Parallel Computation of 3D Morse-Smale Complexes

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Abstract

The Morse-Smale complex is a topological structure that captures the behavior of the gradient of a scalar function on a manifold. This paper discusses scalable techniques to compute the Morse-Smale complex of scalar functions defined on large three-dimensional structured grids. Computing the Morse-Smale complex of three-dimensional domains is challenging as compared to two-dimensional domains because of the non-trivial structure introduced by the two types of saddle criticalities. We present a parallel shared-memory algorithm to compute the Morse-Smale complex based on Forman’s discrete Morse theory. The algorithm achieves scalability via synergistic use of the CPU and the GPU. We first prove that the discrete gradient on the domain can be computed independently for each cell and hence can be implemented on the GPU. Second, we describe a two-step graph traversal algorithm to compute the 1-saddle-2-saddle connections efficiently and in parallel on the CPU. Simultaneously, the extrema-saddle connections are computed using a tree traversal algorithm on the GPU.

Categories and Subject Descriptors (according to ACM CCS):
I.3.5 Computational Geometry and Object Modeling

1. Introduction

The Morse-Smale (MS) complex partitions the domain of a scalar function into regions with uniform gradient behavior. It provides an abstract representation that enables multi-scale analysis and visualization of 2D and 3D scalar functions [LBM\(^*\)06, GDN\(^*\)07, RKG\(^*\)11, KRHH11]. Motivated by the increasing data sizes, recent approaches towards the computation of MS complexes focus on memory efficiency and scalability in addition to performance [GBPH08, GRWH11, PRG\(^*\)11]. We present a parallel shared-memory algorithm to compute the MS complex of a three-dimensional scalar function. Our algorithm makes a synergistic use of the multi-core CPU and GPU of a desktop computer to compute the MS complex of large three-dimensional data sets. Experiments indicate that this approach can process data consisting of over 1 billion vertices within minutes while utilizing less than 2GB of memory.

1.1. Related work

MS complexes were initially introduced in the context of dynamical systems [Sma61a, Sma61b]. Later, Edelsbrunner et al. [EHZ03] studied the problem of computing the MS complex for two-dimensional piecewise linear functions. They computed the cells of the MS complex while restricting the bounding arcs of the complex to edges of the input mesh. This approach was extended to construct MS complexes of three-dimensional functions [EHNP03]. These early approaches traced the gradient paths from saddle critical points and produced a boundary representation of cells in the MS complex. Bremer et al. [BEHP04] also followed this approach and developed a multi-resolution representation of 2D scalar functions via controlled topological simplification, and demonstrated the application of the MS complex to feature identification, noise removal, and view-dependent simplification.

Gyulassy et al. [GDN\(^*\)07, GNP\(^*\)06] focused on three-dimensional functions and employed an approach based on repeated cancellations of critical point pairs applied on an artificial complex created from the input mesh by including dummy critical points. The cancellations were appropriately scheduled in order to remove the dummy critical points leaving behind the true critical points and cells of the MS complex. The order of cancellation determines the quality of the resulting MS complex and the algorithm efficiency.
Another approach to compute the MS complex is based on a discrete analog of Morse theory [Mil63] introduced by Forman [For02] to study discrete functions defined on cells of a cell complex. Reininghaus et al. [RH11, RLH11] discussed an application of discrete Morse theory to analyze vector fields. Bauer et al. [BLW11] computed simplified two-dimensional scalar functions while ensuring that the input function is modified by no more than a threshold δ and all surviving critical point pairs have persistence greater than 2δ. Caflisch et al. [CCL03] and Lewiner et al. [LLT04] successfully employed Forman’s discrete Morse theory to compute MS complexes of piecewise-linear functions and demonstrated applications to segmentation, visualization, and mesh compression. Gyulassy et al. [GBPH08] also used a discrete Morse theory based formulation to develop an efficient algorithm for computing MS complexes of large 3D data that do not fit in main memory. They partition the data into blocks called “parcels” that fit in memory, compute gradient flows on the boundary of the parcels, propagate the flows to the interior and compute the MS complex restricted to a parcel. Critical cells that are created on the boundary are canceled during a subsequent merge step resulting in the MS complex of the union of the parcels. This serial method scales well for large data but the geometry of the MS complex is sensitive to the order of cancellations chosen during the merge step.

Robins et al. [RWS11] proposed an algorithm to compute the Morse complex of 2D and 3D gray-scale digital images modeled as discrete functions on cubical complexes. The algorithm computes the Morse complex with provable guarantees on its correctness with respect to the critical cells. However, the algorithm does not guarantee polynomial time execution since they use a modified breadth first search algorithm that traverses all possible paths between two nodes, which can be exponential in the number of nodes.

