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Approximating Lower-Star Persistence via 2D Combinatorial Map Simplification

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Abstract

Filtration simplification consists of simplifying a given filtration while simultaneously controlling the perturbation in the associated persistence diagrams. In this paper, we propose a filtration simplification algorithm for orientable 2-dimensional (2D) manifolds with or without boundary (meshes) represented by 2D combinatorial maps. Given a lower-star filtration of the mesh, faces are added into contiguous clusters according to a “height” function and a parameter $\epsilon$. Faces in the same cluster are merged into a single face, resulting in a lower resolution mesh and a simpler filtration. We prove that the parameter $\epsilon$ bounds the perturbation in the original persistence diagrams, and we provide experiments demonstrating the computational advantages of the simplification process.

Keywords: Persistent homology computation; 2D combinatorial map; mesh simplification.

1. Introduction

Topological data analysis (TDA) is a relatively new subfield of computer science. One of the most useful tools in TDA is persistent homology, which is an algebraic method for associating topological features with discrete data. In particular, persistent homology requires two crucial components: (1) a cell complex denoted $K$ to structure the data; and (2) a filtration, or a nested sequence of indexed subcomplexes where the initial complex is the empty complex and the terminating complex is $K$. See Dey et al. (1999); Bern et al. (1999); Carlsson (1999) for initial reports and Edelsbrunner and Harer (2010) for a modern exposition of the field.

Much work has been done to simplify complexes and filtrations in order to quickly compute persistence. In Günther et al. (2012), the authors proposed an efficient algorithm that computes persistent homology for 3D gray-scale images by first obtaining the Morse-Smale complex, which is much smaller than the input complex $K$ but contains all necessary information. The authors first computed a combinatorial gradient vector field using an algorithm from Robins et al. (2011), which induces a Morse-Smale complex. The persistence given by this new complex is then exactly the same as that given by the input complex. More recently, in Boissonnat et al. (2018), the authors first simplify a filtration of an arbitrary simplicial complex with “strong collapses”. The persistence associated with the reduced filtration is then the same as the original one.

Unlike the previous approaches, we approximate the persistence given by a filtration associated with some 2D mesh (i.e. a piecewise-linear orientable 2D manifold with or without boundary) within a user specified tolerance $\epsilon$. This is accomplished by merging faces in the mesh that meet a criterion. Other authors have taken similar approaches. In Botnan and Spreemann (2015), the authors gave two different approaches for approximating Čech persistence by building an approximation to the Čech filtration. In Dey and Slechta (2018), the authors developed a notion of $(p, \epsilon)$-admissible, where upon contracting a $(p, \epsilon)$-admissible edge in some complex $K$, the difference in $p$-dimensional persistence diagrams given by the initial and the simplified filtration is bounded above by $\epsilon$. We attain a similar goal: upon merging clusters of faces which meet our criterion, called $\epsilon$-permissible, the difference in the 0-, 1-, and 2-dimensional persistence diagrams given by the lower-star filtration on the initial and “merged” meshes is bounded by $\epsilon$.

In Damiand and Gonzalez-Diaz (2016), the authors proposed an efficient algorithm for computing the homology of meshes represented by 2D combinatorial maps, thereby avoiding the time-consuming step of constructing and modifying boundaries and coboundaries of cells. The process consists of merging faces if they share a common edge, guaranteeing that the structure of the combinatorial map and the homology of the mesh is preserved throughout the process.

This paper extends the work in Damiand and Gonzalez-Diaz (2016) and Damiand and Gonzalez-Diaz (2019) by giving an algorithm to approximate the lower-star persistent homology of meshes within a specified tolerance $\epsilon$. First, faces are grouped into clusters according to the parameter $\epsilon$. Faces in the same cluster are subsequently merged. In Damiand and Gonzalez-Diaz (2019), cluster membership was determined by a distance between faces, and there was no theoretical guarantee that persistence diagrams corresponding to the initial and simplified filtrations were “close.” In this paper, membership is determined by the faces’ “height”, and as our main contribution, we provide such a guarantee in Section 4. Experiments in Section 5 demon-
strate the utility of our approach, and we conclude with a brief discussion of possible directions for future work in Section 6.

2. Preliminary Notions

In this section we review combinatorial maps and persistent homology.

2.1. 2D combinatorial maps

Roughly speaking, a 2D combinatorial map (Lienhardt (1994); Damiand and Lienhardt (2014)), called a 2-map, is a representation of a mesh (see Fig. 1). A mesh \( M \) is a 3-tuple \((V, E, F)\) where \( V \) is a set of vertices or 0-cells, \( E \) is a set of edges or 1-cells, and \( F \) is a set of faces or 2-cells.

