# Rapid and Accurate Computation of the Distance Function Using Grids

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E-mail: ytsai@math.ucla.edu Abstract

We present two fast and simple algorithms for approximating the distance function for given isolated points on uniform grids. The algorithms are then generalized to compute the distance to piecewise linear objects. By incorporating the geometry of Huygen's principle in the reverse order with the classical viscosity solution theory for the eikonal equation  $|\nabla u| = 1$ , the algorithms become almost purely algebraic and yield very accurate approximations. The generalized closest point formulation used in the second algorithm provides a framework for further extension to compute the distance accurately to smooth geometric objects on different grid geometries, without the construction of the Voronoi diagrams. This method provides a fast and simple translator of data commonly given in computational geometry to the volumetric representation used in level set methods.

### 1. INTRODUCTION

Given a compact set K in space, the distance function for K at a point x is defined to be the smallest Euclidean distance of x to the points in K. Fast and accurate computation of the distance function is an important step in many applications. For example, level set methods [14] rely heavily on it to represent an interface, and, in particular to keep the level set function well behaved during time evolution. In the surface interpolation model of [28], a surface interpolating a given points set K is constructed by first computing the distance function to K. In the Island Dynamics model of [6] in materials science, one needs the distance function to help understand the aggregation of atoms into islands. Generally, applications in physics, chemistry, molecular biology, and even urban planning that use point pattern analysis require construction of the distance function or (at least) Voronoi diagrams [12] [5]. Furthermore, in applications such as ray tracing and surface rendering where one needs information about the normals or other geometric quantities of the surfaces of the given objects, the distance function is essential.

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In computational geometry, the extraction of distance information is dealt with through the construction of the Voronoi diagram. There is an abundance of literature about this topic. For isolated points, those algorithms usually start out using geometric properties (usually Euclidean distance between points and some ad hoc observations) and intricate sorting algorithms to construct a representation (e.g. the "winged-edge" data structures [12]) of the Voronoi diagram that is optimally linear in the number of data points [24][1]. For configurations of slightly more general geometric objects such as isolated points and linear segments (triangles in the planes), there are  $\mathcal{O}(M \log M)$  algorithms for its construction and point location [27], where M is the number of segments. However, if we need to evaluate the distance at all the grid points, we still have to go through the data structures on each grid point to compute the information. The overall complexity is at least  $\mathcal{O}(N)$  where N is the number of grid points. We would like to emphasize that a lot of applications can be solved more efficiently, both in terms of development time and computation time, if we accept the use of grids. One reason is that these applications are formulated and solved on the grids and require only the distance values on the grid nodes. In [22], the author proposed a fast tree-based method that efficiently uses a fast construction of Voronoi diagrams for accurate distance computation. Another efficient approach that uses Voronoi diagram for distance computation on the grids is reported in [11], in which the author considered mainly a single triangulated object.

Thus we want to provide a framework that can be generalized to compute the distances to a variety of interfaces often encountered in the PDE-level set formulations of the problems. From the perspective of the grid-based numerical methods for PDEs, our method can thus be adopted almost effortlessly. Of course many PDE based applications are discretized on grids, e.g. the surface interpolation of [28]. Even problems involving the solution of PDEs on surfaces can use the level set formulation [3][10]. In these cases, our algorithms provide fast and simple procedures to accomplish the task of generating the distance function. With our algorithms, the user only has to visit each grid point and perform some simple calculations. The complexity is linear in the numer of grid points in the grid that resolves the interface. Only the data structures very similar or identical to the original ones used in the original applications are required. One can also think of our approach as a fast volumetric preprocessing of the distance.

We also note that once we have a second or higher order approximate distance function  $d(\bar{x})$ , we can easily find the Voronoi generator  $\bar{x}^*$  of any given point  $\bar{x}$  by  $\bar{x}^* = \bar{x} - d(\bar{x})\nabla d(\bar{x})$ , which requires  $\mathcal{O}(1)$  operations.

The rest of this paper is organized as follows: We begin by presenting the standard PDE approximation methods for finding the distance function with an iterative scheme [23] and the Dijkstra like fast marching method [26]. Next we give motivation for the new method based on the geometric consideration of the reverse Huygen's principle. We employ a special sweeping strategy similar to the ones used in [4][8][25] to update each grid point. We then reformulate the computation of the distance function by storing the closest points on the interface on each grid point. Finally, we generalize the closest point algorithm to compute the distance accurately to triangulated surfaces commonly used in computational geometry to the embedded surface formulation used by the level set methods.

For simplicity, we will first discuss the problem and setup in  $\mathbb{R}^2$ . This can be directly extended to  $\mathbb{R}^3$ .