More recently, Peterka et al. [PRG∗11] introduced a set of building blocks for implementing parallel algorithms, which leverage high performance computing clusters. In particular, they discuss a parallel implementation of the discrete Morse theory based algorithm proposed by Gyulassy et al. [GBPH08] using their framework. Unlike the work of Peterka et al., we focus on a parallel implementation on a desktop computer. Günther et al. [GRWH11] described a memory efficient algorithm to compute the MS complex for 3D data and use the complex to compute persistent homology groups. The discrete gradient field is computed using a parallel variant of the method proposed by Robins et al. followed by an efficient computation of the boundary map that represents the MS complex. They employ a modified breadth first search based algorithm to traverse the gradient field, the maintained of visited flag per cell for each traversal. Thus the number of traversals that can be launched in parallel is limited by available memory. In comparison, our parallel algorithm for computing the discrete gradient field is based on a novel design of the discrete Morse function followed by a two-step algorithm to compute the cells of the MS complex. Our hybrid multi-core approach for implementing this algorithm results in a method that is fast in addition to being memory efficient.

1.2. Contributions
The above mentioned methods for computing the MS complex are either slow or not easily applicable because of one or more of the following reasons: (a) they compute and trace the gradient serially, (b) they do not guarantee that they trace the correct geometry of the gradient flow, or (c) they require specialized computing resources such as HPC clusters.

Our contributions to solve the above problems for scalar functions on three-dimensional structured grids is based on a characterization of MS complexes using Forman’s discrete Morse theory that leads to a parallel algorithm. We describe an extension of the parallel discrete gradient field construction algorithm introduced earlier [SMN11] to compute 2D MS complexes. The presence of two types of saddle criticalities in 3D introduces considerable complexity into the geometry of the MS complex. The discrete gradient paths between saddles may split and merge causing an explosion in the number of paths to be traced. Moreover, the data sizes are much larger compared to 2D scalar functions. The key contributions of this paper towards efficient computation of 3D MS complexes are

- An extension of the independent gradient pairing algorithm discussed in [SMN11] that significantly reduces the number of cells that remain unpaired, thereby producing fewer artifacts in the form of spurious critical cells.
- An efficient two-step algorithm to compute gradient paths between 1-saddles and 2-saddles.
- Synergistic use of CPU and GPU to ensure maximum utilization of computational resources.
- Implementation of a split and merge approach that supports computation of MS complexes for large datasets.

We present experimental results on both synthetic and real-world data sets to demonstrate the efficiency of the algorithm in terms of performance and memory requirement.

2. Background
This section reviews the necessary background on Morse functions and discrete Morse functions required for the algorithm description. While the focus of the paper is on 3D domains, we often consider 2D domains for ease of illustration and to describe prior work.

2.1. Morse functions
Consider a smooth scalar function \( f : \mathbb{R}^3 \to \mathbb{R} \). A point \( p \in \mathbb{R}^3 \) is called a critical point with respect to \( f \) if the gradient of \( f \), \( \nabla f \), is identically zero at \( p \). A critical point is non-degenerate if the Hessian of \( f \), equal to the matrix of second
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Maxima.

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edional manifold, where as its descending manifold is a d-dimensional manifold. A Morse function f is called a

Morse-Smale function if all of its critical points are non-degenerate. The index of a critical point is the number of negative eigenvalues of the Hessian matrix. An integral line passing through a point p is a maximal curve in the domain, whose tangent at every point equals the gradient of f at that point. The function f increases along the integral line and its limit points are the critical points of f.

The set of all integral lines that share a common source to-

tgether with the point p, is called the ascending manifold of p and the set of all integral lines that share a common destination together with the point p is called the descending manifold of p, see Figure ?? . The ascending manifolds (similarly, the descending manifolds) of all critical points partition the domain. The Morse-Smale complex is a partition of the domain into cells formed by the collection of integral lines that share a common source and a common destination. The ascending manifold of a critical point of index d is a \((n-d)\)-dimensional manifold, where as its descending manifold is a d-dimensional manifold. A Morse function f is called a Morse-Smale function if all ascending and descending manifolds of two critical points intersect transversally. Thus, if the index of two critical points differ by one then their as-

cending / descending manifolds either do not intersect or intersect along a one-dimensional manifold connecting the critical points. The critical points, referred to as nodes, together with the 1-manifolds that connect them, referred to as arcs, form the 1-skeleton of the MS complex, which is referred to as the combinatorial structure of the complex.

