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Parallel redistancing using the Hopf–Lax formula

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A B S T R A C T

We present a parallel method for solving the eikonal equation associated with level set redistancing. Fast marching [1,2] and fast sweeping [3] are the most popular redistancing methods due to their efficiency and relative simplicity. However, these methods require propagation of information from the zero-isocontour outwards, and this data dependence complicates efficient implementation on today’s multiprocessor hardware. Recently an interesting alternative view has been developed that utilizes the Hopf–Lax formulation of the solution to the eikonal equation [4,5]. In this approach, the signed distance at an arbitrary point is obtained without the need of distance information from neighboring points. We extend the work of Lee et al. [4] to redistance functions defined via interpolation over a regular grid. The grid-based definition is essential for practical application in level set methods. We demonstrate the effectiveness of our approach with GPU parallelism on a number of representative examples.

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1. Introduction

The level set method [6] is a powerful technique used in a large variety of problems in computational fluid dynamics, minimal surfaces and image processing [7]. In general these methods are concerned with the transport evolution ⪿ = + · of a level set function : → in velocity field . While the essential property of is typically the location of its zero iso-contour ( = { ∈ | } = 0 ), in practice many applications additionally require that it be a signed distance function ( = 1). This property will not generally be preserved by the transport evolution, but it can be recovered without modifying the location of the zero isocontour. This process is commonly referred to as redistancing or reinitialization [8–10].

We describe the redistancing problem in terms of the signed distance function that is obtained from an arbitrary non-signed-distance level set function (while preserving the zero isocontour of ). Mathematically, the process obeys the eikonal equation as

\[ |\nabla \phi(x)| = 1 \]

\[ \text{sgn}(\phi(x)) = \text{sgn}(\phi^0(x)). \]

There is extensive previous work related to the solution of (1). The most commonly used methods are the fast marching method (FMM) [1,2] and the fast sweeping method (FSM) [3]. First proposed by Tsitsiklis [1] using optimal control, the
fast marching method was independently developed by Sethian in [2] based on upwind difference schemes. It is similar to Dijkstra’s method [11] for finding the shortest path between nodes in a graph. The fast marching method uses upwind difference stencils to create a discrete data propagation consistent with the characteristics of the eikonal equation. Sorting is used to determine a non-iterative update order that minimizes the number of times that a point is utilized to create a strictly increasing (or decreasing) propagation. The operation count is $O(N \log(N))$ where $N$ is the number of grid points and the $\log(N)$ term is a consequence of the sorting. Fast sweeping is similar, but it uses a simpler propagation. Rather than using the optimal update ordering that requires a heap sort, a Gauss–Seidel iterative approach is used with alternating sweep directions. Typically, grid axis dimensions are used as the sweep directions. For $\mathbb{R}^n$ it only requires $2^n$ propagation sweeps of updating to properly evaluate every point.

Notably, both the FMM and FSM approaches create data flow dependencies since information is propagated from the zero isocontour outwards and this complicates parallel implementation. Despite this, various approaches have achieved excellent performance with parallelization. The Gauss–Seidel nature of FSM makes it more amenable to parallelization than FMM. Zhao initially demonstrated this in [12] where each sweep direction was assigned to an individual thread with the final updated nodal value being the minimum nodal value from each of the threads. This method only allowed for a low number of threads and further scaling was achieved by splitting the individual sweeps into subdomain sweeps with a domain decomposition approach. However, this strategy can require more sweep iterations than the original serial FSM and the required iterations increase with the number of domains which reduces parallel efficiency. Detrixhe et al. [13] developed a parallel FSM that scales in an arbitrary number of threads without requiring more iterations than in the serial case. Rather than performing grid-axis-aligned Gauss–Seidel sweeps, they use Cuthill–McKee ordering (grid-diagonal) to decouple the data dependency. Since the upwind difference stencil only uses grid axis neighbors, nodes along a diagonal do not participate in the same equation and can thus be updated in parallel trivially. They extended these ideas to hybrid distributed/shared memory platforms in [14]. They use a domain decomposition strategy similar to Zhao [12] to divide the grid among available compute nodes and a fine grained shared memory method within each subdomain that utilizes their approach in [13] to achieve orders of magnitude performance increases.