### 2. THE PROBLEM

Let  $\Omega$  be a compact connected subset of  $\mathbb{R}^2$  and  $\Gamma \subset \Omega$  closed. The distance function  $u(\bar{x})$  to  $\Gamma$  is defined by

$$u(\bar{x}) = {\rm dist} \ (\bar{x}, \Gamma) := \min_{p \in \Gamma} |\bar{x} - p|$$

as a deterministic optimization problem. However, it can also be formulated as the viscosity solution of the Eikonal equation

$$|\nabla u| - 1 = 0 \text{ with } u(\bar{x}) = 0 \text{ for } \bar{x} \in \Gamma;$$
(1)

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i.e.  $u: \Omega \to \mathbb{R}$  is continuous,

$$u(\bar{x}) = \lim_{\epsilon \to 0} u^{\epsilon}(\bar{x})$$

where  $u^{\epsilon}$  solves the parabolic equations

$$|\nabla u^{\epsilon}| - \epsilon \Delta u^{\epsilon} = 1 \ (\epsilon > 0).$$

The existence and uniqueness of such viscosity solution  $u(\bar{x})$  is well known, see e.g [2]. Additionally, from classical PDE theory, it follows that u is increasing along the characteristics in regions where the characteristics do not cross. This is crucial to the approximation procedures described later in this paper.

Before describing our algorithms, we present some notation which shall be used later.

DEFINITION 2.1. Given  $P \in \Omega$  and  $\Gamma \subset \Omega$  compact, we define

source
$$(P) = \{Q \in \Gamma : \operatorname{dist}(P, \Gamma) = |P - Q|\}.$$

Clearly, if P is in the interior of a Voronoi region, source(P) contains only a unique point, and we shall call this point  $P^*$ . If P is on the Voronoi boundaries, we shall choose one element in source(P), and also call this predetermined point  $P^*$  for convenience. Since we are concerned with the distance, any such  $P^*$  works. We will also adopt the conventional term used in computational geometry with  $P^*$  being called *a Voronoi generator* [12].

We consider  $\Omega \subset \mathbb{R}^2$  a rectangular region and  $\Omega_d$  its discretization with uniform grid size h in both the x- and y- directions.  $u_{i,j}$  refers to u(ih, jh).

DEFINITION 2.2. (neighbors) Let  $E = (x_i, y_j) \in \Omega_d$ . We say P is a neighbor of E if  $P \in \{(x_i, y_{j\pm 1}), (x_{i\pm 1}, y_j)\}$ .

DEFINITION 2.3.  $(\sqrt{2}h \text{ neighbors})$  We say Q is a  $\sqrt{2}h$  neighbor of E if  $Q \in \{(x_{i\pm 1}, y_{j\pm 1})\}$ .

# 3. APPROXIMATION OF THE DISCRETIZED EQUATION — GODUNOV'S HAMILTONIAN

To put our proposed methods into context, we review briefly the distance computation from the numerical PDE point of view. This provides the motivation and the foundation of our algorithms.

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Let u(x) be the viscosity solution of (1). Rouy and Tourin[18] proved convergence to the viscosity solution of an iterative method solving the steady state equation (1) with the Godunov Hamiltonian approximating  $|\nabla u|$ :

$$H_G(p_-, p_+, q_-, q_+) = \sqrt{\max\{p_-^+, p_+^-\}^2 + \max\{q_-^+, q_+^-\}^2}$$
(2)

where  $p_{\pm} = D_{\pm}^{x} u_{i,j}$ ,  $q_{\pm} = D_{\pm}^{y} u_{i,j}$ ,  $D_{\pm}^{x} u_{i,j} = \pm (u_{i\pm 1,j} - u_{i,j})/h$  and accordingly for  $D_{\pm}^{y} u_{i,j}$ . Here  $u_{i,j}$  is the approximation to the exact solution, and  $x^{+} = \max(x, 0)$ ,  $x^{-} = -\min(x, 0)$ . The Godunov Hamiltonian yields a convergent method that is first order accurate in grid size.

Osher[13] proved that the *t*-level set of u(x) is the zero level set of the viscosity solution of the evolution equation at time *t* 

$$\phi_t = |\nabla\phi| \tag{3}$$

with appropriate initial conditions. So one can try to solve the time-dependent equation by the level set formulation [16] with a consistent, monotone Hamiltonian [17]. Crandall and Lions proved that the discrete solution obtained this way converges to the desired viscosity solution [7].

Later, Tsitsiklis [26] combined the heap sorting procedure with a variant of Dijkstra's algorithm to solve the steady state equation of the more general problem

$$|\nabla \phi| - R(\bar{x}) = 0 \text{ for } R(\bar{x}) > 0.$$

$$\tag{4}$$

This was later rederived in [20] and in [9] and has become known as the fast marching method. Its complexity is  $\mathcal{O}(N \log(N))$ , where N is the number of grid points.

