Introduction to the R package TDA

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Abstract

We present a short tutorial and introduction to using the R package TDA, which provides some tools for Topological Data Analysis. In particular, it includes implementations of functions that, given some data, provide topological information about the underlying space, such as the distance function, the distance to a measure, the kNN density estimator, the kernel density estimator, and the kernel distance. The salient topological features of the sublevel sets (or superlevel sets) of these functions can be quantified with persistent homology. We provide an R interface for the efficient algorithms of the C++ libraries GUDHI, Dionysus, and PHAT, including a function for the persistent homology of the Rips filtration, and one for the persistent homology of sublevel sets (or superlevel sets) of arbitrary functions evaluated over a grid of points. The significance of the features in the resulting persistence diagrams can be analyzed with functions that implement the methods discussed in Fasy, Lecci, Rinaldo, Wasserman, Balakrishnan, and Singh (2014), Chazal, Fasy, Lecci, Rinaldo, and Wasserman (2014c) and Chazal, Fasy, Lecci, Michel, Rinaldo, and Wasserman (2014a). The R package TDA also includes the implementation of an algorithm for density clustering, which allows us to identify the spatial organization of the probability mass associated to a density function and visualize it by means of a dendrogram, the cluster tree.

Keywords: Topological Data Analysis, Persistent Homology, Density Clustering.

1. Introduction

Topological Data Analysis (TDA) refers to a collection of methods for finding topological structure in data (Carlsson 2009). Recent advances in computational topology have made it possible to actually compute topological invariants from data. The input of these procedures typically takes the form of a point cloud, regarded as possibly noisy observations from an unknown lower-dimensional set $S$ whose interesting topological features were lost during sampling. The output is a collection of data summaries that are used to estimate the topological features of $S$.

One approach to TDA is persistent homology (Edelsbrunner and Harer 2010), a method for studying the homology at multiple scales simultaneously. More precisely, it provides a framework and efficient algorithms to quantify the evolution of the topology of a family of nested topological spaces. Given a real-valued function $f$, such as the ones described in Section 2, persistent homology describes how the topology of the lower level sets $\{x : f(x) \leq t\}$ (or superlevel sets $\{x : f(x) \geq t\}$) changes as $t$ increases from $-\infty$ to $\infty$ (or decreases from $\infty$ to $-\infty$). This information is encoded in the persistence diagram, a multiset of points in the plane, each corresponding to the birth and death of a homological feature that existed for
some interval of $t$.

This paper is devoted to the presentation of the R package TDA, which provides a user-friendly interface for the efficient algorithms of the C++ libraries GUDHI (Maria 2014), Dionysus (Morozov 2007), and PHAT (Bauer, Kerber, and Reininghaus 2012).

In Section 2, we describe how to compute some widely studied functions that, starting from a point cloud, provide some topological information about the underlying space: the distance function (distFct), the distance to a measure function (dtm), the k Nearest Neighbor density estimator (knnDE), the kernel density estimator (kde), and the kernel distance (kernelDist). Section 3 is devoted to the computation of persistence diagrams: the function gridDiag can be used to compute persistent homology of sublevel sets (or superlevel sets) of functions evaluated over a grid of points; the function ripsDiag returns the persistence diagram of the Rips filtration built on top of a point cloud.

One of the key challenges in persistent homology is to find a way to isolate the points of the persistence diagram representing the topological noise. Statistical methods for persistent homology provide an alternative to its exact computation. Knowing with high confidence that an approximated persistence diagrams is close to the true—computationally infeasible—diagram is often enough for practical purposes. Fasy et al. (2014), Chazal et al. (2014c), and Chazal et al. (2014a) propose several statistical methods to construct confidence sets for persistence diagrams and other summary functions that allow us to separate topological signal from topological noise. The methods are implemented in the TDA package and described in Section 3.

Finally, the TDA package provides the implementation of an algorithm for density clustering. This method allows us to identify and visualize the spatial organization of the data, without specific knowledge about the data generating mechanism and in particular without any a priori information about the number of clusters. In Section 4, we describe the function clusterTree, that, given a density estimator, encodes the hierarchy of the connected components of its superlevel sets into a dendrogram, the cluster tree (Kpotufe and von Luxburg 2011; Kent 2013).