2.2. Simplification

A Morse-Smale function f can be simplified to a smoother function by repeated application of a cancellation operation that removes a pair of critical points connected by an arc in the MS complex. Critical point pairs correspond to topological features and are ordered based on the notion of persistence, which is equal to the absolute difference in function value between the two critical points. Persistence measures the importance of a critical point pair [ELZ02]. The least persistent critical point pair is always connected by an arc in the MS complex [EHZ03].

Cancellation of a pair of critical points is achieved by a local smoothing of the function within the ascending / descending manifolds containing the critical points. For example, consider the case of a two-dimensional Morse-Smale function after a maximum-saddle cancellation (see Figure 2). The 1-skeleton is updated by deleting the two corresponding nodes, deleting the arcs incident on the saddle, and re-routing the arcs incident on the maximum to the surviving maximum adjacent to the saddle. The embedding of a new arc is obtained by extending the old arc along the arc between the removed maximum and the saddle. We allow only cancellations between a pair of critical points that are connected by a single arc. Canceling a pair of critical points

order partial derivatives, is non-singular. We call f a Morse function if all of its critical points are non-degenerate. The index of a critical point is the number of negative eigenvalues of the Hessian matrix. An integral line passing through a point p is a maximal curve in the domain, whose tangent at every point equals the gradient of f at that point. The function f increases along the integral line and its limit points are the critical points of f.

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Figure 1: (a)-(d) Behavior of a Morse-Smale function near the critical points in a 3D domain. (e) The reversed integral lines of a 2D function. (f) The descending manifold of a maximum shown as the closure of the set of reversed integral lines that originate from the critical point. (g) Combinatorial structure of the MS complex where nodes are critical points and connecting integral lines are arcs.

Figure 2: (a) MS complex for a simple height function. (b) Canceling a pair of critical points, \(q_i, p_{i+1}\), of index \(i, i+1\) that are connected by a single arc. (c) Combinatorial realization: compute the union of the descending manifold of \(p_{i+1}\) with the descending manifolds of all index \(i+1\) critical points connected to \(q_i\). Compute the union of the ascending manifold of \(q_i\) with the ascending manifolds of all index \(i\) critical points connected to \(p_{i+1}\).
3: A 2D discrete Morse function. Gradient pairs are shown as arrows oriented towards higher dimensional cells. Dashed red arrows denote edge-quad pairs and solid blue arrows denote vertex-edge pairs. Critical cells are shown in red (maxima), green (saddle), and blue (minima).

that are connected by two distinct arcs in the Morse-Smale complex results in a strangulation, which cannot be realized by a local smoothing of the function [GNP’06].

2.3. Discrete Morse functions

Discrete Morse theory was developed by Forman [For02] to study the topology of cell complexes. A d-cell $α^d$ is a topological space homeomorphic to a d-ball $B^d = \{ x \in \mathbb{R}^d : |x| \leq 1 \}$. For example, a vertex is a 0-cell, an edge between two vertices is a 1-cell, a polygon is a 2-cell, and in general a d-dimensional polytope is a d-cell. We will restrict our attention to cells of the above kind, which can be represented by a set of vertices. A cell $α$ is a face of $β$, denoted $α < β$, if $α$ is represented by a subset of vertices of $β$. The cell $β$ is called a coface of $α$. A face $α$ is called the set of $β$ if $α < β$ and $\text{dim}(α) + 1 = \text{dim}(β)$. In this case $β$ is a cofacet of $α$ denoted by $α < β$. The set of zero-dimensional faces of a cell $α$ is called the vertex set of $α$ denoted by $V_α$.

A cell complex $K$ is a collection of cells that satisfies two properties: (a) If $α$ belongs to $K$ then so do all faces of $α$, and (b) If $α_1$ and $α_2$ are two cells in $K$ then either they are disjoint or they intersect along a common face. A regular cell complex is a cell complex in which, given two incident cells, $β^{d+1}$ and $α^{d−1}$, there are exactly two cells $α^d$, $α^d_1$ such that $γ^{d−1} < α^d_1$, $α^d_2 < β$. In this paper, we consider only finite regular cell complexes.

Given a regular cell complex $K$ representing the domain, a function $f : K \rightarrow \mathbb{R}$ is said to be a discrete Morse function if for all $d-\text{cells} \; α^d \in K$, (a) at most one of its cofacets has a lower function value, and (b) at most one of its facets has a higher function value (see Figure 3). A cell is critical if none of its cofacets has a lower function value and none of its facets has a higher function value. A discrete vector, referred to as a gradient pair, is a pairing between two incident cells that differ in dimension by one. A discrete vector field on $K$ is a set of discrete vectors such that every cell in $K$ is represented in at most one pair of the field. A V-path is a sequence of cells $α^d_0\;β^{d+1}_0\;α^d_1\;β^{d+1}_1\ldots\;α^d_{r−1}\;β^{d+1}_{r−1}$ such that $α^d_0$ and $α^d_{r−1}$ are faces of $β^{d+1}_0$ and $β^{d+1}_{r−1}$ is a vector for all $i = 0..r$. A V-path is called a gradient path if it contains no cycles. A discrete gradient field is a discrete vector field that contains no non-trivial closed V-paths.