Later, Osher and Helmsen gave a criterion on general Hamiltonians so that the fast marching algorithm is applicable, and extended the fast marching methods to these more general problems [15].

In [4], Boué and Dupuis suggest a "sweeping" approach to solve the steady state equation, which, in our experience, results in a  $\mathcal{O}(N)$  algorithm for the simple problem at hand. We point out here that this strategy when applied to compute the distance to a set of isolated points is very similar to that described in an orginal and quite early paper of Danielsson [8]. However, the author in [8] considered only points lying on the grid and did not provide extension to the three dimensional case. This iterative approach has been used successfully in [25] in both 2D and 3D. We shall incorporate it into our algorithms. Using this "sweeping" approach, the complexity of our algorithms drops to  $\mathcal{O}(N)$  from  $\mathcal{O}(N \log N)$  in the fast marching case and the implementation of the algorithms becomes a bit easier than the fast marching method that requires heap sort. (Sweeping is not recommended for general  $R(\bar{x})$ in equation [4])

Since the fast marching method is, by now, well known, we will not give much detail on its implementation in this paper. In general, this involves a sorting procedure and the solution of

$$H_G(D_-^x u_{i,j}, D_+^x u_{i,j}, D_-^y u_{i,j}, D_+^y u_{i,j}) = 1$$
(5)

for  $u_{ij}$  in terms of its four neighboring values.





**FIG. 2.** The isosurface  $\{u = 0.1\}$  computed by the solution of the Godunov Hamiltonian



It is important to note that this method yields the exact distance of a point inside the Voronoi regions induced by lines in 2D and planes in 3D. The normal  $n(x,y) = \nabla u(x,y)/|\nabla u|$  of the distance to the lines is constant. So any consistent divided difference of u gives an exact evaluation of its derivatives.

However, this method is inconsistent for computing the distance to isolated points. For instance, if  $u_{l,m} = 0$ , then the computed  $u_{l+1,m} = u_{l,m+1} = h$ , and  $u_{l+1,m+1} = (1 + 1/\sqrt{2})h$ , which is the distance between  $u_{l+1,m+1}$  to the line joining  $u_{l+1,m}$  and  $u_{l,m+1}$  plus h. That is, the Godunov solver treats the values  $u_{l+1,m}$  and  $u_{l,m+1}$  as though they come from a plane wave, resulting in an early arrival time (figure 1). The level sets of the solution look like diamond shaped polyhedra, as seen in figure 2, instead of perfect circles/spheres. This leads us to take more specific consideration of the geometry of the problem.

#### 4. APPROXIMATION BY GEOMETRIC CONSIDERATIONS

Since in our algorithms the two dimensional problem is only a special case of the three dimensional case, we will now discuss our algorithms in the three dimensional setting. To facilitate our discussion, we first introduce some more notation.

NOTATION 4.1. Let  $P \in \Omega \subset \mathbb{R}^2$ , and u(P) be the value of the viscosity solution of (1). We use the following notation for brevity

$$C_P := \{ \bar{x} \in \Omega : |\bar{x} - P| = u(P) \},\$$

which is the circle with radius u(P) centered at P. In 3D, we will use  $S_P$  to denote the sphere centered at P with radius u(P).

NOTATION 4.2. Let  $E \in \Omega$  be represented by  $(i, j, k) \in \Omega_d \in \mathbb{R}^3$ . Depending on the situation, we will use u(E) and u(i, j, k) interchangeably to represent the value of u at E.

DEFINITION 4.1. (Sweeping directions in 3D) We define the following iterations going through the uniform grids (1 : Nx - 1, 1 : Ny - 1, 1 : Nz - 1).

```
(x+,y+,z+) sweeping:
for k=1:Nz-1
for j=0:Ny-1
for i=0:Nx-1
update u_{i,j,k}
(z-,y-,x-) sweeping:
for i=Nx-1:1
for j=Ny-1:1
for k=Nz-1:1
update u_{i,j,k}
```

Following the above examples, we can define the rest of the 6 iterations: (y+,x+,z-), (y-,x-,z+), (x-,y+,z+), (z-,y-,x+), (x+,y-,z+), and (z-,y+,x-).

