbf{P}^T \mathbf{Q}$  and

$$\mathbf{Q} = \frac{1}{\sin^2 \gamma} \left( \begin{array}{cc} 1, & -\cos \gamma \\ -\cos \gamma, & 1 \end{array} \right).$$

Inserting  $\nabla T(C)$  into the Hamilton-Jacobi equation at the mesh point C, we have a consistent discretization of the equation in the triangle  $\triangle ABC$ :

$$H(C, \nabla T(C)) \approx 1,$$
 (2.22)

$$\hat{H}\left(C, \frac{T_C - T_A}{b}, \frac{T_C - T_B}{a}\right) = 1, \qquad (2.23)$$

or

$$H(C, T_C, T_A, T_B) = 1.$$
 (2.24)

Since in general this is a nonlinear equation for  $T_C$ , we have to numerically solve the nonlinear equation to obtain  $T_C$  if  $T_A$  and  $T_B$  are given. For example, an anisotropic eikonal equation will result in a quadratic equation for  $T_C$  (see Section 4).

It is possible that we have multiple solutions of  $T_C$  when solving the nonlinear equation (2.24); thus we have to choose the one that satisfies causality, the so-called characteristic condition.

The fundamental idea is the following. If there is no solution to equation (2.24) it means that this triangle does not support a consistent  $T_C$ . If there is one or more solutions we need to check the causality condition. Using the computed  $T_C$  and the above equation, we get a  $\nabla T(C)$  which can be used to compute the ray direction  $\nabla_{\mathbf{p}} H(C, \nabla T(C))$ . The computed  $T_C$  satisfies the causality condition if the characteristic starting from C against the direction  $\nabla_{\mathbf{p}} H(C, \nabla T(C))$  intersects the line segment AB, as illustrated in Figure 2.2. If there is no  $T_C$  satisfying this causality condition it means that this triangle does not support a  $T_C$  that is both consistent and causality satisfying. If after the causality check, there are multiple  $T_C$ 's from this triangle, then we choose the smallest one using the first-arrival time principle.

Now we rigorously establish the above causality principle, the so-called characteristic condition.

Since  $H(\mathbf{x}, \mathbf{p})$  is strictly convex in the **p** argument, for a given **x** any straight line in the slowness space intersects the slowness surface defined by  $H(\mathbf{x}, \mathbf{p}) = 1$  at two points at most; see Figure 2.3. In general, there are three cases:

- 1. two different intersections;
- 2. two repeated identical intersections;
- 3. no intersection.

In the first case two outward normals  $\nabla_{\mathbf{p}} H$  at those two intersections give two possible characteristic directions according to equation (2.4) at the given point  $\mathbf{x}$ . In the second case we have two repeated identical intersections, and the outward normal at the intersection point also points into a characteristic direction at that point  $\mathbf{x}$ .

Now consider the straight line equation (2.21) parametrized by  $T_C$  in the slowness space, which is rewritten as

$$\mathbf{p} = \mathbf{r} T_C + \mathbf{r}_0 \tag{2.25}$$



Figure 2.3: A straight line intersects the convex slowness surface at two points at most.

where

$$\mathbf{r} = \mathbf{P}^{-1} \begin{pmatrix} b^{-1} \\ a^{-1} \end{pmatrix} = \frac{a - b \cos \gamma}{a \ b \sin^2 \gamma} \mathbf{r}_1 + \frac{b - a \cos \gamma}{a \ b \sin^2 \gamma} \mathbf{r}_2, \tag{2.26}$$

$$\mathbf{r}_0 = \mathbf{P}^{-1} \begin{pmatrix} -T_A \ b^{-1} \\ -T_B \ a^{-1} \end{pmatrix} = \frac{b \ T_B \ \cos \gamma - a \ T_A}{a \ b \ \sin^2 \gamma} \mathbf{r}_1 + \frac{a \ T_A \ \cos \gamma - b \ T_B}{a \ b \ \sin^2 \gamma} \mathbf{r}_2.$$
(2.27)

This straight line intersects the slowness surface in exactly the same way as described above in terms of the three cases; therefore we are able to establish the following causality principle:

The first case: two different outward normals. These two outward normals must have *either* opposite signs in the second component, which correspond to downgoing rays and upgoing rays, respectively, *or* have opposite signs in the first component which correspond to left-going and right-going rays, respectively.

When the straight line is vertical or horizontal, these claims have been proved in [21]; moreover, one can generalize the proof to handle the general case that the straight line is inclined.

