TopoMap: A 0-dimensional Homology Preserving Projection of High-Dimensional Data

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Abstract— Multidimensional Projection is a fundamental tool for high-dimensional data analytics and visualization. With very few exceptions, projection techniques are designed to map data from a high-dimensional space to a visual space so as to preserve some dissimilarity (similarity) measure, such as the Euclidean distance for example. In fact, although adopting distinct mathematical formulations designed to favor different aspects of the data, most multidimensional projection methods strive to preserve dissimilarity measures that encapsulate geometric properties such as distances or the proximity relation between data objects. However, geometric relations are not the only interesting property to be preserved in a projection. For instance, the analysis of particular structures such as clusters and outliers could be more reliably performed if the mapping process gives some guarantee as to topological invariants such as connected components and loops. This paper introduces TopoMap, a novel projection technique which provides topological guarantees during the mapping process. In particular, the proposed method performs the mapping from a high-dimensional space to a visual space, while preserving the 0-dimensional persistence diagram of the Rips filtration of the high-dimensional data, ensuring that the filtrations generate the same connected components when applied to the original as well as projected data. The presented case studies show that the topological guarantee provided by TopoMap not only brings confidence to the visual analytic process but also can be used to assist in the assessment of other projection methods.

Index Terms—Topological data analysis, computational topology, high-dimensional data, projection.

1 INTRODUCTION

Multidimensional Scaling (MDS) accounts for the problem of embedding data in a Cartesian space while preserving intrinsic properties of the data. A particularly important task in the context of MDS is dimensionality reduction, which aims to map data from a d-dimensional to a k-dimensional Cartesian space where $k << d$. In the context of visualization, where the embedding space is 2D or 3D, MDS is typically called multidimensional projection (MDP).

Over the last decades, a multitude of MDP methods have been developed to map high-dimensional data to a visual space while preserving geometric properties such as the Euclidean distance between data objects. A main issue shared by all those methods is that the preservation of geometric properties can only be guaranteed under very particular conditions. Thus, errors and distortions are highly likely in the resulting mapping, introducing uncertainties to analytical procedures carried out from projection layouts [68]. For instance, structures observed in the point cloud resulting from a projection such as neighborhood relations might not be the ones existing in the original data, thus potentially leading inexperienced practitioners to wrong conclusions.

Although a number of alternatives have been proposed to render the analysis of projection layouts more reliable [54], few are focused on developing MDP methods with theoretical guarantees as to properties...
preserved by the mapping. Guaranteeing that a certain property is preserved exactly by the mapping makes the analytical process more reliable and meaningful, ensuring that what is seen is indeed what takes place in the high-dimensional space.

This work introduces TopoMap, a novel MDP technique that is guaranteed to preserve topological structures during the dimensionality reduction process. Specifically, TopoMap maps high-dimensional data to a visual space while preserving 0-homology (Betti-0) topological persistence as defined by a Rips filtration over the input data points. Intuitively, a Rips filtration grows a high dimensional ball around the data points, and adds an edge (or a high dimensional simplex) to the filtration when two (or more) balls intersect. In other words, the proposed method ensures that the topological filtrations over both the original as well as the projected data generate the same connected components at the same instances of the respective filtrations. The topological guarantee provided by TopoMap allows analysts to confidently explore high-dimensional data by visualizing which groups of objects are more tightly connected in the high-dimensional space. As we show in the provided case studies, visualizing persistent components via Betti-0 preserving projections enables an intuitive analytical process, making the identification of objects with similar properties an easier task. Moreover, in contrast to many distance-based (dissimilarity-based) projections, there is no uncertainty in the visual identification of groups (clusters) in the layout produced by TopoMap.

Besides enabling reliable mechanisms for data exploration, the proposed methodology can be used to assess and better understand distance-based projections. Since TopoMap is guaranteed to preserve the connected components of a particular neighborhood graph structure, one can rely on it to analyze how those connected components are mapped by other projection methods. As a result, one can further understand how distance-based MDP methods split or merge components, thus revealing regions with distortion.

In summary, the main contributions of the work are:

- A theoretical framework to support the design of a dimensionality reduction technique called TopoMap, which is guaranteed to preserve the Betti-0 topological persistence defined by the Rips filtration over the data.
- An optimization procedure that ensures the correct mapping of the connected components resulting from the filtration process.
- An exhaustive evaluation using both labeled data as well as case studies over unlabeled data showing the potential of TopoMap to support the analysis of high-dimensional data as well as distance-based projection techniques.

To the best of our knowledge, TopoMap is the first dimensionality reduction technique to provide guarantees as to the preservation of the topological properties of the Rips filtration of the data under analysis.

2 Related Work

In order to better contextualize the proposed methodology, we organize the related work in two main parts, topological data analysis (which describes related work in topological data representations) and topology-based multidimensional projection (which describes how these topological data representations can drive projection methods).

2.1 Topological data analysis

Topology-based methods [24] have been very popular in the last two decades to support advanced data analysis and visualization tasks [27]. By providing a concise, structural representation of the data, these techniques greatly help in the visualization and analysis of the data. They have been applied successfully to a variety of domains, such as astrophysics [69, 73], biological imaging [2, 9, 14], chemistry [8, 32, 59], fluid dynamics [41], material sciences [34, 35, 72], or turbulent combustion [11, 33, 44].