#### 4.1. Motivation and algorithm

We return briefly to two dimensions for simplicity for this part of the exposition. Let us first consider a single Voronoi generator  $E^*$  whose location is not explicitly known at the start of the algorithm. Suppose  $r_P$  and  $r_Q$  are the distances of two distinct points P and Q to  $\Gamma$ , respectively: i.e.  $u(P) = r_P$ ,  $u(Q) = r_Q$ . Let E = (x, y) be a point of interest with

$$||E - E^*|| \ge \max\{u(P), u(Q)\}$$
(6)

a priori. Consider  $C_P$  and  $C_Q$  as in the definition:  $C_P$  is the circle of radius  $r_P$  centered at P and  $C_Q$  of radius  $r_Q$  centered at Q. The intersections of  $C_P$  and  $C_Q$  are the solutions of a quadratic equation. Denote the two intersections by V and W. It is then easy to see that either V or W is  $E^*$ . From the hypothesis that E is farther from  $E^*$  than P or Q is, we deduce that  $u(E) = \max\{||E - W||, ||E - V||\}$ .

Thus, by enforcing this sort of upwinding decision in selecting intersections to maintain monotonicity of the solution "from"  $\Gamma$ , we are able to approximate the distance function. In fact, this is the key property for the success of the aforementioned fast marching and sweeping methods.

Now we can generalize to a full 3D algorithm for distance approximation of a given set of isolated points.

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Algorithm 1 (Spherical intersection solver).

1. Initialize: give the exact distance to u at grid points that are

 $\sqrt{2}$ -neighbors of  $\Gamma \subset \Omega_d$ . Mark them so they will not be updated. Mark all other grid values as  $\infty$ .

2. Iterate through each grid point E with index (i,j,k) in each sweeping direction or according to the fast marching heap sort;

• Select stencils: At each grid point (i,j,k), find its neighbors  $P_1, P_2$ , and  $P_3$  in x-, y-, and z-direction respectively such that  $u(P_1) \le u(i \pm 1, j, k), u(P_2) \le u(i, j \pm 1, k)$ , and  $u(P_3) \le u(i, j, k \pm 1)$ . Let  $NBD = \{P_1, P_2, P_3\}$ .

– Discard from NBD the element(s) on which u is  $\infty$ ,

- If  $NBD = \emptyset$ . Goto 3.

- Otherwise, let  $Q \in NBD$  such that  $u(Q) = \min\{u(P) : P \in NBD\}$ . Discard from NBD those  $P_l$  that are too large:  $u(P_l) - u(Q) > ||P_l - Q||$ .

• If all  $P_1, P_2$ , and  $P_3$  remain in NBD :

- If  $S_{P_1}, S_{P_2}$ , and  $S_{P_3}$  intersect: Let V and W be the two points of their intersection, and set  $u(E) = \max \{ ||E - V||, ||E - W|| \}.$ 

\* If the monotone requirement  $u(E) \ge u(P_i)$ , i = 1, 2, 3 is satisfied, skip to step 3.

\* Otherwise the intersections contain false information. Treat as if  $S_{P_1}$ ,  $S_{P_2}$ , and  $S_{P_3}$  do not intersect.

- If  $S_{P_1}$ ,  $S_{P_2}$ , and  $S_{P_3}$  do not intersect: Discard the element from NBD on which u has the greatest value, i.e. discard from NBD the P with  $u(P) = \max_{p_l \in NBD} u(P_l)$ . Go to the next case.

• If there are only two  $P'_l s$  remaining in NBD (assuming they are  $P_2$ ,  $P_3$  without loss of generality):

– If the circles  $C_{P_2}$  and  $C_{P_3}$  lying on the y-z plane intersect, let V and W be the intersections. Set  $u(E) = \max \{||E - V||, ||E - W||\}$ . Check if the monotonicity requirement  $u(E) \ge u(P_2), u(P_3)$  is satisfied. If yes, go to step 3, otherwise, treat as if  $C_{P_2}$  and  $C_{P_3}$  do

not intersect.

- If  $C_{P_2}$  and  $C_{P_3}$  do not intersect: discard from NBD the P with  $u(P) = \max\{u(P_1), u(P_2)\}$ .

• If there is only one  $P_l$  remaining, simply set  $u(E) = u(P_l) + h$ .

3. Go to step 2.

Note that in the updating procedure, we never update the boundary of  $\Omega_d$  which is initially set to  $\infty$ . This is due to the fact that the characteristics only flow towards the boundaries, therefore the upwinding stencils never contain the grid points on the boundaries.