FMM is more difficult to implement in parallel, however even Tsitsiklis [1] developed a parallel FMM algorithm using a bucket data structure. A number of approaches use domain decomposition ideas similar to Zhao [12] and Detrixhe et al. [14] to develop parallel FMM [15–18]. In these approaches the grid is typically divided into disjoint sub grids with a layer of ghost nodes continuing into adjacent neighbors. Each sub grid is updated in parallel with the FMM update list typically leading to rather elaborate communication between threads. Jeong et al. [17] developed the fast iterative method (FIM), which is a parallel approach using domain decomposition but with a looser causal relationship in the node update list to localize updates for Single Instruction Multiple Data (SIMD) level parallelism. Simplifications to the update list in FMM improve parallel scaling, but tend to increase the number of worst case iterations. Dang et al. [19] extended FIM to a coarse/fine-grained approach based on domain decomposition with load balancing via master/worker model that allowed for efficient performance on heterogeneous platforms.

Recently an interesting alternative to FMM and FSM has been proposed. Darbon and Osher [5] and Lee et al. [4] utilize the Hopf–Lax formulation of the solution to the Hamilton–Jacobi form of the eikonal equation. Notably, the signed distance at an arbitrary point is obtained without the need of distance information from neighboring points. This allows for the solution at any given point in any order and prevents the need for communication across cores which greatly simplifies parallel implementation. Furthermore, this inherently allows for updates done only in a narrow band near the zero-isocoutour. FSM must solve over the entire domain, and while FMM can be done in a narrow band, FMM methods are generally more difficult to implement in parallel. These aspects make the Hopf–Lax approaches in [4,5] very compelling for parallel architectures. In this paper, we extend the work of Lee et al. to handle functions defined via interpolation over a regular grid. Lee et al. demonstrated compelling results with abstractly defined functions. However, treatment of grid-based functions is essential for practical application in level set methods. We demonstrate the effectiveness of our approach with Graphics Processing Unit (GPU) parallel implementation.

### 2. Method

Following Lee et al. [4] we use the Hamilton–Jacobi formulation of the eikonal equation (1)

\[
\frac{\partial}{\partial t} \tilde{\phi}(x,t) + \| \nabla \tilde{\phi}(x,t) \|_2 = 0
\]

(2)

for $x \in \mathbb{R}^n, t > 0$. We assume that $\phi^0$ is constructed such that

\[
\begin{align*}
\phi^0(x) < 0 & \quad x \in \Omega \setminus \partial \Omega \\
\phi^0(x) > 0 & \quad x \in (\mathbb{R}^n \setminus \Omega) \\
\phi^0(x) = 0 & \quad x \in \partial \Omega
\end{align*}
\]

for some set $\Omega \subset \mathbb{R}^n$. As in Lee et al. [4] we assume that the set $\Omega$ is closed and non empty. Isocontours of the time dependent solution $\phi$ progress from the boundary $\partial \Omega$ in its normal direction at a rate of $1$. To know the distance to the
Algorithm 1 Modified secant method.

```
while |φ(X_i, t^{k+1})| > ε do
    Δt = |φ(X_i, t^k) - φ(X_i, t^{k-1})| / |φ(X_i, t^k) - φ(X_i, t^{k-1})|
    if |Δt| > t₀ then
        if φ(X_i, t^k) > 0 then
            Δt = -Δmax
        else
            Δt = -Δmax
        end if
    end if
    t^{k+1} = t^k + Δt
end while
```

boundary, we simply need to know at which time \( \hat{t} \) the zero-isocontour of \( \hat{\phi} \) has progressed to the point \( x \). In other words, the signed distance \( \phi(x) \) from the point \( x \) to the boundary \( \partial \Omega \) is given by the time \( \hat{t} \) with \( \phi(x, \hat{t}) = 0 \): \( \phi(x) = \hat{t} \). Note that we only consider the case here of positive \( \phi \) since the case of negative \( \phi \) is trivially analogous.

As in Lee et al. [4], we treat the problem as root finding and use the secant method. However, unlike Lee et al. we are specifically interested in redistancing grid based functions. Thus we assume that the initial function is defined in terms of its interpolated values from grid nodes as \( \phi^0(x) = \sum_i \hat{N}_i(x) \) where the function \( \hat{N}_i \) is the bilinear interpolation kernel associated with grid node \( x_i \) and \( \phi^0_i = \phi^0(x_i) \). Also, when we solve for the redistanced values, we do so only at grid nodes (i.e. we solve for \( \hat{\phi}_I = \phi(x_I) = \hat{t}_I \). Thus the secant method only requires the evaluation of the function \( \phi(x_I, t^k) \) for iterative approximation \( t^k \rightarrow \hat{t} \). We next discuss the practical implementation of the secant method and evaluation of \( \phi(x_I, t^k) \) for grid based data.