4: The weighted discrete function is defined recursively as a weighted sum of the function value at facet $G_0$ and face $G_1$. Facets $G_0$ (in red) and faces $G_1$ (in blue) for an edge, triangle, and quad cell are shown. The function value at vertices increases along the vertical axis.

Ascending / descending manifolds are similarly defined for discrete Morse functions.

3. 2D MS complex Algorithm

In this section, we briefly describe the algorithm for 2D MS-complexes by [SMN11], without proof and detailed justifications. In the next section, we describe how the algorithm can be extended to three-dimensional scalar functions focusing on the additional challenges present in 3D. Given an input scalar function $f$, it is extended to a discrete Morse function $F_w$ which imposes a total ordering on all cells of the domain $K$. The gradient field is interpreted as a directed acyclic graph and gradient paths as traversals on this graph. The gradient field decomposes into a union of trees and the graph traversal is performed in parallel as an iterative search procedure towards the root of the trees.

3.1. Weighted discrete function

The weighted discrete function $F_w$ is equal to $f$ at vertices. The value of $F_w$ on a d-dimensional cell $α^d$ is recursively determined by its facet with highest function value, while ensuring that the value of $F_w$ at two different cells are never equal. Specifically,

$$F_w(α^d) = F_w(G_0(α^d)) + ε^d \times F_w(G_1(α^d)),$$

where $ε$ is an infinitesimally small positive real number,

$$G_0(α^d) = \arg \max_{γ < α^d} F_w(γ),$$

$$G_1(α^d) = \arg \min_{γ < α^d} F_w(γ).$$

$G_0(α)$ represents the highest facet of $α^d$ and $G_1(α)$ represents the highest face of $α$ that is disjoint from $G_0(α)$, $V_{G_0(α)}$ is the vertex set of $G_0(α^d)$, and $G_{max}$ denotes the value of the argument $γ$ that maximizes the function. $F_w$ is equal to $f$ at mesh vertices, $F_w$ ensures that when two cells share their highest facet, then the tie is broken using their respective second maximum face whose vertex sets are disjoint from the common facet. Figure 4 shows the definition of the weighted discrete function for some common cell types. Subsequent
algorithms require only the order on the cells induced by $F_w$. This order may be computed in terms of $f$ instead of explicitly computing $F_w$, assuming that the function values at the vertices are totally ordered. We use simulation of simplicity [EM90] to handle datasets that do not satisfy this requirement. In the following discussion we will use the analogy of height while referring to the value of $F_w$ at a cell.

The $\varepsilon$-lower-star of a $d$-cell $\alpha$ is defined as the set of cells, including $\alpha$, that have $\alpha$ as their highest $d$-face. As shown in [SMN11], the cells in the $\varepsilon$-lower-star of a cell appear contiguously in the ordering induced by $F_w$.

### 3.2. Computing gradient pairs

All cells are critical with respect to $F_w$. Algorithm 1 uses the ordering based on the weighted discrete function defined above to determine the gradient pairs of a simpler discrete Morse function that contains fewer critical cells. The gradient pairs are unique and independent of the order in which cells are processed. Algorithm 1 processes all cells $\alpha \in K$ individually to compute gradient pairs. The cell $\alpha$ belongs to the complex $K$, and $\beta$ is a cofacet of $\alpha$. The set $P_\beta$ is the collection of cofacets, $\beta$, of $\alpha$ such that $\alpha$ is the highest facet of $\beta$, i.e., $\alpha = G_0(\beta)$. The cell $\alpha$ is paired with the lowest cell in $P_\alpha$.

**Algorithm 1** \texttt{ASSIGNGRADIENT} (Cell complex $K$)

1: for all $\alpha \in K$ do  
2: $P_\alpha = \{\beta | \alpha \prec \beta$ and $\alpha = G_0(\beta)\}$  
3: if $P_\alpha \neq \emptyset$ then  
4: $\hat{\beta} = \min_{F_w}(P_\alpha)$  
5: pair_cells ($\alpha$, $\hat{\beta}$)

### 3.3. Computing the 2D MS complex

Once the discrete gradient field is computed, the descending / ascending manifolds and hence the combinatorial MS complex are extracted as a closure of the set of gradient paths that originate at a critical cell / closure in the dual of the set of inverted gradient paths that originate at a critical cell. These are computed using a breadth first traversal of gradient paths. A combinatorial connection between any two critical cells is established if there is a gradient path that connects them. For multi-core environments, multiple BFS traversals are launched, one from each critical point. The gradient paths of a 2D domain decomposes to a union of trees [CCL03]. In massively parallel environments, each cell iteratively queries its parent for its parent, to locate the root.