Consider an acute triangle  $\triangle ABC$  in Cartesian coordinates with the origin at the mesh point C. We show that only one of those two ray directions may satisfy the causality condition, since the acute triangle must belong to one of the following cases

- 1. if  $\triangle ABC$  is completely located in the upper half plane, then a downgoing ray emanating from the segment AB should be selected to hit point C;
- 2. if  $\triangle ABC$  is completely located in the lower half plane, then a downgoing ray emanating from the segment AB should be selected to hit point C;
- 3. if  $\triangle ABC$  is located completely in the right half plane, then a left-going ray emanating from the segment AB should be selected to hit point C;

4. if  $\triangle ABC$  is located completely in the left half plane, then a right-going ray emanating from the segment AB should be selected to hit point C.

Finally to ensure that the characteristic indeed emanates from the segment AB, one has to verify that the selected outward normal allows a ray to start from point C and intersect with the segment AB and to accept the corresponding  $T_C$  accordingly; if not the rays hitting point C have to be enforced to travel along the edges AC and BC, which will be dealt with in the **third case**.

The second case: two identical outward normals. We can use the similar arguments as in the first case to decide whether to accept the corresponding characteristic or not; if not we go to the third case.

The third case: no valid outward normals. In this case we force rays to travel along either edge AC or edge BC. Since the ray direction is given and the Hamiltonian is convex we can find the corresponding group speed by inversion of the relation (2.4) (see [17] for more details). Then the traveltime at C is given by

$$T_{C} = \min\left\{T_{A} + \frac{|AC|}{v_{g}^{AC}}, T_{B} + \frac{|BC|}{v_{g}^{BC}}\right\}$$
(2.28)

where  $v_g^{AC}$  and  $v_g^{BC}$  denote the group speed along the edge AC and BC, respectively. **Remark:** Rectangular grids can be considered as special cases. There are two

**Remark:** Rectangular grids can be considered as special cases. There are two possible virtual triangulations on a rectangular grid as illustrated in Figure 2.4 for two spatial dimensions. In case (a) four triangles are created which is similar to a five-point stencil used in finite-difference schemes. In case (b) eight triangles are connected which result in a nine-point stencil. Case (b) will give more accurate solutions on the same Cartesian grid than Case (a) due to better directional resolution. As we will show in our numerical examples, the gain in accuracy of case (b) justifies the extra cost compared to case (a).



Figure 2.4: Triangulation based on regular meshes.

We can summarize the above into an algorithm. A 2-D local solver: (given  $T_A$  and  $T_B$ , determine  $T_C = T_C(T_A, T_B)$ .)

- 1. Solve equation (2.24) for two possible roots (more roots would be the same),  $T_C^1$  and  $T_C^2$ , either analytically or numerically.
- 2. If there are two roots,  $T_C^1$  and  $T_C^2$ , then
  - (a) if  $T_C^1$  satisfies the characteristic condition, then

$$T_C = \min\{T_C, T_C^1\}$$

(b) if  $T_C^2$  satisfies the characteristic condition, then

$$T_C = \min\{T_C, T_C^2\};$$

(c) if none of the two roots satisfies the characteristic condition, then

$$T_C = \min\left\{T_C, T_A + \frac{|AC|}{v_g^{AC}}, T_B + \frac{|BC|}{v_g^{BC}}\right\};$$

3. else

$$T_C = \min\left\{T_C, T_A + \frac{|AC|}{v_g^{AC}}, T_B + \frac{|BC|}{v_g^{BC}}\right\}$$

**Remark:** The same local solver applies to vertices on the computational boundary for outflow boundary conditions by using those triangles which are in the interior of the computational domain only.

## 2.4 Consistency and monotonicity

Considering a triangle  $\triangle ABC$  in which  $T_A$  and  $T_B$  are given, we update the travel-time  $T_C$  at the vertex C. Denoting

$$q_1 = \frac{T_C - T_A}{b}, \ q_2 = \frac{T_C - T_B}{a}, \ q_3 = \frac{T_B - T_A}{c}$$

we adopt the framework given in [4] to show consistency and monotonicity of the Godunov numerical Hamiltonian resulting from our local solver.

**Lemma 2.1 (Consistency and Monotonicity)** The numerical Hamiltonian  $\hat{H}$  is consistent:

$$\hat{H}\left(C, \frac{T_C - T_A}{b}, \frac{T_C - T_B}{a}\right) = H(C, \mathbf{p})$$
(2.29)

if  $\nabla T_h = \mathbf{p} \in \mathcal{R}^2$ . The numerical Hamiltonian  $\hat{H}$  constructed in the local solver is monotone if the causality condition holds.

**Proof.** By  $\nabla T_h = \mathbf{p} \in \mathcal{R}^2$ , we have

$$\begin{pmatrix} \frac{T_C - T_A}{b} \\ \frac{T_C - T_B}{a} \end{pmatrix} = \mathbf{P}\mathbf{p}.$$
(2.30)

Inserting this into the numerical Hamiltonian, we have equation (2.29).