#### 4.2. Relation to "reversing" Huygen's principle and the computation of envelopes

In the above algorithms, we determine the location of one Voronoi generator in  $\Gamma$  by finding the intersections of two circles in 2D and three spheres in 3D. In the 2D case, we





are asking basically "where is the point whose distances to the stencils  $P_1$  and  $P_2$  are as prescribed by  $u(P_1)$  and  $u(P_2)$ ?" If we consider the points in a neighborhood of E in the continuous case, we know that  $E^*$  has to be the intersection of all the corresponding circles centered at the neighboring points. More precisely, one can consider  $\Gamma$  as the envelope of the circles centered at each point in space with radius prescribed by the distance function evaluated at that point. This can be viewed as sending circular waves from each point in the neighborhood of E at times prescribed by the distance function, and asking for the envelope of the waves at a certain time. It is as if we are running Huygen's principle "backwards." This concept can certainly be extended to general  $\Gamma$  such as curves and surfaces.

#### 4.3. A hybrid algorithm

As noted above, solving the Godunov Hamiltonian will give the exact distance to hyperplanes in their Voronoi regions. On the other hand, our methods give the exact distance for some simple point configurations and at least first order approximation in certain circumstances (please see the section about the error analysis). We would like to create a method that makes use of the advantages of both approaches.

The idea is to adopt a consistency check of the solution obtained from the spherical method. At grid points where the result of the check implies an inconsistency of the distance information, we return to solving Godunov's Hamiltonian (2). Here, the inconsistency of the distance information means either an ambiguity (defined below) or the *disagreement* in the intersection computed from using another set of neighboring grid points as stencil. Figure 3 delineates what *disagreement* means: let S be the  $\sqrt{2}$ -neighbor with  $u(S) = \min\{u(T) : T \text{ is a } \sqrt{2} \text{ neighbor of } E\}$ , and P and Q be the stencils selected from the regular neighbors of E by the spherical solver. If the three circles  $C_S$ ,  $C_P$ , and  $C_Q$  do not intersect, we switch to the Godunov method.

DEFINITION 4.2. We say we have ambiguity when any of the following cases happen:

C<sub>1</sub> and C<sub>2</sub> do not intersect.
 u(E) = max{|E - W|, |E - V|} < min{u(P), u(Q)}</li>
 u(E) = u(P) = u(Q).

Algorithm 2 (A 2D hybrid solver).

1. Initialize: Set  $u_{ij} \equiv \infty$  for  $(i, j) \notin \Gamma$ . For each  $\gamma \in \Gamma$ , set  $u(\gamma) = 0$ . For  $z \in \Gamma$  not in  $\Omega_d$ , compute the exact solution at the vertices of the grid cell in which z. lies.

2. For each sweep direction in  $\{(x+,y+), (y-,x-), (x+,y-), (x-,y+)\}$ , iterate through each grid point according each of the sweep directions or according to the fast marching heap sort.

3. At each grid E with index (i,j)

- if all the neighbors are infinity, skip
- if there is at least one non-infinity neighbor in both the

x- and y- direction,

let P and Q be the smallest one in each direction.

Without loss of generality, assume u(P) > u(Q). Discard P if

u(P) - u(Q) > ||P - Q||.

• if there is only one finite neighbor  $\tilde{z}$ , set  $u(E) = u(\tilde{z}) + h$ .

• if both P and Q remain, find the points of intersection of circles  $C_P$  and  $C_Q$ .

- If ambiguous, solve equation (5) for u(E) instead.

- If not ambiguous, find the smallest neighbor S from the  $\sqrt{2}h$  neighbors  $\{i \pm 1, j \pm 1\}$ , and find the points of intersection of the circle  $C_S$  and  $C_P$  or  $C_Q$ . If there are no common points of intersection, solve equation (5) for u(E) instead.

– Otherwise, set  $u(E) = \max\{||E - W||, ||E - V||\}$ , where W and V are the intersects.

We emphasize here that if the normals of the line segments in  $\Gamma$  do not have slopes which are integer multiples of  $\pi/4$ , we need to adopt the  $\sqrt{2}$ -neighbors of a given grid point as the regular stencils. Otherwise, the sweeping method would simply fail because during each iteration, we cannot compute and propagate the exact distance away from  $\Gamma$ . However, this problem may be resolved if we adopt a "generalization" of the closest point formulation (which is a current research direction, and shall be mentioned at the end of this paper).

#### 4.4. The closest point formulation

We see that the above algorithm tries to determine the closest point  $E^*$  of each grid point E. So if we store the generator of each grid point and propagate this closest point information somewhat "along" the characteristics, the computation of the intersections of spheres at each grid point becomes unnecessary. All we have to do is to maintain the monotonicity of the solution. This formulation is motivated by the work of Sean Mauch[11] and the Dynamic Surface Extension Method of Steinhoff et al.[21][19].

DEFINITION 4.3. Given  $\Gamma$ , a closest point function for  $\Gamma$  is  $V : \mathbb{R}^3 \to \mathbb{R}^3$  such that

$$V(E) = E^{\dagger}$$

for  $E^*$  predetermined as described in definition 1.