2.1. Secant method for roots of \( \hat{\phi}(x_I, \hat{t}) = 0 \)

In order to use the secant method to solve for the root in this context we use the iterative update

\[
t^{k+1} = t^k - \frac{\hat{\phi}(x_I, t^k) - t^k - t^{k-1}}{\hat{\phi}(x_I, t^k) - \hat{\phi}(x_I, t^{k-1})}.
\]

The initial guess \( t^0 \) can either be set from neighboring nodes that have been recently updated, or generally from a priori estimates of the distance (see Section 2.3). However, when no such information is possible or when it would negatively affect parallel performance we use \( t^0 = 0 \). We set \( t^1 = t^0 + \epsilon \) where \( \epsilon \) is proportionate to the grid cell size.

The main concern with using the secant method in this context is that while \( \phi(x_I, t) \) is monotonically decreasing in \( t \), it is not strictly monotone. This means that there can be times \( t \) where \( \frac{d}{dt} \phi(x_I, t) = 0 \). For example, if the minimum of \( \phi^0(x_I) \) over the ball centered at \( x_I \) of radius \( t \) is in the interior of the ball (at a point of distance \( s \) from \( x_I \)), then \( \frac{d}{dt} \phi(x_I, r) = 0 \) for \( s \leq r \leq t \) (see Section 2.2). The secant method is not well defined if we have iterates with equal function values. To compensate for this, if secant would divide by zero, and we have not already converged, we simply increase or decrease \( t^{k+1} = t^k \pm \Delta t_{\text{max}} \) in the correct direction. The correct direction is trivial to find, because if \( \phi(x_I, t^k) > 0 \) then we need to increase \( t^k \). Otherwise we need to decrease \( t^k \). In practice, we use \( \Delta t_{\text{max}} = 5 \Delta x \) where \( \Delta x \) is the grid cell size.

Another issue is that errors in the approximation of \( \phi(x_I, t^k) \) can lead to more secant iterations. This can be reduced by solving for \( \phi(x_I, t^k) \) to a higher tolerance. However, requiring more iterations to approximate \( \phi(x_I, t^k) \) more accurately can be more costly than just using more secant iterations with a less accurate (but more efficient) approximation to \( \phi(x_I, t^k) \). We discuss the process and cost of solving for \( \phi(x_I, t^k) \) in Section 2.2. Our modified secant algorithm is summarized in Algorithm 1.

2.2. Hopf–Lax formula for \( \phi(x_I, t^k) \)

As in Lee et al. [4] we obtain the solution of Equation (2) with the Hopf–Lax formula

\[
\hat{\phi}(x_I, t^k) = \min_{y \in \mathbb{R}^n} \left\{ \phi^0(y) + t^k H^*(\frac{x_I - y}{t^k}) \right\}
\]

where \( H^* \) is the Fenchel–Legendre Transform of \( H = \| \cdot \|_2 \)

\[
H^*(x) = \begin{cases} 0 & \|x\|_2 \leq 1 \\ \infty & \text{otherwise} \end{cases}
\]

or equivalently

\[
\hat{\phi}(x_I, t^k) = \min_{y \in B(x_I, t^k)} \phi^0(y)
\]
where $B(x_i, t^j)$ is the ball of radius $t^j$ around grid node $x_i$. Thus the problem of evaluating $\hat{\phi}(x_i, t^j)$ amounts to finding the minimum of the initial $\phi^0$ over a ball. While Lee et al. [4] use Split Bregman iteration to solve this, we instead simply use projected gradient descent. We used a few hundred projected gradient iterations in practice since this was faster than Split Bregman in parallel implementations due to its relative simplicity. Using $y_k^j$ as an initial guess for the argmin of $\phi^0$ over the ball $B(x_i, t^j)$, we iteratively update the approximation from

$$y_k^{j+1} = y_k^j - \gamma \nabla \phi^0(y_k^j)$$

$$y_k^{j+1} = \text{PROJ}_{B(x_i, t^j)}(y_k^{j+1})$$

where

$$\text{PROJ}_{B(x_i, t^j)}(y) = \begin{cases} y & \|x_i - y\|_2 \leq t^j \\ x_i - \frac{x_i - y}{\|x_i - y\|_2} t^j & \text{otherwise} \end{cases}$$