Differentiating  $\hat{H}(C, q_1, q_2)$  with respect to  $q_1$  and  $q_2$ , the monotonicity of the Hamiltonian requires

$$\frac{\partial \hat{H}(C, q_1, q_2)}{\partial q_1} \ge 0, \qquad \frac{\partial \hat{H}(C, q_1, q_2)}{\partial q_2} \ge 0; \tag{2.31}$$

these can be satisfied if and only if the following holds component-wise:

$$\mathbf{P}^{-1}\nabla_{\mathbf{p}}H \ge \mathbf{0}.\tag{2.32}$$

The last inequality means that the characteristic direction  $\nabla_{\mathbf{p}} H$  at C is a convex combination of the two vectors CA and CB, which is exactly the condition that we have imposed in choosing the characteristic direction in the local solver, the upwinding condition.  $\Box$ 

## 2.5 How to compute the group speed?

In the above discussions we have to compute the group speed from a given ray direction. Since in general anisotropic media we cannot find an explicit formula for the group speed in terms of a given ray direction, we have to use a numerical procedure to determine the group speed approximately.

Fortunately for the static Hamilton-Jacobi equation with the Hamiltonian H being convex in the gradient argument we can easily modify a shooting method presented in [17] to compute the group speed from a given ray direction which is uniquely determined by two given points in a homogeneous medium. Since the algorithm is well explained in [17], we will not pursue it any further.

## 2.6 Acute versus obtuse triangles

In developing our local solver we have assumed that the triangulation does not consist of obtuse triangles. What happens if the triangulation does have obtuse triangles? We illustrate the consequences by using the paraxial eikonal theory and geometrical argument.

#### 2.6.1 Isotropic cases

Consider the isotropic eikonal equation. Assuming that we have an obtuse triangle  $\triangle ABC$  in which  $T_A$  and  $T_B$  are given, we update the travel-time  $T_C$  at the vertex C. Let the unit directional vector  $\mathbf{r}_1$  along edge CA and  $\mathbf{r}_2$  along edge CB be in the second and fourth quadrant, respectively; see Figure 2.5.

Then according to equation (2.26) we have vector  $\mathbf{r}$  located in between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , as illustrated in Figure 2.5, and the straight line defined by equation (2.25) has  $\mathbf{r}$  as its directional vector. Depending on  $\mathbf{r}_0$ , i.e. on  $T_A$ ,  $T_B$  and the triangle, the straight line may have no intersection, two identical intersections, and two different intersections with the slowness surface defined at the node C:

$$|\mathbf{p}|F(C) = 1; \tag{2.33}$$

see Figure 2.5.



Figure 2.5: An obtuse triangle and its consequences in isotropic wave propagation.



Figure 2.6: An obtuse triangle and its consequences in anisotropic wave propagation.

By the characteristic condition, if the triangle supports a consistent discretization then we will choose the intersection with the outward normal satisfying that the characteristic starting from C against the direction provided by the outward normal intersects the edge AB. As  $\mathbf{r}$  varies in between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , we have two extreme intersections defined by extending  $\mathbf{r}_1$  and  $\mathbf{r}_2$  to the isotropic slowness surface; they are  $D_1$  and  $D_4$  as illustrated in the figure. However, as observed from the figure the ray direction defined by the outward normal at  $D_1$  has negative first-component and positive second-component while the ray direction defined by the outward normal at  $D_4$ has positive first-component and negative second-component; similar observations can be made about  $D_2$  and  $D_3$ .

In fact we can rigorously prove that as  $\mathbf{r}$  varies from  $\mathbf{r}_1$  to  $\mathbf{r}_2$ , the ray direction defined by the corresponding outward normal changes its signs from (+, -) to (-, +), noting that the ray direction hitting point C is opposite to the outward normal direction in the figure. According to the proposed local solver, varying  $\mathbf{r}$  between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , the possible ray direction hitting point C will change its sign from (+, -) to (-, +) going through

$$(+,-) \rightarrow (0,-) \rightarrow (-,-) \rightarrow (-,0) \rightarrow (-,+)$$

or

$$(+,-) \rightarrow (+,0) \rightarrow (+,+) \rightarrow (0,+) \rightarrow (-,+)$$

due to the convexity of the slowness surface.

Therefore according to the paraxial eikonal theory [17] one cannot define a locally stable, uni-directional propagation problem to update the traveltime at node C by using traveltimes at node A and B.

#### 2.6.2 Anisotropic cases

Next we consider the anisotropic eikonal equation. Assuming that we have an obtuse triangle  $\triangle ABC$  in which  $T_A$  and  $T_B$  are given, we update the travel-time  $T_C$  at the vertex C.