Cells are in relation together. Two cells are said to be incident if one cell belongs to the boundary of the other. Two i-cells are adjacent if there exists a \((i-1)\)-cell incident to both. Two i-cells are neighbors if they are both in the boundary of an \((i+1)\)-cell.

Some cells with specific properties are identified. An edge is dangling if it is incident to a vertex which is incident to no other edge. An edge is isolated if it has no adjacent edge. An edge is inner if it is incident to two different faces. An edge is a border if it is contained in the boundary of the mesh.

A 2D combinatorial map (or simply, a 2-map) is a 3-tuple \((D, \beta_1, \beta_2)\) where \( D \) is a finite set of darts and \( \beta_1 \) and \( \beta_2 \) are one-to-one mappings \( D \) onto \( E \) and \( F \), respectively.

Meshes can be represented by 2-maps as follows. A dart \( d \) represents an orientation of an edge; \( \beta_1(d) \) is the dart following \( d \) and belonging to the same face as \( d \); \( \beta_2(d) \) denotes the opposite orientation of \( d \). In Fig. 1(b), darts 8 and 11 represent the two possible orientations of a single edge, so \( \beta_2(8) = 11 \) and \( \beta_2(11) = 8 \).

A border dart belongs to exactly one vertex, one edge, and one face, and thus each cell of the mesh is described by a set of darts in the 2-map. For example, in Fig. 1(b), vertex \( v_1 \) is described by the set of darts \( \{2, 5, 8, 12\} \). Even isolated edges (like \( e_7 \) in Fig. 1(a)) belong to a degenerate face (which is why there are 5 faces in Fig. 1(a)). Inner, dangling, and isolated edges are described always by two darts \( d_1, d_2 \). They can be detected in a 2-map thanks to particular configurations of darts and \( \beta \) links (for example an edge \( \{d_1, d_2\} \) is isolated if \( \beta_1(\beta_1(d_1)) = d_1 \)).

A border edge is described by one dart \( d \) (like dart 16 in Fig. 1(b)). In such a case, \( \beta_2(d) = \emptyset \) and \( \beta_2 \) is a partial bijection.

2.2. Persistent Homology

In this subsection we review persistent homology, an important tool used to assign structure to discrete data. We assume that the reader has some knowledge of homology (for a reference, see Hatcher (2002)). For additional information on persistent homology, we encourage the reader to consult Edelsbrunner and Harer (2010).

Persistent homology captures the topological changes occurring in a growing sequence of meshes \( M_1 \subseteq M_2 \subseteq \ldots \subseteq M_\alpha = M \), called a filtration. As one progresses through the filtration, homology classes of different dimensions may appear (be born) and disappear (die). Some homology classes may be present in \( M_\alpha \) as \( \alpha \to \infty \). Such classes give the homology of \( M \).

Filtrations are frequently constructed on a mesh \( M \) via a real-valued function \( h \) on the vertices of \( M \). One such example is the lower-star filtration \( \{M_\tau\}_{\tau \in \mathbb{R}} \) where a cell \( \sigma \) is in \( M_\tau \) if for all vertices \( v \) which are incident to \( \sigma \), \( h(v) \leq \tau \). Intuitively, this filtration corresponds to taking sublevel sets of \( M \) under \( h \).

If a homology class is born at \( M \) and dies entering \( M_\tau \), then \( \tau - \tau' \) is the persistence of the homology class. If it is born at \( M \) but never dies then its persistence is set to infinity. Intuitively, homology classes with low persistence are topological noise and the ones that persist correspond to features of the mesh.

The information obtained when computing persistent homology of a filtration can be summarized by a set of persistence diagrams, each of which is a multi-set of (birth, death) pairs in the extended real plane, where each pair represents the birth and death of a homology class. For a filtration \( M := \{M_\tau\}_{\tau \in \mathbb{R}} \), there is a persistence diagram for each dimension \( p \) denoted \( \text{Dgm}_p(M) \). As expected, \( \text{Dgm}_p(M) \) contains persistence information for \( p \)-dimensional homology classes. In addition, all points on the diagonal given by \( y = x \) are considered to be in each persistence diagram with infinite multiplicity. This enables comparing diagrams with a different number of homology classes with positive persistence. One of the most popular tools for comparing persistence diagrams is the bottleneck distance.