We thus arrive at the following algorithm:

Let u be the distance function on the grids, and V be the corresponding closest point function.

Algorithm 3 (The closest point solver).

1. Initialize: give the exact distance to u, and the exact closest points to V at grid points near  $\Gamma$ . Mark them so they will not be updated. Mark all other grid values as  $\infty$ .

2. Iterate through each grid point E with index (i,j,k) in each sweeping direction or according to the fast marching heap sort.

 For each neighbor P<sub>l</sub> of E, compute u<sup>tmp</sup><sub>l</sub> = ||E − P<sup>\*</sup><sub>l</sub>||<sup>2</sup>.
 If ||E − P<sup>\*</sup><sub>l</sub>||<sup>2</sup> < min<sub>l</sub> u(P<sub>l</sub>), set u<sup>tmp</sup><sub>l</sub> = ∞. This is to enforce the monotonicity of the solution.

5. Set  $u(E) = \min_l u_l^{tmp} = u_{\lambda}^{tmp}$  and  $V(E) = P_{\lambda}^*$ .

# 4.5. A generalized closest point formulation

In the spirit of Steinhoff's Dynamic Surface Extension, we can define functions that map each point in  $\mathbb{R}^3$  to the space of representations of surfaces. Given a proper initialization of this "generalized" closest function, we can then compute the values of the function on the whole computational domain using the "sweeping" approach mentioned above.

In computational geometry and related fields such as computer graphics and computer vision, surfaces are often triangulated and stored as a set of triangles in 3D. This motivates our next generalization of the closest point algorithm.

Given a triangle T (or line segment S) in  $\mathbb{R}^3$  and a point P, the following algorithm explores the fact that we can easily compute the exact distance from P to T (or S) using elementary Euclidean geometry.

DEFINITION 4.4. (Piecewise linear surface element)

We define a piecewise linear surface element S to be a set of three ordered points in  $\mathbb{R}^3$ , and interpret the three points as the three vertices of a triangle. In case where two of the three points are identical, we say that S is a line segment. In the degenerate case where the three points are identical, we say that S is a point in  $\mathbb{R}^3$ .

We can further define the distance function of a point P and a surface element S as the minimum distance between P and the points on the triangle represented by S.

$$\operatorname{dist}(P,\mathcal{S}) := \min_{y \in T}(P,y)$$

where T is the triangle defined by S. Notice that this definition is valid even for the degenerate cases when S is a line segment or a point.

DEFINITION 4.5. (Generalized closest point function) We say W is a generalized closest point function if

 $W : \mathbb{R}^3 \to \{\text{the space of piecewise linear surface elements }\}.$ 

Let  $\Gamma$  be represented by a set of piecewise linear surface elements in  $\mathbb{R}^3$ . Let u be the distance function on the grids, and W be the corresponding generalized closest point function.

Algorithm 4 (The generalized closest point solver).

1. Initialize: give the exact distance to  $u_1$ , and the exact surface elements to W at grid points near  $\Gamma$ . Mark them so they will not be updated. Mark all other grid values as  $\infty$ . 2. Iterate through each grid point E with index (i,j,k) in each sweeping direction or

according to the fast marching heap sort.

3. For each neighbor  $P_l$  of E, compute  $u_l^{tmp} = \text{dist}(E, W(P_l))$ 4. If  $\text{dist}(E, W(P_l)) < \min_l u(P_l)$ , set  $u_l^{tmp} = \infty$ . This is to enforce the monotonicity of the solution.

5. Set 
$$u(E) = \min_l u_l^{tmp} = u_{\lambda}^{tmp}$$
 and  $W(E) = W(P_{\lambda})$ .

We show the results at the end of this paper. See figure 11 and figure 12.

#### 4.6. Analysis of the algorithms

Our experience shows that a satisfactory approximation can be reached after one set of sweeping iterations. We have not yet been able to prove convergence. However, we do see the limits to grid based computation of distance functions.

Both the spherical intersection method and the closest point approach described above are in essence trying to determine the closest point of E using the closest points of the neighbors of E. In order to get the exact solution  $(u(E) \text{ or } E^*)$ , we need the following property:

 $E^* \in \{P_i^* : P_i \text{ the neighbors of } E\}.$ 

However, this is not always possible. In fact, near the Voronoi boundary, we might reach the limits of our methods using the standard 4-point stencil. See figure 4.