The Rips filtration [24] is often used to analyze the topology of high dimensional point clouds, and is motivated by the work by Chazal and Oudot [16] who showed that Rips filtrations can provably capture the homology of the manifold sampled by the point cloud. Among the popular representations in topological data analysis, the Reeb graph [63] is obtained by contracting to a single point each of the connected components of level sets of an input scalar field, resulting in a characteristic skeleton-like representation of the input data. For discrete point cloud data, the Mapper [71] is an approximation of the Reeb graph of some user-defined function (often called lense function) defined over a nearest neighbor graph of the input point cloud. Another popular abstraction is the Morse-Smale complex [21], which is a cellular decomposition of the domain of an input scalar field, such that all the points of a given cell admit the same gradient integration extremities. For discrete point cloud data, the Morse-Smale complex has been used over the k-nearest neighbor graph of the input point cloud for clustering purposes [15]. As discussed next, all of these representations (Mapper, Reeb graph, Morse-Smale complex) have also been used as a driving data representation for dimensionality reduction. Given the increasing popularity in using topology-based techniques for data analysis, it is not surprising that there are several open source tools and libraries available [1, 12, 27, 49, 52, 53, 75].

2.2 Topology-based Multidimensional Projection

Multidimensional projection has long been a fundamental analytical tool, mainly in the context of visualization [24, 54]. In fact, the visualization community has not only proposed a number of MDP methods tailored to visual analytic tasks [39], but has also developed methodologies to facilitate the analysis of MDP distortions [4, 50] and to enrich MDP layouts so as to uncover information hidden in the projections [11, 33]. The extensive literature about MDS/MDP techniques has been organized over several books [10, 47] and surveys [19, 54, 77]. In order to emphasize our contribution, we focus on techniques that explicitly rely on topological concepts to perform and assess multidimensional projections, disregarding distance preserving methods such as the classical MDS [37] and neighborhood preserving techniques such as LLE [66], t-SNE [76], and Lamp [39]. We refer interested readers to the above books and surveys for a broader discussion about MDS/MDP methods.

Isomap [74] is probably one of the first MDS techniques to resort to topological mechanisms to accomplish dimensionality reduction. Isomap aims to capture the topological (manifold) structure of the data through a graph representation from which geodesic distances are computed. A number of variants of Isomap have been proposed, including Landmark-Isomap [8, 6], out-of-sample [7] and spatio-temporal extensions [19]. An interesting variant of Isomap is the method proposed by Lee and Verleysen [46], which tears a graph representation of the data so as to preserve essential (non-contractable) loops, thus enabling loop preserving manifold unfoldings. The recent work by Yan et al. [79] is another particularly interesting variant of Isomap (precisely, a variant of L-Isomap). Similar to previous work on skeletonization [43], this approach identifies cycles in the original data, but additionally aims at preserving these cycles when projecting the data to 2D. Specifically, it focuses on the Mapper (cf. Sect. 2.1) of a function defined on the KNN-graph structure of the data to select landmark points. The underlying idea is that the topology-based landmark selection captures the structure of the 1-dimensional homology groups of the data, which hopefully are preserved during the dimensionality reduction phase accomplished via regular L-Isomap. However, their approach does not take into account 0-homology groups which are therefore not preserved. In contrast, the TopoMap method proposed in this work provides theoretical guarantees as to 0-dimensional homology group preservation, thus ensuring that the connected components visualized in the projection layout are the same as in the original high-dimensional data, according to its Rips filtration. Similar to Yan et al. [79], Gerber et al. [29, 30] introduced projection methods driven by topological data representations. In particular, it differs from our work in the sense that the introduced projections are driven by the network of cells of maximum dimension (called crystals) of the Morse-Smale complex (Sect. 2.1). They do not aim at specifically preserving the persistence diagram of the Rips complex as studied in this paper and therefore encode a different information, specifically tailored for regression tasks.

In scientific visualization, Weber et al. [78] introduced a terrain
Wang [36] proposed algorithms to generate an ensemble of terrains whose elevation is carefully designed, such that the contour tree of the density (which are not taken into account in terrain metaphors). For instance, in the resulting elevation can also be displayed as a planar heat map visual results which are highly similar to t-SNE. For this reason, it is evaluated to the quality of dimensionality reduction techniques. A good example is the work by Rieck and Leitte [55], which assesses the quality of a projection technique from the 2nd Wasserstein distance between persistence diagrams computed from the original and projected data. In a follow up work, Rieck and Leitte [55] proposed the use of persistent homology to compare quality measures for dimensionality reduction, making possible to analyze the agreement of multiple quality measures, thus identifying regions where different quality measures disagree the most. Persistent homology has also been employed by Paul and Chalup [60] as a mechanism to validate dimensionality reduction methods when applied to particular benchmark data. As we shall prove later, TopoMap is guaranteed to preserve connected components under filtration, and is therefore exact (no error) when comparing the 0-dimensional homology persistence diagrams generated by a filtration in the visual and original spaces respectively.

### 3 Topology Preserving Projection

Given a data set that is a collection of high-dimensional points in $\mathbb{R}^d$, a common topology-based approach to analyze this data is to study the evolution of cycles in the simplicial complexes resulting from a Euclidean distance based Rips filtration over these points. Our goal is to project the data onto $\mathbb{R}^2$ such that above evolution for a subset of the cycles is preserved in the projected space as well.