**Definition 1.** The bottleneck distance between two persistence diagrams \( \text{Dgm}_p(A) \) and \( \text{Dgm}_p(B) \) is

\[
d_b(\text{Dgm}_p(A), \text{Dgm}_p(B)) := \inf_{\psi \in \Psi} \sup_{a \in \text{Dgm}_p(A)} \|a - \psi(a)\|_\infty
\]

where \( \Psi \) is the set of bijections from \( \text{Dgm}_p(A) \) to \( \text{Dgm}_p(B) \).

From now on, by abuse of notation, we use \( M \) to denote a mesh, the 2-map representing the mesh, or the filtration considered on the mesh. In this paper, persistence diagrams are computed via Algorithm 1, which is a modification of the incremental algorithm given in Gonzalez-Diaz and Real (2005). Given a mesh and an ordering of its cells, Algorithm 1 computes a 3-tuple \((M, H, f)\) where \( M \) is the given mesh represented by a 2-map; \( H \) is the set of surviving cells; and \( f \) is a map from the \( k \)-cells in \( M \) to a sum of surviving cells. For a \( k \)-cell \( \sigma \), \( \partial(\sigma) \) denotes the set of \((k-1)\)-cells in its boundary. Algorithm 1 guarantees that the set of all the surviving \( k \)-cells (for a fixed \( k \)), together with the disjoint union of sets operation \( + \), forms the group \( C_k(H) \) which is isomorphic to \( H_k(M) \). In addition,
Algorithm 1: Computing persistent homology (Algorithm 2 of Gonzalez-Diaz and Real (2005)).

**Input:** An ordering of the cells of \( M: \{\sigma_1, \ldots, \sigma_m\} \).

**Output:** Persistent homology of \( M \) with respect to such ordering.

Initialize \( H \leftarrow \emptyset \) and \( f(\sigma_i) \leftarrow 0 \), for \( 1 \leq i \leq m \).

for \( i = 1 \) to \( m \) do
  if \( f(\partial(\sigma_i)) = 0 \) then
    \( f(\sigma_i) \leftarrow \sigma_i \);
    \( H \leftarrow H \cup \{\sigma_i\} \) (a new homology class was born);
  else
    Let \( \sigma_j \in f(\partial(\sigma_i)) \), \( j = \max\{\text{index}(\mu) : \mu \in f(\partial(\sigma_i))\} \);
    \( H \leftarrow H \cup \{\sigma_i\} \) (an old homology class died);
  endif
  foreach \( x \in M \) such that \( \sigma_j \in f(x) \) do
    \( f(x) \leftarrow f(x) + f(\partial(\sigma_i)) \).
  endfor
endfor

We pay particular attention to step 1. In this step, the 2-map is simplified by dispatching the faces into clusters subject to some constraints according to the parameter \( \epsilon \). Note that since \( h \) is only defined on the vertex set of \( M \), the index at which a face enters the lower-star filtration is precisely the maximum height value of a constituent vertex. Hence, for face \( \sigma \), we use the notation \( \max(\partial(\sigma)) := \max\{h(v) \mid v \text{ is a vertex of } \sigma\} \) to denote the height, while \( \min(\partial(\sigma)) \) is defined as expected. We will ensure that all clusters of size greater than 1 are \( \epsilon \)-permissible.

**Definition 2.** A cluster of faces \( C = \{\sigma_1, \ldots, \sigma_m\} \) is \( \epsilon \)-permissible if \( \max(C) - \min(C) \leq \epsilon \) where \( \max(C) = \max_{\sigma \in C}(\max(\partial(\sigma))) \) (analogously for \( \min \)).

Algorithm 2: Finding \( \epsilon \)-permissible clusters

**Input:** A 2-map \( M \); a parameter \( \epsilon \)

**Output:** A set of \( \epsilon \)-permissible clusters \( \Lambda = \{C_1, \ldots, C_n\} \).

\( k \leftarrow 1 \);

foreach face \( \sigma \) of \( M \) not yet assigned in a cluster do
  add a new cluster \( C_k \) to \( \Lambda \);
  add \( \sigma \) into \( C_k \);
  \( T \leftarrow \) all faces adjacent to \( \sigma \);
  while \( T \) is non empty do
    \( \tau \leftarrow \) one face in \( T \);
    remove \( \tau \) from \( T \);
    if \( \tau \) is not yet assigned in a cluster and
      \( \max(\max(h(\tau)), \max(\partial(C_2))) - \min(\min(h(\tau)), \min(\partial(C_2))) \leq \epsilon \) then
        add \( \tau \) into \( C_2 \);
        add all faces adjacent to \( \tau \) into \( T \);
    endif
  endwhile
  \( k \leftarrow k + 1 \);
endfor

We partition \( M \) into \( \epsilon \)-permissible clusters via Algorithm 2. The first face of each cluster is chosen arbitrarily, and the rest are added using neighborhood relations satisfying \( \epsilon \)-permissibility. If \( \sigma \) is not yet assigned into a cluster, we create a new cluster \( C_k \) and assign \( \sigma \) to \( C_k \). Then, while \( C_k \) is \( \epsilon \)-permissible, \( C_k \) is grown by progressively adding faces which are adjacent to the cluster. Each cluster is associated with a value corresponding to the minimum \( \min \) of all faces in the cluster and a value corresponding to the maximum \( \max \) of all faces in the cluster, which makes it easy to check if adding a new face would violate the \( \epsilon \)-permissible condition.