In practice, we set the step size $\gamma$ equal to the grid spacing $\Delta x$. Note that the gradients $\nabla \phi^0(y_k^j)$ are computed using the bilinear interpolation kernels $N_l(x)$ as $\nabla \phi^0(y_k^j) = \sum_l \phi_l^0 \nabla N_l(y_k^j)$. We emphasize that for efficiency the sum over $l$ can be taken over only the four grid nodes surrounding the cell that the argument $y_k^j$ is in. We further note that the index for the cell containing the argument $y_k^j$ can be found in constant time using $\text{floor}(\frac{x}{\Delta x})$ where $y_{nk}^j$ are the components of $y_k^j$. In general, $\phi^0$ is a non-convex function defined from the grid interpolation and projected gradient descent will only converge to a local minimum. We illustrate this in Fig. 1. Failure to converge to a global minimum can lead to large errors in the approximation of $\hat{\phi}(x_i, t^j)$. While it is impractical to ensure we achieve a global minimizer, it is possible to find multiple local minimizers increasing the probability we find a global minimizer. We solve (4) multiple times with different initial guesses $y_k^0$ and then take the minimum over these solutions to come up with a final answer that is likely to be close to a global minimizer. We found in practice, on the order of one guess per grid cell in the ball $B(x_i, t^j)$ is sufficient to find a global minimizer. For problems without many local extrema the number of initial guesses can be reduced. In general when finding $\hat{\phi}(x_i, t^j)$ we use as initial guesses $\text{PROJ}_{B(x_i, t^j)}(y_{k-1})$ where

$$y_{k-1} = \text{argmin}_{y \in B(x_i, t^{k-1})} \phi^0(y)$$

is the argmin of $\phi^0$ used in the previous secant iteration as well as a small number of random points in $B(x_i, t^j)$. (See Fig. 2.) We use this strategy because $\text{PROJ}_{B(x_i, t^j)}(y_{k-1})$ tends to be a good guess for the global minimum. In general, it is very likely that at the next step, the minimum will either be the same point, or along the boundary. Therefore, we prioritize random initial guesses near the boundary of the ball. In fact, for $t^{k-1} < t^j$ we know that the argmin will be at a distance $s$ from $x_i$ with $t^{k-1} \leq s \leq t^j$ so in theory we should only sample in the annulus. However, in practice we do not have full confidence in the argmin attained at iteration $k - 1$ since our procedure is iterative. Allowing for initial guesses at some locations closer to $x_i$ than $t^{k-1}$ admits the possibility of finding a more accurate argmin. Thus, we found that skewing the sampling density to be higher towards the boundary of the ball struck a good balance between efficiency and accuracy. We illustrate this process in Fig. 4.
Fig. 2. The figure illustrates representative random initial guesses used in solving for \( \tilde{\phi}(x_i, t^1) \). In addition we use an initial guess equal to the minimizer computed in the previous secant iteration shown in magenta. (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)

Fig. 3. The plots above are the points \((\tilde{\phi}(x_i, t^1), t^1)\) found when running our algorithm with different choices of random guess and gradient descent iterations on circle initial data. The left most plot was run with 100 random guesses, and 1 gradient descent iteration. The middle plot was run with 1 random guess, and 100 gradient descent iterations. The right plot was run with 1 random guess and 5 gradient descent iterations. Note that in all cases, the correct root was found.

Failure to find the global minimum over the ball can cause unpredictable behavior in the secant iteration for \( \hat{t} \). This includes false positives where a \( t^k \) is incorrectly interpreted as a root. However, continuing to run secant to a fixed large number of iterations usually corrects for this. In general, there is a tradeoff between the number of initial guesses and iterations of projected gradient and the number of secant iterations. We illustrate this in Fig. 3 which shows the path to convergence for a few choices of iteration parameters. When \( \tilde{\phi}(x_i, t^k) \) is solved with high accuracy, the secant iteration converges with minimal overshoot in 7 iterations. When \( \tilde{\phi}(x_i, t^k) \) is not solved to high accuracy, secant overshoots by a large margin, and takes 16 iterations to converge, but notably still converges. However because each iteration is cheaper, the total runtime is lower to reach the same convergence for \( \hat{t} \). In practice we found that a few hundred projected gradient iterations combined with our initial guess sampling strategy struck a good balance between accuracy and efficiency.