In this section, we first introduce the necessary notations and formalize the problem that is of interest in this work. We refer the reader to Edelsbrunner and Harer [24] for a comprehensive discussion on these topics. Next, we describe a high level approach for solving the problem, and discuss different choices that can be made in the implementation of the high level solution.

#### 3.1 Problem Formulation

**VietorisRips complex.** Let $P = \{p_1, p_2, \ldots, p_n\}$ be a set of points in $\mathbb{R}^d$. Given a distance threshold $\delta$, the VietorisRips complex (or Rips complex), is defined as the set of all $k$-simplices $K \subseteq \mathbb{R}^d$, $|K| = k + 1, k \geq 0$, such that $d(p_i, p_j) \leq \delta, \forall p_i, p_j \in K$. Here, $d(\cdot, \cdot)$ is the Euclidean distance. Intuitively, the Rips complex for a distance threshold $\delta$ captures the shape of the data when each point $p_i$ is replaced with a $d$-dimensional ball of diameter $\delta$ centered around it. For example, consider the 6 points in $\mathbb{R}^3$ shown in Fig. 2(a). Fig. 2(b)-(f) illustrates this shape for 5 different values of the distance threshold $\delta$.

**Rips Filtration.** Consider a model where the distance threshold $\delta$ is increased from 0 to $\infty$. That is, the $d$-dimensional balls are gradually grown in size. A Rips filtration captures this growth model.

Consider an ordered set of simplexes $\mathcal{P} = \{K_0 = \emptyset, K_1, K_2, \ldots, K_m\}$. Let $\delta_i, i \in [0, m]$, be the smallest distance threshold such that simplex
$K_i$ is part of the Rips complex defined for $\delta_i$. Then, the above ordered set is a Rips filtration if $\forall i, j, i < j$:

1. $\exists l \leq i$ s.t. $K_l \cap K_j = K_i$; and
2. $\delta_l \leq \delta_j$.

**Topological Persistence and Persistence Diagram.** Consider the growth as defined by the Rips filtration, wherein the simplices from the filtration are added one at a time. That is, the $i^{th}$ iteration in this growth will consist of the subset $S_i = \{K_0, K_1, \ldots, K_i\}$. The addition of each new simplex can change the topology of the underlying data, where the topology is captured by the set of cycles in the simplicial complex defined by $S_i$. More specifically, a new $k$-cycle, $k > 0$, can either be created or an existing $k$-cycle can be destroyed \[25\]. Informally, a 0-cycle corresponds to a connected component, a 1-cycle to a loop, 2-cycle to a void, and so on. Given one such $k$-cycle, let $\delta_k$ be the threshold at which this cycle is created, and $\delta_k$ the threshold at which it is destroyed. Then the topological persistence \[25\] of this $k$-cycle is defined as $\delta_k - \delta_k$, and intuitively captures the lifetime of this cycle in the given filtration. Note that a cycle that is not destroyed has a persistence equal to infinity.

The persistence diagram \[17\] plots all the cycles created during the filtration as a scatter plot, where the coordinates of the point corresponding to a cycle is its creation and destruction thresholds (i.e., the $x$- and $y$-axes of this plot corresponds to the creation and destruction thresholds).

**Problem Definition.** Let $PD_k^p$ denote the persistence diagram restricted to $k$-cycles computed using the Rips filtration over the point set $P$. Given a set of points $P = \{p_1, p_2, \ldots, p_n\}$ in $\mathbb{R}^d$, our goal is to compute the corresponding set of points $P' = \{p'_1, p'_2, \ldots, p'_n\}$ in $\mathbb{R}^2$ such that $PD_k^p = PD_k^{P'}$, where there is a one to one correspondence between the connected components or 0-cycles (i.e., a point $p_i$ belongs to a 0-cycle w.r.t. $P$ if, and only if, the point $p'_i$ will belong to the corresponding 0-cycle w.r.t. $P'$).

In other words, the Rips filtration over the projected points $P'$ not only has the exact same connected components during each iteration of the growth, but even the iterations at which they are created and destroyed are the same when compared to the Rips filtration over the high-dimensional points $P$.

### 3.2 Approach

Since we are interested only in the evolution of the set of connected components, it is sufficient to consider only the 0- and 1-simplices (vertices and edges respectively) of the filtration. Consider the set of edges in the above filtration. Only a subset of these edges result in a change in topology, or in other words, merge two disconnected components into a single component. The following lemma bounds the number of such topology changing edges in the filtration.

**Lemma 1.** Given a Rips filtration defined over a set of $n$ points, there is exactly $n - 1$ topology changing edges that result in reducing the number of $0$-cycles.

**Proof.** Consider an input with $n$ points. At the beginning of the filtration, say at an infinitesimally small threshold $\varepsilon > 0$, there are a total of $n$ components corresponding to an input point. The addition of each topology changing edge reduces the count of connected components by one. Thus, there exists exactly $n - 1$ such edges until there exists just a single connected component.

The 5 edges in Figures 2(b)–2(f) correspond to the topology changing edges in the Rips filtration over the 6 points in Fig. 2(a).

Let $\mathbb{K}_0 = \{p_1, p_2, \ldots, p_n, e_1, e_2, \ldots, e_{n-1}\} \subset \mathbb{K}_0$ be the subset of a filtration, where $p_i, 1 \leq i \leq n$, are the set of input points and $e_i, 1 \leq i < n$, corresponds to topology changing edges (in order of their appearance in $\mathbb{K}_0$). Note that we ignore all other edges in $\mathbb{K}_0 \setminus \mathbb{K}_0$, since they do not change the topology with respect to 0-cycles.