We repeat this process until each face is assigned into a cluster. After dispatching all of the faces to clusters, we apply Algorithm 3 which simplifies the 2-map by merging each cluster. The simplification process is depicted in Fig. 2. Critically, the result of merging faces via Algorithm 3 always results in a topological disk with a connected boundary. The complexity of Algorithm 3 is \( O(na(n)) \), where \( n \) being the number of darts in the 2-map and \( a(n) \) is the slow growing inverse Ackermann function (cf. Damiani et al. (2006); Damiani and Gonzalez-Diaz (2016) for more details). Note that since \( a(n) \leq 5 \) for all practical purposes, Algorithm 3 is functionally linear.

Once the mesh is simplified, the only remaining steps are
to compute the lower-star filtration of the simplified mesh, and then to compute the persistent homology using Algorithm 1.

The complexity of the whole process is \( O(n + O(m^2)) \), \( n \) being the number of darts of the given 2-map, and \( m \) being the number of cells in the simplified 2-map. This shows the interest of simplifying the 2-map before computing the persistent homology.

4. Persistence Diagram Stability

Let \( M \) be a polygonal mesh represented as a 2-map, and let \( M' \) denote the mesh obtained by merging clusters as explained in Section 3. In this section, we move to showing that merging \( \epsilon \)-permissible clusters bounds the perturbation in the persistence diagrams associated with the lower-star filtration. Throughout this section, we will use \( \text{Dgm}_p(M) \) to denote the \( p \)-dimensional persistence diagram given by the lower-star filtration of \( M \). We will also use the notation \( M_\alpha \) to denote \( \{ \sigma \in M \mid \max h(\sigma) \leq \alpha \} \). In the interest of simplicity, we abuse notation and do not distinguish between vertices in \( M \) and \( M' \).

If \( v \) is a vertex in \( M' \), then there is a natural corresponding vertex in \( M \). We will use \( v \) to refer to both of these vertices. Likewise for edges and faces. In addition, we will assume that there is a total order \( < \) on the simplices in \( M \) which respects the height function \( h \) and dimension. That is, for \( \sigma, \tau \in M \), if \( h(\sigma) < h(\tau) \), then \( \sigma < \tau \). Similarly, if \( h(\sigma) = h(\tau) \) and \( \dim(\sigma) < \dim(\tau) \), then \( \sigma < \tau \). Hence, for each set of simplices in \( M \), there exists a unique minimal simplex. This total order allows us to consider all those simplices in \( M \) which precede simplex \( \sigma \) under \( < \). We denote such a set as \( M_{<\sigma} \).

Lemma 1. Let \( u, v \in M \) be vertices which are also in \( M' \), where \( M' \) is obtained by merging a collection of \( \epsilon \)-permissible clusters. If \( u \) and \( v \) are in the same connected component in \( M_\alpha \), then \( u \) and \( v \) are in the same connected component in \( M'_{<\alpha} \).

Proof. Note that clusters are merged into a single face which is homeomorphic to a disk. Hence, if \( u \) and \( v \) are connected in \( M_\alpha \) by a path of edges which includes edges that are removed by cluster merging, then the segment of edges between two vertices on the boundary of a cluster can be replaced by instead traversing the boundary of the cluster. If \( w \) is a vertex removed by cluster merging and which is also the face of an edge that connected \( u \) and \( v \), then all edges \( e \) on the boundary of the cluster containing \( w \) must have height \( h(e) \leq h(w) + \epsilon \leq \alpha + \epsilon \). Hence, there is a path between \( u \) and \( v \) in \( M'_{<\alpha} \).

Theorem 1. If \( M' \) is obtained from \( M \) by merging a collection of \( \epsilon \)-permissible clusters, then \( d_0(\text{Dgm}_0(M), \text{Dgm}_0(M')) \leq \epsilon \).