Let the unit directional vectors  $\mathbf{r}_1$  along edge CA and  $\mathbf{r}_2$  along edge CB be in the second and fourth quadrant, respectively; the slowness surface defined at node C is given as illustrated in Figure 2.6. Then according to equation (2.26) we have vector  $\mathbf{r}$  located in between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , as illustrated in Figure 2.6, and the straight line defined by equation (2.25) has  $\mathbf{r}$  as its directional vector. Depending on  $\mathbf{r}_0$ , i.e. on  $T_A$ ,  $T_B$  and the triangle, the straight line may have no intersection, two identical intersections, and two different intersections with the convex slowness surface defined at the node C:

$$H(C, \mathbf{p}) = 1; \tag{2.34}$$

see Figure 2.6.

We note that this particular configuration as illustrated in Figure 2.6 has the following property: the sector of the slowness surface subtended by  $\mathbf{r}_1$ ,  $\mathbf{r}$  and  $\mathbf{r}_2$  has both horizontal and vertical tangent lines, and they are both unique due to the strict convexity.

By the characteristic condition, if the triangle supports a consistent discretization then we will choose the intersection with the outward normal satisfying that the characteristic starting from C against the direction provided by the outward normal intersects the edge AB. As **r** varies in between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , we have two extreme intersections defined by  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , respectively; they are  $D_1$  and  $D_4$  as illustrated in the figure. However, as observed from the figure the ray direction defined by the outward normal at  $D_1$  has negative first-component and positive second-component while the ray direction defined by the outward normal at  $D_4$  has positive first-component and negative second-component; similar observations can be made about  $D_2$  and  $D_3$ .

In fact we can rigorously prove that as  $\mathbf{r}$  varies from  $\mathbf{r}_1$  to  $\mathbf{r}_2$ , the ray direction defined by the corresponding outward normal changes its sign from (+, -) to (-, +). According to the proposed local solver, varying  $\mathbf{r}$  between  $\mathbf{r}_1$  and  $\mathbf{r}_2$ , the possible ray direction hitting point C will change its sign from (+, -) to (-, +) going through

$$(+,-) \rightarrow (0,-) \rightarrow (-,-) \rightarrow (-,0) \rightarrow (-,+)$$

or

$$(+,-) \rightarrow (+,0) \rightarrow (+,+) \rightarrow (0,+) \rightarrow (-,+)$$

due to the convexity of the slowness surface.

Therefore according to the paraxial eikonal theory [17] one cannot define a locally stable, uni-directional propagation problem to update the traveltime at node C by using traveltimes at node A and B.

# 2.7 Generalizations to higher dimensions

All the above procedure can be easily extended to higher dimensions. The design principle for the local solver still holds; namely, we first use consistency to find possible candidates and then check the causality condition. The only thing to which we need to pay attention is how to compute the group speed in higher dimensions if we are given two points. In this case although we have to solve an implicit nonlinear system, the problem still has a unique solution by the convexity of the slowness surface. Therefore, we can use a similar shooting method as the one in [17].

# 3 The Fast Sweeping Algorithm

We now describe the complete algorithm combining the local solver explained in the previous section with the fast sweeping strategy that we developed in [23].

- Step 1, Sorting: Sort all the nodes (vertexes) according to the  $l^p$  distance to a few reference points. In all our tests we use  $l^1$  distance.
- Step 2, Initialization: Assign large positive values to all nodes except those that belong to or near the boundary (the initial front). Those boundary nodes are assigned exact values or approximated values using shooting methods and these values are fixed in later iterations.
- Step 3, Fast sweeping: Start Gauss-Seidel iterations using the local solver described in the previous section with alternating sweeping orderings according to the increasing and decreasing of distances of nodes to the chosen reference points. During the iterations, it is crucial that the newly updated value is accepted only if it is smaller than the current value. This updating rule will guarantee that (1) the scheme is monotone and the value at any node is always improving; (2) once

the value is correct, i.e., reaches the possible minimum value, it will not change in later iterations.

Remark: If a rectangular grid is used then the sorting step is not needed as we can alternate the natural ordering by indexes in different directions.

Given the consistency and monotonicity of the local solver by Lemma 2.1, one can easily follow the proof in [32, 23] to show the following:

**Theorem 3.1** There exists a unique solution for the discretized nonlinear system and the fast sweeping iteration converges.

Also using similar arguments from [32, 23] one can show that the total number of sweepings needed depends only on the specific properties of the PDE, such as the turns of characteristics, which can be computed from the characteristic equations (2.4). This tells us how many sweepings are need to cover the tangent directions along a characteristic curve in the computational domain consecutively. On the discrete level, this indicates into how many connected regions we can divide all nodes such that in each region all nodes have the same dependence pattern on their neighbors; as a result this dependence pattern can be covered by one of the orderings in the upwind fashion.