Consider only the ordered set of topology changing edges $\mathbb{K}_0 = \{e_1, e_2, \ldots, e_{n-1}\}$ from the above filtration. By definition, the length of these edges satisfies $|e_1| < |e_2| < \ldots < |e_{n-1}|$. While these inequalities might not hold in practice (two consecutive edges could have the same length), a simulated small perturbation \[25\] of the points can ensure this property holds. The following lemma, which shows the equivalence between $\mathbb{K}_0$ and the Euclidean distance minimum spanning tree (EMST) computed over $P$, provides the basis for our projection algorithm.

**Lemma 2.** Given a set of points $P = \{p_1, p_2, \ldots, p_n\}$, let $G$ be the complete weighted graph defined over $P$ such that the weight of each edge $(p_i, p_j)$ is equal to the Euclidean distance $d(p_i, p_j)$ between the corresponding end points. Then, the ordered set of topology changing edges $\mathbb{K}_0 = \{e_1, e_2, \ldots, e_{n-1}\}$ is precisely the set of edges of the minimum spanning tree (MST), in increasing order of weight, computed over $G$.

**Proof.** We prove this by showing, through induction, that the ordered set of topology changing edges are the same as those added by the Kruskal’s algorithm \[18\].

Consider the first edge $e_1$ of the filtration. By definition, it is the edge with the smallest length, and thus also the first edge that is added by the Kruskal’s algorithm. Let, edges $e_1, e_2, \ldots, e_{n-1}$ be the first $i - 1$ edges added by the Kruskal’s algorithm. The induction hypothesis is that at this stage, the connected components created by the filtration is exactly the same as the set of connected trees created by the Kruskal’s algorithm.

Now, consider the $i^{th}$ topology changing edge of the filtration $e_i$. For sake of contradiction, say the $i^{th}$ edge added by the Kruskal’s algorithm is $e'_i \neq e_i$. This implies that the edge $e_i$ has length less than that of $e'_i$, and connects two connected subtrees together. Then, by definition, $e_i$ will occur before $e_i$ in the filtration, and will also be a topology changing edge. Thus, the case of $e_i \neq e'_i$ is not possible.

The following proposition follows from the above lemmas and it guarantees the existence of a mapping (projection) that retains the topology with respect to 0-cycles in the projected space.

**Proposition 1.** Let $P$ be a set of points in $\mathbb{R}^d$, $\mathbb{K}_0 = \{e_1, e_2, \ldots, e_{n-1}\}$ be the ordered subset of topology changing edges in $\mathbb{K}_0$ and $\mathbb{C}_p$ be the set of connected components obtained during the filtration over $\mathbb{K}_0$ after the addition of the first $i$ topology changing edges $\{e_1, e_2, \ldots, e_i\}$. Let $\mathcal{M} : \mathbb{R}^d \rightarrow \mathbb{R}^2$ be a mapping that maps points in $P$ to $P'$, and $\mathbb{K}_0' = \{e_1', e_2', \ldots, e_{n-1}'\}$ be the set of topology changing edges in $\mathbb{K}_0'$. Then, there exists at least one mapping $\mathcal{M}$ satisfying the following properties:

(a) edge lengths $|e'_i| = |e_i|, \forall i \in [0, n-1]$;

(b) the components generated by the filtrations are identical, i.e., $\mathbb{C}_p = \mathbb{C}_p' \forall i \in [0, n-1]$; and

(c) $PD_0' = PD_0$, where $PD_0'$ and $PD_0$ are the persistence diagrams of $\mathbb{K}_0'$ and $\mathbb{K}_0$ respectively.

We abuse notation in the above proposition when stating that $\mathbb{C}_p = \mathbb{C}_p'$. What this notation means is that the mapping $\mathcal{M}$ establishes a one-to-one relation between the components in $\mathbb{C}_p$ and $\mathbb{C}_p'$, that is, every point in a component $C' \subset \mathbb{C}_p'$ is the image of a point in the corresponding component $C \subset \mathbb{C}_p$. Note that in the above proposition, guaranteeing properties (a) and (b) is a sufficient condition for (c).

The proof of Proposition 1 is constructive and is provided in Sect. 3.3. In fact, using the above lemmas, we design the iterative algorithm shown in Procedure TopoMap that projects a set of high-dimensional points onto $\mathbb{R}^2$ while guaranteeing the properties stated in Proposition 1. The algorithm places the points onto a plane such that the minimum spanning tree edges are preserved. In other words, it “draws” the minimum spanning tree maintaining the edge lengths.

The algorithm initially maintains all the points as a separate component, and stores the minimum spanning tree edges as an ordered set. In each iteration, the algorithm then adds the smallest edge from this ordered set to connect two components, thus reducing the number of maintained components by one. The length of the edge is preserved during this step, that is, its placement is such that distance between the
two components is equal to the length of the connecting edge that, in turn, has the same length as its counterpart in the original space. This edge is then removed from the edge set and the process repeated until all edges are appropriately placed. The key aspect of the algorithm is Line 8 that places the points based on the minimum spanning tree edge lengths. We describe different ways of accomplishing this in the next section. The maintenance of the set of connected components is accomplished using the union-find data structure [18].