Proof. We proceed by constructing a matching \( \gamma : \text{Dgm}_0(M) \rightarrow \text{Dgm}_0(M') \). Consider \((b, d) \in \text{Dgm}_0(M)\). The point \((b, d)\) corresponds to the lifetime of some homology class created by vertex \( v \in M \). We use the notation \([v]_M\) to refer to the set of vertices in \( M \) which are in the same connected component as \( v \) prior to the introduction of the edge which kills the homology class created by \( v \) at index \( d \) (if there is such an edge). If \( d = \infty \), then \([v]_M\) refers to those vertices in the same
connected component as $v$ in $M$. For $v' \in M'$, we define $[v']_{M'}$ in an analogous way.

In order to define $\gamma$, we consider two cases. First, we assume that $d$ is finite. In this case, the homology class $[v]_M$ dies when an edge $e$ is introduced to $M_d$ such that the vertices $v$ and $u$ are in the same connected component, where $u < v$. We associate with $v$, $u$ a unique vertex in $M'$, denoted $v'$, $u'$. In particular, we let $v'$ denote the minimal vertex in $[v]_M \cap M'$ under $\prec$. If there is no such $v'$, then $[v]_M$ only includes vertices which are removed via face merging. Hence, $[v]_M$ dies when an edge $e = \{w_0, w_1\}$ joins the connected component containing $v$ with the one containing $u$. In particular, both $w_0$ and $w_1$ must be in the cluster containing $v$, as if $w_0$ is not in the cluster, then $w_0$ must be on the boundary of the cluster containing $v$ which would imply that $v'$ exists. Ergo, by $\epsilon$-permissibility, $h(v) = b \leq h(e) = d \leq h(v) + \epsilon$. Hence, we let $\gamma(b, d) = \left(\frac{d - b}{2}, \frac{h(v)}{2}\right)$, which satisfies $\|\gamma(b, d) - (b, d)\|_{\infty} = \max\left\{\left|\frac{d - b}{2} - b\right|, \left|\frac{h(v)}{2} - d\right|\right\} = \frac{d - b}{2} \leq \epsilon/2$.

We let $u'$ be the minimal element in the set $[u]_M \cap M \cap \infty \cap M'$. If $u'$ does not exist, then when $[u]_M$ merges with $[v]_M$, the connected component $[u]_M$ contains only vertices which are removed during cluster merging. By identical reasoning as in the case where $v'$ was undefined, the connected component $[v]_M$ must contain a vertex $w$ which is in the same cluster as $u$. Hence, as $[u]_M$ is joined to $[v]_M$ upon the introduction of an edge $e$ in $M_d$, where $e$ is in the cluster containing $u$, it follows that $h(u) \leq h(e) \leq h(u) + \epsilon$ by $\epsilon$-permissibility. Therefore, as $h(v) = h(e)$, it follows that $h(e) - h(v) = d - b \leq \epsilon$. Hence, we let $\gamma(b, d) = \left(\frac{d - b}{2}, \frac{h(v)}{2}\right)$, which we have already shown to satisfy $\|\gamma(b, d) - (b, d)\|_{\infty} \leq \epsilon/2$.

Hence, we assume that both $u'$ and $v'$ exist. By Lemma 1, the vertices $u'$, $v'$ are in the same connected component in $M'_{\tau}$, $d' \leq d + \epsilon$. It follows that $d \leq d'$, else $u$ and $v$ would be in the same connected component prior to $M_d$. Note that $u'$, $v'$ satisfy $h(u) \leq h(u') \leq h(u) + \epsilon$ and $h(v) \leq h(v') \leq h(v) + \epsilon$. This is because if $u$ is not removed, then $u' = u$ and $h(u) = h(u')$. If $u$ is removed and $u'$ exists, then the connected component containing $u$ prior to joining with $v$ must contain some vertex $w$ on the boundary of the merged cluster, which satisfies $h(u) \leq h(w) \leq h(u) + \epsilon$, and since $h(u) \leq h(u') \leq h(w)$, it follows that $h(u) \leq h(u') \leq h(u) + \epsilon$. The same reasoning applies to $v$, $v'$.