2.3. Computing in a narrow band

In many applications, only data close to the interface is needed. Since each grid node can be solved for independently, the Hopf–Lax approach naturally lends itself to narrow banding strategies for these applications. We provide a narrow banding strategy based on a coarse initial grid computation followed by a fine grid update in the narrow band. We first redistance the function on the coarse grid, then we interpolate values from the coarse grid to the fine grid. We then only recompute values on the fine grid that are smaller than a threshold value and we use the value interpolated from the coarse nodes as an initial guess \( t_0 \) for the computation on the fine grid. As an example see Fig. 5.
2.4. Computing geometric quantities

The Hopf–Lax formulation naturally allows us to compute normals \( \mathbf{n} = \nabla \phi \) and curvatures \( \nabla \cdot \mathbf{n} \) at the grid nodes. As pointed out in Lee et al. [4], as the argmin \( \mathbf{y}^k \) from Equation (5) is iterated to convergence, it approaches the closest to point to the grid node \( x_i \) on the zero isocontour of \( \phi^0 \). Therefore, recalling that when \( t^k \) has converged (within a tolerance) to the root \( t \) of \( \phi(x, t) = 0 \), \( t^k \) is approximately the distance to the zero isocontour, we can compute the unit normal at the grid node from

\[
\mathbf{n}(x_i) = \frac{x_i - y^k}{t^k}.
\]

Notably, this approximation is very close to the exact signed distance function with zero isocontour determined by the bilinearly interpolated \( \phi^0 \). It is as accurate as the argmin \( \mathbf{y}^k \) so it essentially only depends on the accuracy of the secant method. We get this very accurate geometric information for free. Moreover, the curvature \( \nabla \cdot \mathbf{n} \) can be computed accu-
3. Results

All of the following results were run on an Intel 6700k processor with an Nvidia GTX 1080. The domain for each problem was set to be [0, 1] x [0, 1] and was run on a 512 x 512 grid. To ensure efficient performance on the GPU, both projected gradient descent and the secant method were run for a fixed number of iterations rather than to a specific tolerance. All timings listed in this section are averages over 5 separate runs, with the exception of the Vortex problem which is already
Union of circles: the initial data is \( \phi^0 = \max(0.25 - \| (0.3, 0.5) - (x, y) \|_2, 0.75 - \| (0.7, 0.5) - (x, y) \|_2) \). The zero level set is a union of two circles both with radius .25, one centered at (0.3, 0.5) and the other centered at (0.7, 0.5).

Many local minima: the objective function is \( \phi^0 = \sin(4 \pi x) \cdot \sin(4 \pi y) - 0.01 \). The zero level set is a group of rounded squares that almost touch.

an average. This is due to the fact that we noticed in practice variations of up to 10% in the individual runtimes of a given problem

<table>
<thead>
<tr>
<th>Problem</th>
<th>num_secant</th>
<th>num_rand</th>
<th>num_proj</th>
<th>Timing (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circle</td>
<td>10</td>
<td>5</td>
<td>100</td>
<td>47.567</td>
</tr>
<tr>
<td>Two points</td>
<td>10</td>
<td>5</td>
<td>100</td>
<td>45.199</td>
</tr>
<tr>
<td>Vortex (per frame)</td>
<td>10</td>
<td>5</td>
<td>200</td>
<td>73.248</td>
</tr>
<tr>
<td>Square</td>
<td>10</td>
<td>4</td>
<td>200</td>
<td>67.582</td>
</tr>
<tr>
<td>Sine</td>
<td>10</td>
<td>5</td>
<td>200</td>
<td>71.429</td>
</tr>
</tbody>
</table>

Fig. 6 shows initial data \( \phi^0 \) with a zero-isocountour given by a circle with radius .25. Fig. 7 shows a more complicated test. The zero-isocountour is a square bounded between [.25, .75] in x and y. The corners present rarefaction fans, and the inside
Table 1
Timing and error at different grid resolutions using a square as our zero level set. The average L2 error is calculated by calculating the L2 distance between the computed answer and the analytic solution.