Letting M be the total number of nodes, the complexity of the above algorithm is the following:

- on rectangular grids: O(M);
- on unstructured meshes.  $O(M \log M)$ .

One important note is that the constant in the complexity formula does not depend on the anisotropy of the Hamiltonian. As an example, for elliptical anisotropic eikonal equations, if the coefficients are constant, then the characteristics are straight lines; the number of iterations needed for the fast sweeping method to converge is independent of the anisotropy of the Hamiltonian.

**Remark:** On unstructured meshes the log M factor comes from the initial sorting of all nodes. Once the sorting is done the complexity of the fast sweeping iterations is O(M). Usually the sorting can be incorporated into the mesh generation easily with little extra cost.

# 4 Application: an elliptical anisotropic eikonal equation

We apply the above procedure and derive an explicit local solver for anisotropic eikonal equations of the following type

$$[\nabla T(\mathbf{x})M(\mathbf{x})\nabla T(\mathbf{x})]^{\frac{1}{2}} = 1, \quad \mathbf{x} \in \mathbb{R}^d,$$
(4.35)

where  $M(\mathbf{x})$  is a  $d \times d$  symmetric positive definite matrix. In particular M can be considered as a specific metric for the medium in which the wave front is propagating or in which we want to compute geodesics.

For simplicity let us consider in two dimensions,

$$H = \sqrt{a(\mathbf{x}) p_1^2 - 2c(\mathbf{x}) p_1 p_2 + b(\mathbf{x}) p_2^2} = 1, \qquad (4.36)$$

where a > 0, b > 0 and  $c^2 - ab < 0$ . Without abusing notations we use a, b and c to denote the coefficients in the anisotropic eikonal equation in the sequel.

Denote matrix  $\mathbf{P}$  in equation (2.20) by

$$\mathbf{P} = \left(\begin{array}{cc} n_{11}, & n_{12} \\ n_{21}, & n_{22} \end{array}\right),$$

where  $n_{11} = (x_C - x_A)/l_b$ ,  $n_{12} = (y_C - y_A)/l_b$ ,  $n_{21} = (x_C - x_B)/l_a$ ,  $n_{22} = (y_C - y_B)/l_a$ ;  $l_b$ ,  $l_a$  and  $l_c$  are the lengths of edge *CA*, *CB* and *AB*, respectively. Then

$$\mathbf{P^{-1}} = \frac{1}{\sin^2 \gamma} \begin{pmatrix} n_{11} - n_{21} \cos \gamma, & n_{21} - n_{11} \cos \gamma \\ n_{12} - n_{22} \cos \gamma, & n_{22} - n_{12} \cos \gamma \end{pmatrix} \equiv \begin{pmatrix} p_{11}, & p_{12} \\ p_{21}, & p_{22} \end{pmatrix}$$

From (2.21), we have

$$\nabla T(C) \approx \begin{pmatrix} \left(\frac{p_{11}}{l_b} + \frac{p_{12}}{l_a}\right)T_C - \left(\frac{p_{11}}{l_b}T_A + \frac{p_{12}}{l_a}T_B\right) \\ \left(\frac{p_{21}}{l_b} + \frac{p_{22}}{l_a}\right)T_C - \left(\frac{p_{21}}{l_b}T_A + \frac{p_{22}}{l_a}T_B\right) \end{pmatrix} \equiv \begin{pmatrix} g_1T_C + g_2 \\ g_3T_C + g_4 \end{pmatrix}, \quad (4.37)$$

where

$$g_1 \equiv \frac{p_{11}}{l_b} + \frac{p_{12}}{l_a}, \tag{4.38}$$

$$g_2 \equiv -(\frac{p_{11}}{l_b}T_A + \frac{p_{12}}{l_a}T_B), \qquad (4.39)$$

$$g_3 \equiv \left(\frac{p_{21}}{l_b} + \frac{p_{22}}{l_a}\right), \tag{4.40}$$

$$g_4 \equiv -(\frac{p_{21}}{l_b}T_A + \frac{p_{22}}{l_a}T_B). \tag{4.41}$$

Substituting  $\nabla T(C)$  in (4.37) into the Hamilton-Jacobi equation (4.36), we obtain the quadratic equation

$$w_1 T_C^2 + w_2 T_C + w_3 - 1 = 0, (4.42)$$

where

$$w_1 \equiv ag_1^2 + bg_3^2 - 2cg_1g_3, \tag{4.43}$$

$$w_2 \equiv 2ag_1g_2 + 2bg_3g_4 - 2c(g_1g_4 + g_2g_3), \qquad (4.44)$$

$$w_3 \equiv ag_2^2 + bg_4^2 - 2cg_2g_4, \tag{4.45}$$

where  $a = a(\mathbf{x}_{\mathbf{C}}), b = b(\mathbf{x}_{\mathbf{C}}), \text{ and } c = c(\mathbf{x}_{\mathbf{C}}).$ 