Now, we consider two subcases. First, we assume that the class $[v']_{M'}$ merges into some other class $[w]_{M'}$ in subcomplex $M'_D$ where $d \leq D \leq d' \leq d + \epsilon$. In such a case, we let $\gamma(b, d) = (h(v'), D)$. Since $b \leq h(v') \leq b + \epsilon$, and $d \leq D \leq d + \epsilon$, it follows that $\|\gamma(b, d) - (b, d)\|_{\infty} \leq \epsilon$. Second, we assume that there exists no such $D$. However, in $M'_D$, $v'$ and $u'$ are in the same connected component, so it follows that $[u']_{M'}$ merges into $[v']_{M'}$. Since by assumption $h(u) \leq h(v)$ and $h(v') \leq h(u')$, and we have shown that $h(u) \leq h(u') \leq h(u) + \epsilon$, it follows that $h(u') \leq h(v') \leq h(u') \leq h(u) + \epsilon$, which implies that $0 \leq h(u') - h(v') \leq \epsilon$. Hence, we let $\gamma(b, d) = (h(u'), d')$ which satisfies $\|\gamma(b, d) - (b, d)\|_{\infty} \leq \epsilon$. These subcases are illustrated in Fig. 3.

In the second case, we assume $d = \infty$. In $M'$, it follows that $v'$ creates a class $[v']_{M'}$ with lifetime $(h(v'), \infty)$. We let $\gamma(b, d) = (h(v'), \infty)$. We have established that either $h(v') = h(v) + \epsilon$ or $0 \leq h(v') - h(v) \leq \epsilon$. If we have the former, then $\|\gamma(b, d) - (b, d)\|_{\infty} = 0$. In the event of the later, it follows that $\|\gamma(b, d) - (b, d)\|_{\infty} \leq \epsilon$.

We now prove that $\gamma$ is a matching. When $d$ is finite, if $\gamma(b_1, d_1) = \gamma(b_2, d_2) = (b', d')$, then $(b', d')$ has multiplicity at least 2. Otherwise, a single class was killed twice, which is a contradiction. In the case where $d$ is infinite, each connected component in $M$ corresponds to a unique connected component in $M'$, so $\gamma$ must be one-to-one.

To see that $\gamma$ is onto, first note that every point with infinite persistence is the image of a point with infinite persistence, as cluster merging does not destroy connected components. Second, all $(b', d') \in M'$, $d' < \infty$ correspond to the lifetime of some class created by vertex $v'$ denoted $[v']_{M'}$ which is killed when it merges with some class created by $u'$ denoted $[u']_{M'}$. Likewise, $[u']_{M'}$ and $[v']_{M'}$ must merge in $M$ at subcomplex $M_d$. Hence, if $[u']_{M'}$ is the class that dies when merging with $[v']_{M'}$, and $(b, d)$ corresponds to the lifetime of $[u']_{M'}$, then $\gamma(b, d) = (b', d')$. Similarly for if $[v']_{M'}$ dies. Hence, $\gamma$ is onto.

Thus, $\gamma$ is a matching where $\|\gamma(b, d) - (b, d)\|_{\infty} \leq \epsilon$ for all $(b, d) \in D_{gm}(M)$. Ergo, $d_\epsilon(D_{gm}(M), D_{gm}(M')) \leq \epsilon$.  

Although we could have attempted to give a proof using classical results from Cohen-Steiner et al. (2007); Chazal et al. (2016) related to stability of persistence diagrams, we preferred to provide a constructive proof giving the explicit matching between the points in the respective diagrams.

**Theorem 2.** If $M'$ is obtained by merging a collection of $\epsilon$-permissible clusters, then $d_\epsilon(D_{gm}(M), D_{gm}(M')) \leq \epsilon$.

**Proof.** Analogously to the proof of Theorem 1: if an edge $e$ creates a 1-cycle $\pi = \sum_i \epsilon_i$, then we use $[\pi]_M$ to refer to the cycle class containing $\pi$. As in the 0-dimensional case, $\pi'$ refers
to the representative cycle of \([\pi]_M\) that is created at the lowest height value and for which none of the edges are removed when clusters are merged. The matching \(\gamma\) is constructed in the same way as the 0-dimensional case: the points corresponding to those classes for which a representative cycle is contained entirely in a cluster and for which the persistence is less than \(\epsilon\) are paired with the diagonal, points corresponding to those cycles with lifetimes with infinite persistence are paired with canonical choice, and points with finite persistence which do not satisfy the first case are either paired with the diagonal or with the point corresponding to \([\pi']_M\), or, in the event that \([\pi]_M\) kills \([\rho']_M\) where \([\rho]_M\) kills \([\pi]_M\), then the point corresponding to \([\pi]_M\) is paired with the point corresponding to \([\rho']_M\).

The only substantive difference is that in the 0-dimensional case, if \([u]_M\) and \([v]_M\) merge in \(M_d\), then \([u]_M\) and \([v]_M\) merge at \(M'\) where \(d'\) satisfies \(d \leq d' \leq d + \epsilon\). In the 1-dimensional case, if \([\pi]_M\) and \([\rho]_M\) merge in \(M_d\), then \([\pi]_M\) and \([\rho']_M\) merge in \(M'\), where \(d - \epsilon \leq d' \leq d\), as merged faces take the maximum value of the remaining vertices. But this still permits \(\|\gamma(b, d) - (b, d)\|_\infty \leq \epsilon\). The function \(\gamma\) is then a matching by the same reasoning.