Consider the following situation: Suppose  $E^*$  falls into the first quadrant (I). Let P be the grid such that  $dist(P, E^*) < dist(E, E^*)$ . We form a circle  $C_E$  centered at E, passing through  $E^*$ , and  $C_P$  centered at P, also passing through  $E^*$ . Let  $D_E, D_P$  be the corresponding disks. Then if  $\Upsilon = \Gamma \bigcap \{D_P \setminus D_E\} \neq \emptyset$ , then  $P^*$  has to be in it; because any point in  $\Upsilon$  satisfies the property that it is closer to P than  $E^*$  is. Similarly, we can construct the same scenario for Q.

As illustrated in figure 4, E is the only point in the neighborhood that is in the Voronoi region generated by  $E^*$ ; i.e.  $E^* \notin \{P^*, Q^*, (P')^*, (Q')^*\}$ . Hence, our updating procedure will not compute the exact distance at E in this case.

# Local Error Analysis

Let  $r_E := ||E - E^*||$  and  $\tilde{r}_E := ||E - P^*||$ . We want to know what the maximum error  $e := \tilde{r}_E - r_E$  can be, given exact values of u and V at the neighbors of E. From the illustration above, we can easily see that  $\max \tilde{r}_E < \max ||E^* - P|| + h$ , and

$$\max \|E^* - P\| < \sqrt{r_E^2 + h^2} = r_E + h^2 \frac{1}{2r_E} + \mathcal{O}(h^4).$$

Thus,  $0 \leq \tilde{r}_E - r_E < h$ . So the local error is  $\mathcal{O}(h)$ .

### The selection of stencils

In the algorithms described above, we have some criteria which eliminate invalid stencils. In the case where the value on a grid point is still infinity, we know automatically that it cannot lie in a valid stencil. In the case where the grid point carries "obviously incorrect"



FIG. 4. An example of the limit of grid based distance approximation

distance information, we may make use of the observation of the constant travel time of the eikonal equations: if the two neighboring grid points carry finite values and the difference of the values of the distance function on them is too large, we have an incompatibility. Since we want to enforce the upwinding stencil selection, we favor the grid point with the smaller value. Therefore, we have the following propositions:

PROPOSITION 4.1. Let  $P_1$  and  $P_2$  be two neighbors of E, and  $u(P_2) - u(P_1) > ||P_2 - P_1||$ . Then  $P_1^* \neq P_2^*$ .

*Proof.* Assume on the contrary that  $P_1^* = P_2^* = P^*$ , and  $u(P_2) > u(P_1)$ . Let Q be a point on  $\overline{P^*P_2}$  such that  $||Q - P^*|| = u(P_1)$ . Let R be a point on  $\overline{P_1P_2}$  such that  $\Delta P_2QR$  is a right triangle with hypotenuse  $\overline{RP_2}$ . Then  $||P_2 - P_1|| = |\overline{P_1P_2}| \ge |\overline{RP_2}| \ge |\overline{RP_2}| \ge |\overline{QP_2}| = u(P_2) - u(P_1)$ . Thus we have a contradiction. The proof is complete.



This proposition shows that if  $P_1^*$  is correct and  $u(P_2) - u(P_1) > ||P_2 - P_1||$ , then the information we hold on  $P_2$  is not correct.

### 5. RESULTS

We first run our algorithms to approximate the distance function of a single Voronoi generator at the center of a  $11 \times 11 \times 11$  grid. As shown in table 1, the spherical method gives an almost exact result with error close to machine zero. The closest point method gives the exact solution. Since we need to take a square root repeatedly in the first case, we expect to lose some accuracy. However, as demonstrated by the numerical results, the whole process of the spherical method appears to be stable.

A comparison of test case results			
Godunov method Spherical method	0.114 1.53e-15	0.091 2.99e-16	
Closest point method	0	0	

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We then tested our generalized closest point solver in both 2D and 3D with several different configurations. In the test cases our generalized closest point algorithm is exact to machine precision. We will describe our test cases in more detail in a following subsection.

In the following sections, we will "visualize" the distance functions by their level sets. In 2D, we plot the contour lines, and in 3D, we plot the isosurfaces.

FIG. 5. A comparison of results by Godunov method and spherical intersection method in 2D.



FIG. 6. A comparison of results by Godunov method and spherical intersection method in 2D.



# 5.1. A comparison of results in 2D

The following two sets of figures, figure 5 and figure 6, are computed in the sweeping fashion. We can clearly see that the hybrid algorithm gives straight contour lines in the Voronoi regions generated by straight line segments. In other regions, the contour lines have a nice circular shape.