If the quadratic equation (4.42) has real roots

$$T_C = \frac{-w_2 \pm \sqrt{w_2^2 - 4w_1(w_3 - 1)}}{2w_1},\tag{4.46}$$

then we check the causality for the positive roots; i.e., if  $T_C$  is a positive root, we reconstruct  $\nabla T(C) = (p,q)$  by (4.37), and we calculate the characteristic direction

$$\mathbf{d} = \left(\begin{array}{c} ap - cq\\ bq - cp \end{array}\right);$$

next we check whether the characteristic line with direction **d** passing vertex C falls inside the triangle  $\triangle ABC$  or not. If the characteristic line is inside the  $\triangle ABC$ ,

causality is satisfied and we update  $T_C$  if this new value is smaller than the current numerical value of  $T_C$ . Otherwise, if the quadratic equation (4.42) has no positive root satisfying the causality condition, then we have to use the group speed along edges ACand BC, and

$$T_C = \min\left\{T_C, T_A + \frac{|AC|}{v_g^{AC}}, T_B + \frac{|BC|}{v_g^{BC}}\right\}.$$

**Remark:** If  $M(\mathbf{x})$  is a constant matrix, then the characteristics are straight lines. So four sweepings are enough to cover all directions of characteristics like the case of eikonal equations.

# 5 Numerical experiments

In all the examples we choose four corners in 2-D rectangular computational domains as the reference points and sort the nodes according to the  $l^1$ -metric by using the quicksort method, as in [23]. Actually ordering using  $l^1$  distance to two diagonal corners are the same. Essentially our ordering is the same as using two corners on any one side. The convergence of iteration is measured in terms of  $L^1$ -norm as advocated by Lin and Tadmor [15]; i.e., the iteration stops when the successive error satisfies  $||T^{n+1} - T^n||_{L^1} < 10^{-10}$ . In all of our test cases one sweeping means one Gauss-Seidel iteration with a particular ordering through all nodes.

A typical acute triangulation is shown in Fig. 5.1. When we check the accuracy of our methods, we refine the mesh uniformly, i.e., cutting each triangle into four smaller similar ones. Exact solutions needed in initializing the algorithm and checking accuracy are computed by the shooting method [17] whenever possible. In our examples, no special treatment is needed for vertices on the boundary of the computational domain. Their computational stencils (Fig. 2.1) only involves triangles which are in the interior of the computational domain.

## 5.1 Example 1

We consider the following equation

$$\sqrt{a T_x^2 + b T_y^2 - 2c T_x T_y} = 1, \quad (x, y) \in (-2, 2) \times (-2, 2), \tag{5.1}$$

$$T(0,0) = 0, (5.2)$$

where a > 0, b > 0 and  $c^2 - ab < 0$ .

In a homogeneous anisotropic metric the eigenvalues of the symmetric positive definite matrix

$$\mathbf{M} = \left(\begin{array}{cc} a, & c \\ c, & b \end{array}\right)$$

characterizes the anisotropy of the metric. According to [26] the anisotropy coefficient of the metric is defined by

$$\eta = \sqrt{rac{\lambda_{\max}(\mathbf{M})}{\lambda_{\min}(\mathbf{M})}},$$

where  $\lambda_{\max}(\mathbf{M})$  and  $\lambda_{\min}(\mathbf{M})$  are the larger and smaller eigenvalues of  $\mathbf{M}$ , respectively.

**Case 1: a homogeneous, mild anisotropic case.** We take a = 1, b = 1, and c = 0.9;  $\eta = \sqrt{19}$ . Since the point source problem has an upwind singularity at the source [18] we have to measure the order of accuracy of the fast sweeping method away from the singularity. Otherwise the accuracy will degenerate to  $h \log h$  [32]. To achieve this we first fix a small region,  $[-0.2, 0.2] \times [-0.2, 0.2]$ , around the source and assign the exact solution to the grid points inside the small region, then the numerical error is computed only for the grid points outside the small region; this is the so-called wrapping technique. As shown in Table 5.1, with this wrapping technique we are able to observe the expected first-order accuracy while without the wrapping technique we are only able to observe the degraded first-order accuracy. In Table 5.1, we also observe that the number of iterations needed for convergence is almost a constant which is independent of mesh sizes. Figure 5.2 shows the contour plot for this test case.

Next we test our algorithm on rectangular grids, which can be considered as special cases. See the discussion in Section 2 and Figure 2.4. The sweeping directions are based on the  $l^1$ -metric. The accuracy and sweeping iteration numbers are listed in Table 5.2 for both the four three-point stencils and the eight three-point stencils (Figure 2.4). Apparently the eight three-point stencils can result in a much better accuracy than the four three-point stencils.