**Theorem 3.** If \(M'\) is obtained by merging a collection of permissible clusters, then \(d_0(Dgm_2(M), Dgm_2(M')) \leq \epsilon\).

**Proof.** This proof follows analogously to the proof of Theorem 1, except the only case that needs to be considered in when \(d = \infty\). Since cluster merging does not destroy 2-cycles or 2-classes, then for each \((b, d) \in Dgm_2(M)\) there is a canonical choice of a point \((b', d') \in Dgm_2(M')\), where \(b - \epsilon \leq b' \leq b\) and \(d = d' = \infty\). Hence, \(\|\gamma(b, d) - (b, d)\|_\infty \leq \epsilon\).

5. Experiments

We have implemented our algorithm for lower-star filtration simplification by using the CGAL implementation of combinatorial maps (see Damiand (2011)) and an additional layer, called linear cell complex, which represents the geometry. All experiments were run on an Intel® i7-4790 CPU, 4 cores @ 3.60GHz with 32 GB RAM. All the computation times given here are averages of 10 consecutive runs on the same mesh.

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<tr>
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<th>#0-cells</th>
<th>#1-cells</th>
<th>#2-cells</th>
<th>#H0</th>
<th>#H1</th>
<th>#H2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blade</td>
<td>882,954</td>
<td>2,648,082</td>
<td>1,765,388</td>
<td>295</td>
<td>330</td>
<td>295</td>
</tr>
<tr>
<td>DrumDancer</td>
<td>1,335,436</td>
<td>4,06,0302</td>
<td>2,670,868</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Neptune</td>
<td>2,003,932</td>
<td>6,011,808</td>
<td>4,007,872</td>
<td>1</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>HappyBuddha</td>
<td>543,652</td>
<td>1,631,574</td>
<td>1,087,716</td>
<td>1</td>
<td>208</td>
<td>1</td>
</tr>
<tr>
<td>Iphigenia</td>
<td>351,750</td>
<td>1,055,268</td>
<td>703,512</td>
<td>1</td>
<td>8</td>
<td>1</td>
</tr>
<tr>
<td>ThaiStatue</td>
<td>4,999,996</td>
<td>15,000,000</td>
<td>10,000,000</td>
<td>1</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

Figure 4: Example of one mesh used in our experiments: the HappyBuddha mesh. The table gives the number of \(i\)-cells, \(i\)-cells, for the six meshes used, and the number of \(Hi\) generators, \(Hi\), for \(i = 0, 1, 2\).

5.1. 3D Meshes

In our first experiment, we tested our algorithm on six meshes\(^1\), one of which is shown in Fig. 4. The sizes of the meshes range between 703,512 and 10,000,000 faces. All these meshes have only one connected component, except Blade which has 295 connected components because of many small isolated closed meshes within the blade.

Our experiments are performed with various values of \(\epsilon\). For each mesh, we computed the height of the mesh \(\alpha\) (i.e. the maximal height minus the minimal one), and let \(\epsilon = (\Phi \cdot \alpha)/100\), \(\Phi\) being a percentage \((0 \leq \Phi \leq 100)\). We start with \(\Phi = 0.32\), and ran five experiments where, with each additional experiment, \(\Phi\) is doubled. We also computed the persistence diagrams corresponding to the lower-star filtration induced by the height value on the vertices of the original mesh (without simplification). Note that as \(\Phi\) increases, \(\epsilon\) increases too. The average number of faces in a single cluster increases and thus the combinatorial map becomes more and more simplified. This tends to correspond to a rise in the bottleneck distance between the persistence diagrams corresponding to the simplified mesh and the original mesh, displayed in Table 1(a).

We can see an illustration of the effect of the \(\epsilon\) parameter on the size of the different clusters in Fig. 5. Data showing the number of cells in the simplified mesh as \(\Phi\) varies is displayed in Fig. 6(a). As expected, increasing \(\epsilon\) greatly affects mesh size.

The effect of \(\epsilon\) on the computation time is illustrated in Fig. 7(a), where our method for computing persistence based

\(^1\)http://www-graphics.stanford.edu/data/3Dscanrep/
https://www.cc.gatech.edu/projects/large_models/
Figure 6: Number of vertices, edges and faces of the simplified 2-maps (in log₂ scale) depending on the value of Φ. These numbers are average values for the six meshes in (a), and for the six images in (b).