### 4. 3D MS Complex computation

In this section, we discuss how the algorithm discussed in Section 3 is extended to work for three dimensional scalar functions. In particular, Algorithm 1 is conservative and may leave multiple cells unpaired thereby declaring them critical (See Figure 5). We introduce an additional pairing procedure that avoids the creation of such $\varepsilon$-persistent pairs within the $\varepsilon$-lower star. This procedure executes during a second pass over the cells and essentially seeks to pair cells with their second highest facet consistently when their highest facet is paired with another cell. Next, we describe a traversal algorithm to compute the incidence of 2-saddles on 1-saddles. This algorithm is adapted to determine the ascending/descending manifolds of 2-saddles and 1-saddles. Finally, the descending manifolds of maxima and ascending manifolds of minima are computed using an iterative search procedure marching towards the root of the tree.

**Algorithm 2** \texttt{ASSIGNGRADIENT2} (Cell complex $K$)

1: for all $\alpha \in K \setminus K(0)$ do  
2: if $\alpha$ not paired by Algorithm 1 then  
3: $P_\alpha = \{\beta | \alpha$ is the second highest facet of $\beta\}$  
4: if $P_\alpha \neq \emptyset$ then  
5: $\hat{\beta} = \min_{F_w}(P_\alpha)$  
6: if $\hat{\beta}$ not paired by Algorithm 1 then  
7: pair_cells ($\alpha$, $\hat{\beta}$)

Figure 5: (left) The $\varepsilon$-lower-star and the lower link of a vertex with the height function defined. (center) Algorithm 1 declares the green edge and the red triangle as critical because the edge is not the highest facet of any of its cofacets. (right) Algorithm 2 pairs these cells because the edge is the second highest facet of the unpaired triangle.
Independence and Correctness. We first state and prove a Lemma to show that the pairs determined by Algorithm 2 are unique and hence the algorithm can be parallelized. The following Lemma states that if a cell is paired by Algorithm 2, then the pairing is unique, i.e., the cell is either paired with one of its cofacets or with its second highest facet, independent of the order in which cells are paired.

**Extended Order Independent Pairing Lemma.** If Algorithm 2 pairs a cell \( \alpha \) with its second-highest facet \( \beta \), then it will not pair \( \alpha \) with its second-highest facet \( \gamma \).

**Proof.** Consider the incidence relationships shown in Figure 6a between a \( d \)-cell \( \beta \), its highest facet \( \alpha' \), second highest facet \( \alpha \), highest \((d-2)\) facet \( \gamma' \), highest \((d-3)\) facet \( \psi' \), and \( \alpha \)'s second highest facet \( \gamma \). Since the input cell complex \( K \) is regular, there exists exactly two facets of \( \beta \), say \( \alpha_1 \) and \( \alpha_2 \), incident on \( \gamma' \). Further \( \gamma' \) is the highest facet of \( \alpha_1 \) and \( \alpha_2 \), which in turn implies that any third facet of \( \beta \) does not contain \( \gamma' \) and is hence lower than both \( \alpha_1 \) and \( \alpha_2 \). So, \( \alpha_1 \) and \( \alpha_2 \) are the highest and second highest facets of \( \beta \), namely \( \alpha' \) and \( \alpha \). A similar argument on \( \alpha \) and its highest \((d-3)\) face \( \psi' \) shows that \( \gamma \) and \( \gamma' \) are incident on \( \psi' \) and \( \psi \) is their highest facet.

We will prove the existence of a facet, \( \alpha'' \), of \( \beta \) that contains \( \gamma \) as its highest or second highest facet. In either case, \( \gamma \) will be paired with a cell different from \( \alpha \). If \( \gamma \) is the highest facet of \( \alpha'' \) then Algorithm 1 would have paired it with a cell different from \( \alpha \) because \( \alpha \) remains unpaired until it is processed by Algorithm 2. If \( \gamma \) is the second highest facet of \( \alpha'' \), then Algorithm 2 will seek to pair it with the lowest cofacet in \( P_\gamma \). The cell \( \alpha'' \) is lower than \( \alpha \) and belongs to \( P_\gamma \). So \( \alpha \) will not be paired with \( \gamma \).