In this case of rectangular grids we also test the i, j orderings as that used in the fast sweeping method on rectangular meshes, and the same accuracy and the same number of sweepings are obtained; we omit the results here.

**Case 2: Homogeneous, strong anisotropic cases.** We carry out a sequence of tests to study the power and the robustness of our sweeping methods.

First we increase the coefficient a successively from 2 to 2000 with b = 1 and c = 0 fixed. Figure 5.3 shows the contour plots of the results for a = 2, 20, 200, 2000 computed by using the same mesh with 5716 nodes and 11264 triangles. In addition for a = 200, 2000 we use a finer mesh with 90625 nodes and 180224 triangles; the results are shown in Figure 5.4, which have higher resolution than the ones shown in Figure 5.3.

The above cases have the symmetrical axis of the slowness surface aligned with the Cartesian axis, which may not be able to test out the full power of the sweeping methods. To do that we take out the two cases, (a = 200, b = 1, c = 0) and (a = 2000, b = 1, c = 0), and apply to the resulting matrices M a similarity transform defined by a rotation with angle  $\frac{\pi}{6}$ . Then we have (a = 150.25, b = 50.75, c = 86.16953) and (a = 1500.25, b = 500.75, c = 865.5924); the resulting anisotropic coefficients are  $\eta = \sqrt{200}$  and  $\eta = \sqrt{2000}$ , respectively.

Figure 5.5 shows the contour plots for the two cases with the computational mesh of 90625 nodes and 180224 triangles. Numerical errors and order of convergence are shown in Table 5.3 and Table 5.4. We have used  $L^1$ -norm to measure convergence order which is advocated by Lin and Tadmor [15].

These results demonstrate that our sweeping method is robust enough to handle the equation with a very high anisotropy coefficient. Unlike other methods with numerical domain of dependency depending on the anisotropy coefficient  $\eta$  which may be very large when  $\eta$  is large, our iterative methods do not have such a shortcoming and thus are efficient and robust. As discussed in Section 3 the number of sweepings is independent of mesh size and anisotropy and is dependent on the behavior of characteristics. For the case of constant coefficients, the characteristics are straight-lines and that is why

		Wrapping	g [-0.2,	No w	rapping	r D	
Nodes	Elements	$L^1$ error	order	iter	$L^1$ error	order	iter
1473	2816	6.45E-2	_	4	7.47E-2	_	4
5716	11264	3.27E-2	0.98	4	4.68E-2	0.68	4
22785	45056	1.75E-2	0.91	4	2.87E-2	0.71	5
90625	180224	8.88E-3	0.97	4	1.71E-2	0.75	5

Table 5.1: The order of convergence; a = 1, b = 1, c = 0.9.

Table 5.2: Comparison of the accuracy between the four three-point stencils and the eight three-point stencils (Fig. 2.4); Wrapping  $[-0.2, 0.2] \times [-0.2, 0.2]$ ; a = 1, b = 1, c = 0.9.

	Four three	e-point	stencils	Eight three-point stencils		
Mesh	$L^1$ error	order	iter	$L^1$ error	order	iter
$40 \times 40$	1.17E-1	_	4	1.57E-2	_	4
$80 \times 80$	6.35E-2	0.88	4	8.18E-3	0.94	4
$160 \times 160$	3.39E-2	0.90	4	4.18E-3	0.97	4
$320 \times 320$	1.78E-2	0.93	4	2.12E-3	0.98	4

we only need 4-5 sweepings.

## 5.2 Example 2

We consider the following equation with variable coefficients

$$\sqrt{a(x,y) T_x^2 + b(x,y) T_y^2 - 2c(x,y) T_x T_y} = 1, \quad (x,y) \in (-1,1) \times (-1,1),$$
  
$$T(x,y) = 0, (x,y) \in \Gamma,$$

where  $\Gamma$  is a unit square in the middle of the domain:  $\Gamma = \{x = \pm 0.5, |y| \le 0.5\} \cup \{y = \pm 0.5, |x| \le 0.5\}$ .