3.3 Building the Topology Preserving Mapping

The TopoMap algorithm starts with $e_1$, the smallest topology changing edge in $E_0$. Placing the end points of $e_1$ (which are individual components at the start of this procedure) in a lower dimensional space such that this distance is preserved is straightforward. In other words, $e_1 = e'_1$. Now suppose that the first $i - 1$ topology changing edges have been added while ensuring Proposition 1. In the $i^{th}$ step, let edge $e'_i$ be added so as to connect two components from $C_{P-1}$ that are counterparts of the components in $C_{P-1}$ linked by $e_i$. As mentioned in Line 8 of the algorithm, the goal now is to place these two components such that the minimum distance between them is equal to $|e_i|$. If this condition is satisfied, then the properties $|e_1| = |e'_1|$ and $C_{P-1} = C_{P}$, for $j \in \{1, \ldots, i\}$, are naturally attested. By repeating the process for all $e_i$, $i \in \{1, \ldots, n-1\}$, we also guarantee that $PD_{P}=PD_{P}$. Therefore, proving Proposition 1 now requires showing that there exists a way to place two components connected by an edge $e'_i$ whose length is $|e'_i|$. In fact, there are several ways in which this can be accomplished as we show next. Note that, there always exists a valid solution to this problem.

A solution in 1-dimensional space. Consider two sets $C_1$ and $C_2$, where each point in this set is associated with a $x$ and $y$ value corresponding to its 2D coordinates. Let $p_{1} \in C_{1}$ be the rightmost point in $C_{1}$. Similarly, let $p_{2} \in C_{2}$ be the leftmost point in $C_{2}$. Let the input edge length be $d$. The trivial solution is to simply translate all points in $C_1$ such that $(p_{x,x}, p_{y}) = (d, 0)$ and $(p_{x,y}) = (d, 0)$.

A geometric 2D solution. Note that the above solution places all the points only along the $x$-axis. Thus, to obtain a more compact solution that also uses the second dimension, we modify the above solution as follows, by arranging components in the plane with local rotations, similarly to circular layout strategies in tree drawing [67]. Let $hull(C_1)$ and $hull(C_2)$ be the convex hulls of the two components. Pick an edge $e_1$ from $hull(C_1)$ and $e_2$ from $hull(C_2)$. Transform (rotate) $C_1$ such that $e_1$ is parallel to the $x$-axis, and is the topmost edge of the convex hull (i.e., has the highest $y$-coordinate). Similarly, transform $C_2$ such that $e_2$ is also parallel to the $x$-axis, but is the bottommost edge. Let left(edge) denote the left endpoint of the edge $e$. Now, translate component $C_1$ such that left($e_1$) = $(0,0)$, and component $C_2$ such that left($e_2$) = $(0,d)$. Alternatively, the right endpoints of the edges $e_1$ and $e_2$ can be used as well to align the two components.

There are different ways in which the edge of the convex hull can be selected (as well as to decide which end point is used for the alignment). We decided to choose the edge that contains one of the end points of the minimum spanning tree edge that is under consideration. In case this is not possible, we choose the edge closest to this point. The intuition here is to not only preserve the connected components after every iteration, but also to try and preserve the end points of the minimum spanning tree edges as much as possible.

Fig. 4 illustrates this procedure for the example points in Fig. 2. Note that the addition of the first two edges result in the same state as in the 1D solution above. However, when the third edge $(p_5, p_6)$ is added, then the point $p_6$ is placed in a perpendicular orientation. When the last edge $(p_5, p_6)$ is processed, since both components have more than two points each, the convex hull is used to perform the alignment by appropriately transforming both components.

An optimization-based 2D solution. During data analysis, in addition to preserving the topology changing edges of the filtration, it might be beneficial to also possibly preserve other properties as much as possible. In this section, we show how our projection approach can be tuned to support such modifications.

For example, it might be natural to consider a case where we are also interested in keeping the resulting projection “compact”, in the sense that we want the points to be as close to each other as possible, while still ensuring that the distance between the two components, $C_1$ and $C_2$, is equal to the given value. One way of doing this is to minimize the sum of squared distances between the points in $C_1$ and $C_2$ after the placement. This can be achieved using the optimization model described next.

First, fix one of the components, say $C_1$, and expand it to contain all points in the plane that have distance to $C_1$ less or equal $d$ (this is the region which should not contain any point from $C_2$). This is achieved by considering lines that are parallel and at a distance $d$ to the edges of $hull(C_1)$. Since this expanded set of lines is also a convex hull, its inner region can be described by a set on linear inequalities on the plane. Let this set of linear inequalities be denoted by $Ax \leq b, A \in \mathbb{R}^{k \times 2}, b \in \mathbb{R}^k$.

The goal then becomes to find the rigid motion (rotation plus translation) that applied to $C_2$ minimizes the sum of the squared distances to the expanded convex hull without penetrating it. Formally, this problem can be mathematically formulated as:
Fig. 5. Solving the optimization model to place two sets of points. (a) Two components that are to be merged. (b) The feasibility region with respect to the highlighted edge of \( hull(C_1) \) is shaded gray. \( C_2 \) is initially placed in this region and the optimization is solved. (c) Solution.

\[
\min_{\theta,t} \sum_{p_1 \in C_1, p_2 \in C_2} \|p_1 - (R(\theta)p_2 + t)\|^2 \\
\text{s.t. } A_i(R(\theta)p + t) \leq b_i, \forall p \in C_2.
\]

where \( \theta \) represents an angle with rotation matrix \( R(\theta) \) and \( t \) is a translation vector. Note that it is possible to consider only the points that define the convex hull of \( C_2 \) above.