We now show the existence of the cell \( \alpha'' \). Consider the \((d-2)\) cell \( \gamma \) as a face of \( \beta \). Since the input is a regular cell complex, there exists exactly two facets of \( \beta \) incident on \( \gamma \). The cell \( \alpha \) is one such facet of \( \beta \). Let \( \alpha'' \) be the other. The regularity of the input cell complex also implies the existence of exactly two facets, \( \gamma \) and \( \gamma'' \), that are incident on \( \psi' \). Further, \( \psi' \) is the highest \((d-3)\) face of \( \alpha'' \). It follows that \( \gamma \) is either the highest facet or second highest of \( \alpha'' \) using the same argument as above to show that \( \alpha \) and \( \alpha'' \) are incident on \( \gamma' \).

We now prove that Algorithm 2 produces a valid gradient field. A pairing between a cell \( \alpha \) and its cofacet \( \beta \) is valid if there exists a corresponding discrete Morse function. Such a function can be realized via a perturbation if none of the cofacets of \( \alpha \) are lower than \( \beta \). Let \( \beta \) be a cofacet of \( \alpha \) different from \( \beta \). The cell \( \alpha \) is clearly not the highest facet of \( \beta \) since it remains unpaired after being processed by Algorithm 1. If \( \alpha \) is the second highest facet of \( \beta \), then \( \beta \) is higher than \( \beta \) because it was not selected by Algorithm 1. If \( \alpha \) is neither the highest nor the second highest facet of \( \beta \), then the highest facet of \( \alpha \), say \( \gamma' \), is not the highest \((d-2)\) face of \( \beta \). This follows from the converse of the argument used in the above proof to show that \( \alpha' \) and \( \alpha \), the highest and second highest facets of \( \beta \), are incident on \( \gamma' \). Cells in the \( \varepsilon \)-lower-star of \( \gamma' \) appear contiguously in the ordering induced by \( F_\varepsilon \). Since \( \beta \) does not lie in the \( \varepsilon \)-lower-star of \( \gamma' \) and \( \beta \) does, it follows that \( \beta \) is not lower than \( \beta \).

4.2. **Saddle connection Algorithm**

The sub-structure of the gradient field consisting of 1-saddles, 2-saddles and \((1,2)\) gradient paths between them can be very intricate. This is because 1-saddle-2-saddle gradient paths in a three-dimensional domain may both split and merge. Figure 7 depicts the sub-structure of a gradient field that originates from a 2-saddle, splits and merges twice, before reaching a 1-saddle.

We trace the \((1,2)\) gradient paths by interpreting the sub-structure as a directed acyclic graph (DAG) induced by them. The number of paths between a 2-saddle and 1-saddle may be counted as the number of paths between 2-saddles and 1-saddles nodes in this DAG. We do not employ the standard breadth first search algorithm to traverse the graph because
this would necessitate the use of an array of flags to maintain if every cell is visited or not. Parallelizing the traversal will require a buffer, whose size equals that of the input, for each thread. This approach is clearly not scalable. We note that the number of critical cells reachable from a given cell tends to be small. Algorithm 3 describes a priority-queue based traversal method to determine the paths between 2-saddles and 1-saddles. The algorithm computes the number of gradient paths from a given 2-saddle to all (1,2) gradient pairs and 1-saddles that are reachable from it.

The algorithm begins by first initializing a priority queue, \( PQ \), that can contain 2-cells and 1-cells of \( K \). The priority queue is ordered based on the simpler discrete Morse function computed by the gradient pairing algorithms. We associate with each element \( \alpha \) of \( PQ \) the number of paths that arrive from \( \sigma \). \( PQ \) is initialized with the pair \( ( \sigma, 1 ) \). The algorithm pops the first cell \( \alpha \) from \( PQ \). It is possible for copies of the same cell to be entered into \( PQ \). Since all these cells have the same priority, \( PQ \) is repeatedly popped until all copies of \( \alpha \) are removed and the number of paths \( npaths \) that reach \( \alpha \) are summed over all copies. If \( \alpha \) is a critical 1-cell, then an arc with \( npaths \) multiplicity is inserted between \( \sigma \) and \( \alpha \). If \( \alpha \) is a 2-cell, then all the 1-saddles and 2-cells of (1,2) pairs incident on the boundary of \( \alpha \) (other than itself) are inserted into \( PQ \). The newly inserted pairs/saddles lie on \( npaths \) number of paths from \( \sigma \) through \( \alpha \). Newly inserted cells are lower than \( \alpha \). So, \( \alpha \) never re-enters \( PQ \). A cell is inserted into \( PQ \) when processing one of its neighboring cells. So, the number of copies of the cell in \( PQ \) is upper bounded by the number of its neighbors.