We choose  $a(x,y) = 150.25(1 + \lambda \sin^2(\pi xy)), \ b(x,y) = 50.75(1 + \delta \cos^2(\pi xy)), \ c(x,y) = 86.16953(1 - \epsilon \sin^2(\pi xy)), \text{ where } \lambda, \delta \text{ and } \epsilon \text{ are constants to be selected};$ 

Table $5.3$ :	The ord	ler of o	convergence;	a = 150.25,	b = 50.75, c =	86.16953.
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		Wrapping	g [-0.2,	No w	vrapping	r D	
Nodes	Elements	$L^1$ error	order	iter	$L^1$ error	order	iter
1473	2816	8.78E-3	_	4	8.87E-3	-	4
5716	11264	4.04E-3	1.12	4	5.38E-3	0.72	4
22785	45056	2.10E-3	0.94	4	3.22E-3	0.74	5
90625	180224	1.04E-3	1.02	4	1.88E-3	0.77	5

		Wrapping	g [-0.2,	No w	rapping	r S	
Nodes	Elements	$L^1$ error	order	iter	$L^1$ error	order	iter
1473	2816	6.23E-3		4	5.72E-3		4
5716	11264	2.89E-3	1.11	4	3.33E-3	0.78	4
22785	45056	1.53E-3	0.92	4	1.93E-3	0.79	5
90625	180224	7.66E-4	1.00	4	1.10E-3	0.80	5

Table 5.4: The order of convergence; a = 1500.25, b = 500.75, c = 865.5924.



Figure 5.1: A typical acute triangulation.



Figure 5.2:  $a = 1, b = 1, c = 0.9, \eta = \sqrt{19}$ ; convergence after 4 sweepings.

Table 5.5: Comparison of iteration numbers of the four three-point stencil (Fig. 2.4 (a)), the eight three-point stencil (Fig. 2.4 (b)) and the regular triangular stencil (Fig. 2.1),  $a(x, y) = 150.25(1 + \lambda \sin^2(\pi xy)), b(x, y) = 50.75(1 + \delta \cos^2(\pi xy)), c(x, y) = 86.16953(1 - \epsilon \sin^2(\pi xy))$  where  $\lambda = 1, \delta = 1, \epsilon = 0.125$ .

Four three-	-point stencil	Eight three	e-point stencil	triangular stencil	
Mesh	iter	Mesh	iter	Mesh Nodes	iter
$40 \times 40$	11	$40 \times 40$	9	1473	9
$80 \times 80$	13	$80 \times 80$	10	5761	7
$160 \times 160$	13	$160 \times 160$	11	22785	8
$320 \times 320$	13	$320 \times 320$	13	90625	9

this is a perturbation of the homogeneous case (a, b, c) = (150.25, 50.75, 86.16953).

We solve this equation on three different sets of stencils: a regular triangular mesh and two virtual meshes constructed from uniform rectangular grids, where one virtual mesh consists of four three-point stencils at each node and the other of eight three-point stencils at each node.

We take  $\lambda = 1$ ,  $\delta = 1$  and  $\epsilon = 0.125$ . The results in terms of mesh refinement are shown in Table 5.5. Figure 5.6 shows contours of the solutions on different meshes.

# 6 Conclusion

We develop a fast sweeping method for static Hamilton-Jacobi equations with convex Hamiltonians. Local solvers and fast sweeping strategies apply to structured and unstructured meshes. With causality correctly enforced during sweepings numerical evidence indicates that the fast sweeping method converges in a finite number of iterations independent of mesh size. Numerical examples validate both the accuracy and the efficiency of the new methods.

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Figure 5.3: (a):  $a = 2, b = 1, c = 0, \eta = \sqrt{2}$ ; convergence after 5 sweepings; (b):  $a = 20, b = 1, c = 0, \eta = \sqrt{20}$ ; convergence after 5 sweepings; (c):  $a = 200, b = 1, c = 0, \eta = \sqrt{200}$ ; convergence after 5 sweepings; (d):  $a = 2000, b = 1, c = 0, \eta = \sqrt{2000}$ ; convergence after 5 sweepings.



Figure 5.4: Refined mesh. (a):  $a = 200, b = 1, c = 0, \eta = \sqrt{200}$ ; convergence after 5 sweepings; (b):  $a = 2000, b = 1, c = 0, \eta = \sqrt{2000}$ ; convergence after 5 sweepings.



Figure 5.5: (a):  $a = 150.25, b = 50.75, c = 86.16953, \eta = \sqrt{200}$ ; convergence after 5 sweepings; (b):  $a = 1500.25, b = 500.75, c = 865.5924, \eta = \sqrt{2000}$ ; convergence after 5 sweepings.



Figure 5.6:  $a(x,y) = 150.25(1 + \lambda \sin^2(\pi xy)), b(x,y) = 50.75(1 + \delta \cos^2(\pi xy)), c(x,y) = 86.16953(1 - \epsilon \sin^2(\pi xy))$  where  $\lambda = 1, \delta = 1, \epsilon = 0.125$ . (a): On a general triangular mesh with 22785 nodes, convergence after 8 sweepings; (b): on the 160 × 160 rectangular mesh, using the eight three-point stencils, convergence after 11 sweepings; (c) on the 160 × 160 rectangular mesh, using the four three-point stencils, convergence after 13 sweepings.