This problem can be cast as a mixed integer nonlinear optimization problem. The integer variables are needed because the constraints in the above model are actually a “union of sets” instead of an “intersection of the sets” that is usual in optimization. Unfortunately, solving such problems for a large number of points is impractical. We therefore decided to use a simplified heuristic that (i) individually optimizes with respect to each edge of the convex hull \( hull(C_1) \); and (ii) minimizes the sum of distances between points in \( C_2 \) to a single point in \( C_1 \). Formally, let \( A_i \) be a row of the matrix \( A \) and \( b_i \) the respective right-hand side. Let \( p' \in C_1 \) be a point of interest. We first solve the following optimization problem for \( i = 1, \ldots, k \).

\[
\min_{\theta,t} \sum_{p \in C_2} \|p - (R(\theta)p + t)\|^2 \\
\text{s.t. } A_i(R(\theta)p + t) \geq b_i, \forall p \in C_2.
\]

We then consider as final solution the one that obtained the smallest objective value. If we also want to preserve the edge from the filtration in the projection, we can apply the objective function above, such that the edge endpoint in \( C_2 \) has higher weight when compared to other points. In our implementation, we consider \( p' \) to be the end point of the edge that is being processed in that iteration.

\[
\text{Fig. 5} \text{ illustrates this optimization process. It shows sets of points } C_1 \text{ (colored violet) and } C_2 \text{ (colored orange) that are to be placed at a distance } d \text{ from each other (Fig. 5a). The points colored red and blue correspond to the filtration edge under consideration. The blue point is chosen as the point of interest in order to minimize the objective function of the optimization. The points in } C_2 \text{ are first randomly placed in the feasibility region corresponding to one of the edges of } hull(C_1), \text{ and the optimization problem is solved (Fig. 5b)). The resulting solution is shown in Fig. 5c).}
\]

While the simplified optimization model is also nonlinear (due to the rotation), it does not have integer variables and can then be solved by standard nonlinear optimization algorithms. Note that the final solution is not guaranteed to have two points, one in \( C_1 \) and the other in \( C_2 \), at exact distance \( d \) (and therefore, do not satisfy the required filtration constraint). However, this can still be ensured by sliding \( C_2 \) parallel to the edge of \( hull(C_1) \) that is associated to the solution obtained in the optimization process.

### Implementation

TopoMap was implemented using C++. It can be divided into two phases. First is to compute the Euclidean distance minimum spanning tree, for which we used the implementation of the dual tree EMST algorithm [48] provided by the mlpack library [20], and has a time complexity of \( O(N \log N) \), where \( N \) is the size of the input. The next phase is to layout the points. Each iteration of TopoMap aligns two components corresponding to the MST edge being processed. We use the union-find data structure to maintain the list of components, which can be accomplished in \( O(N \alpha(N)) \) time. The convex hull of the resulting merged component is then computed using the qhull library [4], which takes \( O(n \log n) \) time to compute the convex hull of \( n \) points. However, since in each iteration, we use only the points in the convex hull of the individual components, \( n < < N \) in practice. On several large data sets, we found that the layout phase using the geometric approach scaled linearly with the input. On the other hand, computing the EMST became the primary bottleneck when increasing dimensions.

For the optimization based approach, we use the Algencan [61, 62] library for solving our optimization model. It is a robust and high performance implementation of the augmented Lagrangian method for nonlinear optimization problems whose code is freely available. As expected, the optimization approach was slower than the geometric approach. However, the main bottleneck was still the EMST phase especially for large point clouds.

### 3.4 Robustness

From the topological perspective, it is well known that the persistence diagram is robust to noise [17], especially in the context of topology inference [16]: small displacements of points in the original space induce small variations in the persistence diagram. Since TopoMap strictly preserves the persistence diagram, topological robustness to noise is guaranteed by definition. On the other hand, the ordering of the filtration might change slightly due to the noise induced perturbation. Thus, with respect to the actual projection itself, the locations of the points in 2D space might vary marginally when using the geometric solution. Note that the optimization-based approach, being non-deterministic, can produce different layouts even for the same input when run multiple times. However, since the connected components (that represent the points in the persistence diagram) are robust, these components are always maintained by the projection.

### 4 Mapping Evaluation and Interpretation

In this section we present the results of applying TopoMap to project high-dimensional data to a visual space (\( \mathbb{R}^2 \)). Our goal in this section is to analyze the properties of the layout produced by TopoMap, the way it visually encodes the information contained in the high-dimensional data, and how much “readable” the TopoMap layout is when compared to the ones produced by dissimilarity preserving projection methods.

To facilitate the above analyses, we use several data sets (see Table 1) having different numbers of instances and dimensions, some of which are labeled (i.e., the classes are known for the instances). We used the implementations provided by scikit-learn (v0.19.0) for existing methods. All experiments were run on a machine with an Intel(R) Xeon(R) CPU E5-2630 v2 running at 2.60GHz and 64 GB of memory.

| Data set | # Instances | # Attributes | # Classes |
|----------|-------------|--------------|-----------|
| Iris     | 150         | 3            | 3         |
| Seeds    | 210         | 8            | 3         |
| Heart    | 261         | 11           | 2         |
| Cancer   | 699         | 11           | 2         |
| Meat     | 20000       | 64           | 10        |
| MNIST    | 20000       | 784          | 10        |
| Urban    | 17520       | 6            | not labeled |
TopoMaps are quite different from the ones produced by dissimilarity-based methods. This is not a surprise, since TopoMap preserves the connected components observed in the original space and to nicely reflect their relative adjacencies. Fig. 6 performs the same comparison with respect to the first five data sets in Table 1.