Each cell enters \( PQ \) only a constant number of times. So, the complexity of the algorithm is \( n \log(n) \), where \( n \) is the number of 2-cell-1-cell pairs and 1-saddles. Descending manifolds of saddles are computed by modifying Algorithm 3 to save the cells popped out of the priority queue at each iteration of the main loop. Ascending manifolds of 1-saddles are computed by employing the same procedure after reversing the priority, and reversing the role of 1-cells and 2-cells.

5. Handling large data

The 3D MS complex algorithm is extended to handle large data that do not fit in memory using a split and merge technique similar to that described in [SMN11] for 2D scalar functions. The MS-complex is computed in a five-stage process:

- Split dataset
- Compute MS complex on sub-domains
- Merge sub-domains
- Traverse merge history
- Extract geometry

The splitting of the data in stage 1 and simplification in stages 2 and 3 are different from the method described in [SMN11]. We describe these steps below. The complete algorithm is described in the Appendix.

First, the dataset is split into sub-domains. The domain is subdivided successively along each axis till sub-domains of manageable size are obtained. Then, we compute the MS complex within each sub-domain using the algorithm discussed in Section 4. To eliminate noisy critical cells early, we perform critical cell pair cancellations within sub-domains directed by topological persistence [ELZ02]. To ensure combinatorial consistency of the MS complex after merging, we allow cancellation of a pair of critical cells only if (a) neither of the two cells belongs to a gradient pair that crosses a shared boundary, (b) neither of the two cells lie on a shared boundary, and (c) at most one of the two cells is connected to a gradient pair that crosses a shared boundary. Next, we merge the MS complexes of sub-domains in reverse order of the subdivision to create the combined MS complex, similar to [SMN11]. The validity and equivalence of the resulting MS complex is a consequence of the ORDER INDEPENDENT CANCELLATION Lemma [SMN11]. After each merge we again perform cancellations directed by persistence to remove low persistent pairs that may have been created. Merge history traversal and extracting geometry are described in the Appendix.

6. Implementation

We implemented the above discussed techniques to leverage both GPU computing, and multi-core CPU architectures. We used the OpenCL framework to implement the gradient algorithm discussed in Section 4.1 on the GPU. We implemented the Algorithm 3 to process individual 2-saddles in
parallel on the CPU. We use the Boost threading [Boo] library to manage multiple threads. We now discuss various implementation and optimization issues with respect to the five stages.

**Gradient pairing and MS complex.** The gradient pairs and MS-complex are computed within each sub-domain. The gradient pairs are computed in two passes using Algorithm 1 and Algorithm 2. A three-dimensional array of bytes is used to store information per cell. Three bits are used to represent the six possible directions of the pair. Three bits are used to represent the six possible directions of the highest facet. One bit is used to mark cells as critical or not. The 3D array is created on the GPU and transferred to the CPU once the gradient computation is completed.

Next, the decomposition of the domain into descending/ascending manifolds of maxima/minima are computed on the GPU. This is done similar to the algorithm described in Section 3. Simultaneously, the CPU executes Algorithm 3 to determine connections between 2-saddles and 1-saddles. This step is optimized to traverse only those paths that reach 1-saddles by executing a single breadth first search traversal that begins from all 1-saddles and marking all (1,2) pairs that are reached when traversing the gradient field upwards. While processing paths that descend from the 2-saddles, only pairs that are marked reachable from a 1-saddle are inserted into the priority queue $PQ$. One bit of the 3D array, which records per cell information, is used to represent visited or not visited state of a cell.

The Morse-Smale complex is represented as a graph with nodes as critical cells. Adjacencies are represented as list of associative arrays, one for each critical point and the multiplicity of paths associated with each adjacency. Hence the complexity to access a particular adjacency of critical cell is $\log(n)$, $n$ being the maximum number of adjacent critical cells.

Merging proceeds by cancellation of gradient pairs that cross a shared boundary. So, the descending connections of lower index critical cell of the pair and the ascending connections of the higher index critical cell are discarded by the cancellation. We further optimize by not recording such connections in the MS complex and not launching Algorithm 3 from 2-saddles that are paired with maxima.

**Merging sub-domains.** The merging procedure proceeds in the reverse order of subdivision of the domain. Two sub-domains are merged into a single MS complex while ensuring that gradient pairs that cross the shared boundary are identified. Then these pairs are canceled out. This procedure ensures that there is no duplication of critical cells (and their combinatorial connections) that lie on the shared boundary. Similar to the previous step, a persistence based simplification is performed on the combined MS complex after each merge.