<table>
<thead>
<tr>
<th>Size</th>
<th>Timing (ms)</th>
<th>Average L2 error</th>
</tr>
</thead>
<tbody>
<tr>
<td>32 × 32</td>
<td>3.303</td>
<td>6.67 × 10⁻⁵</td>
</tr>
<tr>
<td>64 × 64</td>
<td>3.392</td>
<td>1.5917 × 10⁻⁵</td>
</tr>
<tr>
<td>128 × 128</td>
<td>4.477</td>
<td>3.8723 × 10⁻⁶</td>
</tr>
<tr>
<td>256 × 256</td>
<td>17.8400</td>
<td>9.7391 × 10⁻⁷</td>
</tr>
<tr>
<td>512 × 512</td>
<td>67.533</td>
<td>2.4624 × 10⁻⁷</td>
</tr>
<tr>
<td>1024 × 1024</td>
<td>274.216</td>
<td>6.4343 × 10⁻⁸</td>
</tr>
<tr>
<td>2048 × 2048</td>
<td>1185.87</td>
<td>2.3843 × 10⁻⁸</td>
</tr>
</tbody>
</table>

Table 2
Timing in ms showing scaling in number of GPUs. The first two columns show the time it takes to run our problem when we include the timing cost of transferring the grid data to the GPU (approximately 1.2 ms), while the last two columns show the scaling without the cost of transferring data.

<table>
<thead>
<tr>
<th># GPUs</th>
<th>Total (w/)</th>
<th>Per GPU (w/)</th>
<th>Total (w/o)</th>
<th>Per GPU (w/o)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>125.10</td>
<td>125.10</td>
<td>124.05</td>
<td>124.05</td>
</tr>
<tr>
<td>2</td>
<td>126.25</td>
<td>63.13</td>
<td>124.76</td>
<td>62.38</td>
</tr>
<tr>
<td>4</td>
<td>130.17</td>
<td>32.54</td>
<td>124.92</td>
<td>31.23</td>
</tr>
<tr>
<td>8</td>
<td>139.26</td>
<td>17.41</td>
<td>131.10</td>
<td>16.39</td>
</tr>
<tr>
<td>16</td>
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<tr>
<td>32</td>
<td>167.97</td>
<td>5.25</td>
<td>133.07</td>
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</tr>
<tr>
<td>64</td>
<td>206.07</td>
<td>3.22</td>
<td>134.85</td>
<td>2.11</td>
</tr>
</tbody>
</table>

Fig. 10. Parallel speed up is plotted both with and without including the cost of updating memory on the GPU. With 64 GPUs the memory update can take up to 33% of the runtime. However without the memory update [i.e. if the data is already on the GPU] the method scales simply.

needs to handle sharp corners along the diagonals. Because of these difficulties (especially the sharp gradient in our original interpolated \(\phi^0\)), more work is needed in resolving the projected gradient descent step to ensure quick convergence of secant method. The zero-isocountour shown in Fig. 8 is the union of two overlapping circles. Like in Fig. 6 the gradient is fairly smooth and thus requires less computation to successfully converge in gradient descent. In Fig. 9 we demonstrate our algorithm with a large number of local minima. This problem requires more projected gradient iterations than the simpler examples.

Fig. 11 shows our method being used in a level set advection scheme using a simple vortex test. Like previous problems it was run on a 512 × 512 grid. For this problem the average time per frame for redistancing was 73.248 ms.

3.1. Scaling

The results in Table 1 was run with the square given in Fig. 7 with the same parameters. The poor scaling at the low resolutions is due to not using all of the threads possible on the GPU.

For Table 2 we ran our algorithm on the initial data in Fig. 6 with a 1024 × 1024 grid. The problem was broken up into sub domains and each domain was run separately on the GPU. The performance results are shown in Fig. 10. The scaling is nearly optimal, but breaks down at high number of GPUs when we include the time it takes to transfer the data to the GPU. The transfer time takes approximately 1.2 ms. If we ignore the time it takes to transfer data to the GPU we get a result that is close to being perfectly parallel. We also show this in Fig. 10.
Fig. 11. Practical application: vortex advection test at $t = 0, 1, 2, 3, 4, 5$.

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