In Table 1, we compare the layout produced by MDS, Isomap, t-SNE, UMAP, and TopoMap for 3 synthetic data sets with well known properties. The first is simply a set of points sampled from three Gaussians, the second is points sampled from three rings, while the third is sampled from two concentric spheres. This figure illustrates the ability of TopoMap to preserve the connected components observed in the original space and to nicely reflect their relative adjacencies.

One can notice from these examples that the layouts resulting from TopoMap are quite different from the ones produced by dissimilarity-based methods. This is not a surprise, since TopoMap preserves $n - 1$ distances between connected components while dissimilarity-based methods try to preserve $n^2$ distances (or distributions in the case of t-SNE) between instances.

**Star Shaped Ensembles.** TopoMap produces a layout made up of star shaped ensembles with branches connecting and emanating from them. One way of interpreting this layout is through the use of the equivalence between the 0-homology filtration of the Rips complex and hierarchical clustering using the single-linkage criterion [13]. The connected components built during the filtration are exactly the same as the clusters formed when moving up in the hierarchy. In other words, hierarchical clustering with single-linkage produces, by construction, identical results when considering the input (high-dimensional) data and the two-dimensional projection provided by TopoMap. Thus, when interpreting our projections, users should visually identify centers of density-based visualization, it is easy to see that branches stemming from the starred ensembles typically are low density regions.

The density-based visualization makes it easier to identify tightly connected groups. Although density-based visualizations have been used to evaluate dissimilarity-based methods [50], the presence of errors and distortions prevent the analysis from being accomplished with high confidence [3]. Note that the “centers” of the starred ensembles correspond to denser areas of the layout, thus corresponding to tightly grouped data instances. This is evident in the examples involving the Heart and Cancer datasets, where a density analysis in the layout resulting from MDS, Isomap, and t-SNE would be of little use.

**Branches and Outliers.** When considering the above mentioned density-based visualization, it is easy to see that branches stemming from the starred ensembles typically are low density regions. These branches are essentially of two types: those connecting the starred ensembles; and the ones emanating outwards from the stars. The latter is composed of points whose neighborhoods are not tightly connected. From the hierarchical clustering perspective, these can be considered as single point clusters (outliers for example) which merge with already existing large clusters as one moves up the hierarchy.

For example, the TopoMap projection of the Cancer dataset in Fig. 6 clearly pinpoints the compact class (in red), but it splits the blue class into a number of local clusters, increasing the potential for misleading interpretation. TopoMap, on the other hand, shows two star shaped ensembles, one well defined and another more elongated, indicating the presence of two classes, one of them not so compact. The density-based visualization makes it easier to identify tightly connected groups. Although density-based visualizations have been used to evaluate dissimilarity-based methods [50], the presence of errors and distortions prevent the analysis from being accomplished with high confidence [3]. Note that the “centers” of the starred ensembles correspond to denser areas of the layout, thus corresponding to tightly grouped data instances. This is evident in the examples involving the Heart and Cancer datasets, where a density analysis in the layout resulting from MDS, Isomap, and t-SNE would be of little use.
Transitioning Between Clusters. Branches connecting the centers of starred ensembles tend to encapsulate instances that lie between clusters, and represent a transition from one group to another. To illustrate this property, we used a uniform sample of 20,000 instances from the MNIST data set, and projected it using the angular distance as distance metric (see Sect. 4.4 for a discussion on this). Fig. 7 highlights the different transitions between the star ensembles captured by TopoMap. In particular, note that the cluster corresponding to class 8 transitions to 0 (a), 3 (b), as well as 1 (d). Other transitions such as from class 7 to 9 (c) and from 6 to 0 (e) can also be clearly seen. When well defined clusters are far apart from each other, as is the case of cluster 0 and 8, the density visualization, coupled with the guarantee that the topology changing edges’ lengths are preserved by TopoMap, gives us confidence to claim that the longer branches come from the sparser class. Moreover, outliers tend also to be part of the loose branches, mainly in less dense areas of the layout. This fact can be observed in the projection of the Mfeat dataset, where classes become mixed in loose branches (TopoMap column left image), but not in the center of the ensembles.

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Fig. 7. Mnist data projected using TopoMap (using cosine distance). Transitions between the different starred ensembles clusters: (a) 0 and 8. (b) 3 and 8. (c) 7 and 9. (d) 1 and 8. (e) 0 and 6. (f) class 2 while being a cluster, is far from 0 and is connected to it via outliers.

contains one compact and another more sparse class. The sparse class (red) gives rise to a starred ensemble with a small “center” and long branches emanating from it. The density visualization, coupled with the guarantee that the topology changing edges’ lengths are preserved by TopoMap, gives us confidence to claim that the longer branches come from the sparser class. Moreover, outliers tend also to be part of the loose branches, mainly in less dense areas of the layout. This fact can be observed in the projection of the Mfeat dataset, where classes become mixed in loose branches (TopoMap column left image), but not in the center of the ensembles.