**Traversing merge history and extracting geometry.** The traversal of merge histories computes paths from critical cells that enter a sub-domain through a shared boundary. Since we simultaneously perform simplification during merging, we consider critical cells within a sub-domain that are paired with other critical cells, possibly outside the sub-domain, as entry points of gradient flow from surviving critical cells.

For extracting geometry, we require the original gradient field information. To obtain this, we reload the original function and recompute the gradient field, since the time taken to store and load this information is significantly higher than re-computing it. The partition of the domain based on gradient paths from extrema is computed on the GPU and the ascending/descending manifolds of saddles are computed using the modified version of Algorithm 3.

7. Experiments

We performed experiments on two different classes of datasets. First, we evaluated our algorithm with synthetic datasets to analyze its efficiency and scaling behavior with varying parameters such as regions of near flat-gradient and large numbers or gradients crossing shared boundaries. Second, we evaluated our algorithm’s performance on various volume datasets available from [http://www.volvis.org](http://www.volvis.org) and a dataset obtained from the simulation of a 3D Taylor-Green vortex flow on a Cartesian grid. All experiments were performed on an Intel-Xeon 2 GHz CPU with 4 cores and 16 GB of RAM and an Nvidia GeForce GTX 460 GPU with 336 cores and 1GB of memory. Data was split into $256 \times 256 \times 256$ sized sub-domains that fit in memory.

**Synthetic data.** We use a synthetic dataset $WGAUSS$ to stress test the algorithm. The function is defined as the product of a cosine wave with the 3D Gaussian $i.e. f(x,y,z) = \cos(2\pi \nu d_c) \times G_{c,\sigma}(x,y,z)$, where $\nu$ is the frequency of the cosine wave, $d_c$ is the distance of the point from the center, $G_{c,\sigma}$ is the 3D Gaussian centered at $c$ with variance $\sigma$, and the domain is the unit cube. The function is sampled at various grid resolutions to study scalability. This dataset is challenging since it contains multiple flat-regions at concentric spheres, where the cosine wave achieves its maximum or minimum.

We use two variants of this function to study our algorithm. First, we place the Gaussian at the center of the domain. Two sub-domains are merged into a single MS complex while ensuring that gradient pairs that cross the shared boundary are identified. Then these pairs are canceled out. This procedure ensures that there is no duplication of critical cells (and their combinatorial connections) that lie on the shared boundary. Similar to the previous step, a persistence based simplification is performed on the combined MS complex after each merge.

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Figure 8: (left) Scaling behavior with varying sizes of the WGAUSS dataset simplified using a threshold of 1%. (right) Computation times for stages 2-5 for the $1024^3$ datasets. The persistence threshold for each run is shown above the barplots. Time taken to split the data into sub-domains is approximately 3 minutes.

<table>
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<th>Dataset</th>
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<th>#crits.</th>
<th>time (a)</th>
<th>time (b)</th>
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<td>3s</td>
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</table>

Table 1: Timings for datasets available from volvis.org compared with timings to compute the MS complex as reported in (a) [GRWH11] and (b) [GBPH08].

Figure 8 shows the computation time for large data sizes. As can be seen from the figures, our algorithm performs better on the WGAUSS_MULTI dataset which contains fewer flat regions. The time to cancel the gradient pairs that cross common boundaries is lesser than the time taken to perform a persistence based simplification of the MS complex. This is reflected in the time taken to merge the sub-domains of the WGAUSS dataset. Figure 8 also plots running time for increasing data sizes of the WGAUSS dataset. The scaling results are similar for various values of persistence threshold. We note that the curves deviate away from linear scaling with increasing data sizes. On detailed analysis, we observed that stages two and five scale linearly whereas stages three and four did not. This is because of the representation of the combinatorial MS complex as a list of associative arrays.

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8. Conclusions

We have described a parallel algorithm to compute MS complexes of three-dimensional scalar functions. A hybrid multicore implementation results in significant speedup and superior performance as compared to existing approaches. Further, the algorithm is also memory efficient. In future, we plan to extend the algorithm to handle unstructured meshes.

Performance. To verify the benefits of parallelization, we compare our algorithm against existing methods [GBPH08, GRWH11] on datasets available from volvis.org, and a vortex flow data set, see Table 1. The experimental results indicate orders of magnitude improvement in the running time. Further, the memory required by our algorithm is less than 2GB even for the larger WGAUSS dataset. Gyulassy et al. and Günther et al. [GBPH08, GRWH11] report 23h and 5h, respectively, to process data sizes close to $1024^3$. Figure 9 shows the critical 2- and 3-cells along with the ascending manifolds of 2-saddles for various datasets as well as slice visualizations of the WGAUSS dataset along with the decomposition of one sub-domain into descending / ascending manifolds of maxima / minima.
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