(a)  
(b)  

Figure 7: Computation time (in seconds) of our method by using the lower-star filtration with increasing Φ values. For Φ = 0, lower-star filtration is used without simplification step. The graph shows average values for the six meshes in (a), and for the six images in (b), and details time spent in the different parts of the method: computation of the filtration, combinatorial map simplification and persistence computation by using AT-model. Filtration computes the height of each cell (vertices, edges and face) and dispatches faces into clusters according to Φ. Simplification is the 2-map simplification, and AT-model is the persistent homology computation, including sorting the cells.

(a)  
(b)  

Figure 8: (a) One image used in our second experiments. (b) Its representation as mesh, with one color per cluster, for ϵ = 160, (c) and for ϵ = 320. The table gives the size of the images (in pixels), the number of i-cells for the six meshes built from the images, #i-cells. The number of Hi generators is the same for all images, H0=1, H1=0 and H2=0.

<table>
<thead>
<tr>
<th></th>
<th>Size</th>
<th>#0-cells</th>
<th>#1-cells</th>
<th>#2-cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Im1</td>
<td>1834 × 2000</td>
<td>3,668,000</td>
<td>10,996,333</td>
<td>7,328,334</td>
</tr>
<tr>
<td>Im2</td>
<td>2200 × 1313</td>
<td>2,888,600</td>
<td>8,658,775</td>
<td>5,770,176</td>
</tr>
<tr>
<td>Im3</td>
<td>3000 × 2000</td>
<td>6,000,000</td>
<td>17,990,001</td>
<td>11,990,002</td>
</tr>
<tr>
<td>Im4</td>
<td>3456 × 5184</td>
<td>17,915,904</td>
<td>53,730,433</td>
<td>35,814,530</td>
</tr>
<tr>
<td>Im5</td>
<td>5616 × 3744</td>
<td>21,026,304</td>
<td>63,060,193</td>
<td>42,033,890</td>
</tr>
<tr>
<td>Im6</td>
<td>1280 × 853</td>
<td>1,091,840</td>
<td>3,271,255</td>
<td>2,179,416</td>
</tr>
</tbody>
</table>

(a) (b) (c)
on 2-map simplification is ran with different values of $\Phi$. As expected, computation time decreases while $\epsilon$ increases, as this corresponds to a simpler mesh and thus a simpler filtration. We can notice that for sufficiently large $\epsilon$, computation time does not decrease because all the time is taken by the computation of the filtration and the simplification step.

The bottleneck distances between the persistence diagrams corresponding to the lower-star filtration and the filtration obtained when varying $\epsilon$ are given in Table 1(a). As expected, the bottleneck distances are always smaller than $\epsilon$.

### 5.2. 2D Images

For our second experiment, we used six grayscale images from Unsplash (https://unsplash.com/), one depicted in Fig. 8, with size between 1280 $\times$ 853 and 5616 $\times$ 3744 pixels. For each image, we created a 2D mesh having one vertex per pixel, its height value being the intensity of the pixel. This transformation enables application of our algorithm for approximating the persistent homology of 2D meshes to grayscale images. The number of cells in the simplified meshes obtained from the images are given in Fig. 8.

We compared the persistent homology computation of the six meshes for increasing values of $\Phi$. The effect of $\epsilon$ on the number of cells is illustrated in Fig. 6(b).

Like for experiments with meshes, we can observe in Fig. 7(b) that computation time decreases while $\epsilon$ increases. We obtain here a better speed-up compared to the experiment with meshes. This comes from the numbers of cells which is much larger for images than for meshes. In this case, simplifying the 2-map greatly reduces the numbers of cells and thus the computation time spent in the AT-model computation.

Lastly, the bottleneck distances between the persistence diagrams corresponding to the lower-star filtration and the filtration obtained for the different $\Phi$ are given in Table 1(b). As expected, the bottleneck distances are always smaller than $\epsilon$.

### 6. Conclusion

We conclude with a brief discussion of future directions for research. First, we plan to extend our work to non-orientable manifolds by using the generalized maps package (the non-orientable extension of combinatorial maps) of CGAL. We also would like to define a parallel version of our method: the combinatorial map simplification was already defined in parallel in Damiand and Gonzalez-Diaz (2016) but we need now to study if it is possible to parallelize some parts of the AT-model computation algorithm. Extension in $n$D could be given based on the theoretical results for removal and contraction operations in any dimension given in Damiand et al. (2012). Indeed, all basic tools used in this work, combinatorial maps, removal / contraction operations and AT-model computation, are defined in any dimension.

### References