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There are two main scenarios that can result in a loss of information. The first scenario is when exploring the long branches of the star ensembles. The distance between two points adjacent in a long branch is not necessarily the distance between them in the high dimensional space. Rather, it represents the distance between the corresponding connected components. For example, say a point $p_1$ is connected to $p_2$ in the minimum spanning tree (and hence is an edge of the filtration). This does not guarantee that $p_1$ and $p_2$ will form an edge in the projection—the edge will be between the connected components corresponding to $p_1$ and $p_2$. Thus, the two points may be assigned to different branches of a star ensemble depending on the strategy used during the projection. This property must be considered when interpreting the layout.

Layout interpretation guideline. Based on the above observations, we use the following guideline to explore data using TopoMap for the remainder of this paper:

- Use a density-based colormap to visualize the projection.
- Start exploration by looking at centers of stars with high density. These typically represent clearly distinct clusters in the data.
- Use low density stars to study “uncommon behaviors”.
- Explore branches to analyze sparse clusters and outliers.

4.2 Case Study with Unlabeled Data

There are several urban data sets available representing different facets of the city corresponding to its different properties. These are typically studied in isolation, and can sometimes result in missing out on interesting patterns resulting from the interactions between these facets. For example, when analyzing just the count of taxi trips, it is easy to observe that both Times Square as well as Penn Station are identified as hot spots almost throughout the day. Given that the former is a popular tourist attraction, while the latter is a transit hub, one would however expect differences in the way usage patterns of these places change depending on other conditions.

In this experiment, our goal is to see if such patterns do exist. To do so, we generate high dimensional data sets by combining the NYC taxi data and the weather data as follows. We divided two years (2014–2015) into hourly intervals. Then, given a location, we consider the taxi pickups that happened within a 100 m radius of the location. We then create one high dimensional point for every hourly interval having the following dimensions: count of taxi pickups, average fare, average distance, temperature, precipitation (rainfall), and wind speed. Thus, the data set corresponding to each location is a collection of 6D points.

We then projected the data corresponding to Times Square and Penn Station using TopoMap and visually analyzed the patterns present in the projection and analyzed the different starred ensembles by looking at the temporal distribution of the points forming these clusters. The analysis of the data corresponding to Penn Station can be found in the supplemental material.

Times Square. Fig. 8 shows the results obtained for Times Square. In this scenario it is interesting to note that the most dense region (Figs. 8(b) and (c)) correspond to the summer months. This is in turn divided into two clusters: Fig. 8(b) corresponding to main part of the day (10 am to 6 pm), while Fig. 8(c) corresponds to night hours. Similarly the winter months formed its own cluster (Fig. 8(d)). It was interesting to note that there was also a cluster with a smaller number of points corresponding to the Spring and Autumn months (Fig. 8(e)). A curious observation in the above cases was that none of these clusters included the time period 4 am–8 am. We found these points for summer in a separate cluster shown in Fig. 8(f). On further analysis, we found that this is primarily because the taxi activity at these times was not only lower than the other times, but that these trips also had longer distances and fares than normal. Additionally, we also found that the points corresponding to periods when there was rainfall formed a less dense cluster among the outliers (Fig. 8(g)).

4.3 Probing Dissimilarity-based Projections

Since TopoMap bears theoretical guarantees it can be used to probe other projection methods in order to further understand how those methods behave, especially regarding distortions and cluster preservation. To illustrate this, consider the layouts in Fig. 9. The layout on the top is
Fig. 8. Analyzing Times Square using TopoMap. (a) Projection obtained using TopoMap. (b)–(g): Different clusters are selected and the temporal distributions of the selected points visualized as a histogram.

Fig. 9. Connected components (colored groups) guaranteed to exist in the high-dimensional space are broken apart by t-SNE.

With the help of TopoMap, one can realize where t-SNE is placing the different pieces of a cluster. Although experienced users are usually aware about this “breaking cluster” property of t-SNE, we are not aware of any work capable of revealing the extent/intensity of this phenomenon. Revealing this nature of t-SNE is quite important, and can be considered as a side contribution of the present work helping to illustrate the potential of using TopoMap as an analytical tool.

4.4 Discussions
Using alternate distance metrics. As shown in Sect. 4.1 (Fig. 7) TopoMap can also be used with an alternate distance metric. This requires computing the MST using this metric, in which case the running time for computing the MST degenerates to $O(N^2)$ due to the computation of the distance matrix. Note that when another distance metric is used, the filtration in the projected space is still preserved with respect to Euclidean distance in the visual space. This also makes it easier for the user to gauge the projection in the visualized space, allowing for a comparison between the effect of using different distance metrics.

Other 2D and 3D solutions. While our approach provides a solution ensuring that the 0-dimensional homology is preserved, there can be other valid solutions as well. Depending on the application, one can also trade-off preserving the persistence of outliers to preserving neighborhoods, or optimizing for a different property. Similarly, it would be interesting to see how the outliers would behave when moving to 3D.

5 Conclusions
In this paper we presented TopoMap, the first planar projection technique that is guaranteed to preserve the homology of 0-cycles of the Rips filtration. Evaluation of our approach using a variety of data sets demonstrated several key properties that are desirable in a visual analytical tool: the layout is easy to understand while its theoretical guarantees provide confidence to the users. In the future, we would like to explore ways in which 1-cycles can be preserved as well in the projection. Analyzing the effectiveness of TopoMap to assist clustering mechanisms is another direction we will pursue.

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