#### 5.2. A comparison of results

We run the Godunov method and our methods to approximate the distance function to a set of 10 randomly selected isolated points. From table 3-4, we can see that the spherical method converges numerically with a second order rate while the closest point solver gives the exact distance function. There is a loss of accuracy in the solution obtained by using the spherical method. The error comes from two major sources. Within each Voronoi region, the computed solution loses accuracy from solving the quadratic equations. Near the Voronoi boundaries, it is not always true that the grid value we are updating has to be greater than or equal to the values of the stencils. By enforcing monotonicity of our solution near the Voronoi boundaries, we may therefore create a slight perturbation to the true solution. This perturbation propagates outwards from the Voronoi boundaries during later updating iterations. In the closest point algorithm for isolated points, we enforce a slightly relaxed monotonicity (step 4 and 5) on solution. Unless the situation mentioned in previous analysis occurs, we will obtain the exact distance function.

TABLE 2

Godunov method				
Grid	max error	convergence rate	$L_1$ error	convergence rate
10x10x10 (h=0.01)	0.008555		5.223e-6	
20x20x20	0.005723	0.58	3.343e-6	0.64
40x40x40	0.003656	0.65	2.059e-6	0.70
80x80x80	0.002311	0.66	1.248e-6	0.72

In figures 7-10 we show the comparison of the isosurfaces computed from the solutions of the Godunov solver and our methods. We have a 50x50x50 grid with uniform grid size h = 0.01, and 20 randomly selected isolated points. We can clearly see that the spherical intersection method gives much better results both in terms of  $L_1$  error and "visual" examination of the isosurfaces.

Spherical method				
Grid	max error	convergence rate	$L_1$ error	convergence rate
10x10x10 (h=0.01)	0.006869		1.500e-6	
20x20x20	0.002855	1.3	0.201e-6	2.9
40x40x40	0.001355	1.1	0.041e-6	2.3
80x80x80	0.000674	1.0	0.011e-6	1.9

TABLE 3

ТА	BL	Æ 4	1

#### **Closest point method**

Grid	max error	convergence rate	$L_1$ error	convergence rate
10x10x10 (h=0.01)	0		0	
20x20x20	0	$\infty$	0	$\infty$
40x40x40	0	$\infty$	0	$\infty$
80x80x80	0	$\infty$	0	$\infty$



FIG. 7. Isosurfaces computed by the Godunov solver

FIG. 8. Isosurfaces computed by the spherical intersection method





FIG. 9. Isosurfaces computed by the Godunov solver



FIG. 10. Isosurfaces computed by the spherical intersection method

 TABLE 5

 The generalized closest point method

Grid	max error	convergence rate	$L_1$ error	convergence rate
50x50 (h=0.02) 100x100 200x200	0 0 0	8 8	0 0 0	8

FIG. 11. Contour lines for the distance to a piecewise linear objects



# 5.3. Results of the generalized closest point sweeping

Figure 11 shows the contour lines of the computed distance function to a set of a closed polygon, two line segments and two isolated points. Table 5 shows, in turn, that the generalized closest point algorithm computes the distance to this set of objects to the machine precision after one sweeping iteration. Figure 12 shows the isosurfaces rendered from the computed distance function to a set of two polyhedra, three line segments and two isolated points.

# 6. CONCLUSION

In this paper, we have introduced two grid based algorithms for computing the distance function to a given configuration of isolated points, which are then generalized to compute the distance function to piecewise linear objects. They prove to be accurate, fast ( $\mathcal{O}(N)$ ) and easy to implement. The key to the algorithms is the combination of "sweeping," the upwinding-based stencil selection, and smarter geometric interpretation. Putting them all together leads to accurate and viscosity-solution-compliant approximations. The generalized closest point solver provides a quick translator of the triangulated data commonly used in computational geometry to the level set formulation. It also suggests a more general framework for the distance computation to more general surfaces.



FIG. 12. Isosurfaces of the distance to triangulated surfaces and isolated points



**FIG. 13.** The level contours of the distance function to a smooth parametrized curve constructed using Newton's method and sweeping

# 7. FUTURE WORK

Our first remark is that our algorithms (both spherical intersection and (generalized) closest point sweeping) can easily be generalized to other grid geometries since we are not approximating partial derivatives on the stencil. This topic will be investigated by the author.

Furthermore, from the adaptation of the closest point formulation, we have recognized the potential of computing the distance function to more general geometrical objects. We are currently seeking a generalized closest point formulation for computing more general surfaces in 3D. As indicated in the previous section, instead of storing only the closest point, we can also store a representation of the surface element;

 $V(E) \rightarrow (E^*, \text{ surface element }).$ 

The 'surface element' can be for example the curvature, or a NURB description of the surface. The challenge is to compute the exact distance to a given surface element and to derive the "upwinding" criteria for propagating the surface information throughout the grids. A preliminary result of one method using a conbination of Newton's method and sweeping iterations on a smooth parametrized curve segment if show in figure 13.

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