## 2. Distance Functions and Density Estimators

As a first toy example to using the TDA package, we show how to compute distance functions and density estimators over a grid of points. The setting is the typical one in TDA: a set of points $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^d$ has been sampled from some distribution $P$ and we are interested in recovering the topological features of the underlying space by studying some functions of the data. The following code generates a sample of 400 points from the unit circle and constructs a grid of points over which we will evaluate the functions.

```r
> library("TDA")
> X <- circleUnif(400)
> Xlim <- c(-1.6, 1.6); Ylim <- c(-1.7, 1.7); by <- 0.065
> Xseq <- seq(Xlim[1], Xlim[2], by = by)
> Yseq <- seq(Ylim[1], Ylim[2], by = by)
> Grid <- expand.grid(Xseq, Yseq)
```

The TDA package provides implementations of the following functions:
• The distance function is defined for each $y \in \mathbb{R}^d$ as $\Delta(y) = \inf_{x \in X} \|x - y\|_2$ and is computed for each point of the Grid with the following code:

```r
> distance <- distFct(X = X, Grid = Grid)
```

• Given a probability measure $P$, the distance to measure (DTM) is defined for each $y \in \mathbb{R}^d$ as

$$d_{m0}(y) = \left( \frac{1}{m0} \int_0^{m0} (G_y^{-1}(u))^r du \right)^{1/r},$$

where $G_y(t) = P(\|X - y\| \leq t)$, and $m0 \in (0,1)$ and $r \in [1,\infty)$ are tuning parameters. As $m0$ increases, DTM function becomes smoother, so $m0$ can be understood as a smoothing parameter. $r$ affects less but also changes DTM function as well. The default value of $r$ is 2. The DTM can be seen as a smoothed version of the distance function. See (Chazal, Cohen-Steiner, and Mérigot 2011, Definition 3.2) and (Chazal, Massart, and Michel 2015, Equation (2)) for a formal definition of the "distance to measure" function.

Given $X = \{x_1, \ldots, x_n\}$, the empirical version of the DTM is

$$\hat{d}_{m0}(y) = \left( \frac{1}{k} \sum_{i \in N_k(y)} \|x_i - y\|^r \right)^{1/r},$$

where $k = \lceil m0 \cdot n \rceil$ and $N_k(y)$ is the set containing the $k$ nearest neighbors of $y$ among $x_1, \ldots, x_n$.

For more details, see (Chazal et al. 2011) and (Chazal et al. 2015).

The DTM is computed for each point of the Grid with the following code:

```r
> m0 <- 0.1
> DTM <- dtm(X = X, Grid = Grid, m0 = m0)
```

• The $k$ Nearest Neighbor density estimator, for each $y \in \mathbb{R}^d$, is defined as

$$\hat{\delta}_k(y) = \frac{k}{n \cdot v_d \cdot r^d_k(y)},$$

where $v_n$ is the volume of the Euclidean $d$ dimensional unit ball and $r^d_k(x)$ is the Euclidean distance from point $x$ to its $k$th closest neighbor among the points of $X$. It is computed for each point of the Grid with the following code:

```r
> k <- 60
> kNN <- knnDE(X = X, Grid = Grid, k = k)
```
The Gaussian Kernel Density Estimator (KDE), for each \( y \in \mathbb{R}^d \), is defined as
\[
\hat{p}_h(y) = \frac{1}{n(\sqrt{2\pi}h)^d} \sum_{i=1}^{n} \exp \left( \frac{-\|y - x_i\|^2}{2h^2} \right).
\]
where \( h \) is a smoothing parameter. It is computed for each point of the Grid with the following code:

```r
> h <- 0.3
> KDE <- kde(X = X, Grid = Grid, h = h)
```

The Kernel distance estimator, for each \( y \in \mathbb{R}^d \), is defined as
\[
\hat{\kappa}_h(y) = \sqrt{\frac{1}{n^2} \sum_{i=1}^{n} \sum_{j=1}^{n} K_h(x_i, x_j) + K_h(y, y) - 2 \frac{1}{n} \sum_{i=1}^{n} K_h(y, x_i)},
\]
where \( K_h(x, y) = \exp \left( \frac{-\|x-y\|^2}{2h^2} \right) \) is the Gaussian Kernel with smoothing parameter \( h \). The Kernel distance is computed for each point of the Grid with the following code:

```r
> h <- 0.3
> Kdist <- kernelDist(X = X, Grid = Grid, h = h)
```

For this 2 dimensional example, we can visualize the functions using `persp` from the `graphics` package. For example the following code produces the KDE plot in Figure 1:

```r
> persp(Xseq, Yseq, +     matrix(KDE, ncol = length(Yseq), nrow = length(Xseq)), xlab = "", +     ylab = "", zlab = "", theta = -20, phi = 35, ltheta = 50, +     col = 2, border = NA, main = "KDE", d = 0.5, scale = FALSE, +     expand = 3, shade = 0.9)
```
2.1. Bootstrap Confidence Bands

We can construct a \((1 - \alpha)\) confidence band for a function using the bootstrap algorithm, which we briefly describe using the kernel density estimator:

1. Given a sample \(X = \{x_1, \ldots, x_n\}\), compute the kernel density estimator \(\hat{p}_h\);
2. Draw \(X^* = \{x^*_1, \ldots, x^*_n\}\) from \(X = \{x_1, \ldots, x_n\}\) (with replacement), and compute \(\theta^* = \sqrt{n}\|\hat{p}_h(x) - \hat{p}_h(x)\|_{\infty}\), where \(\hat{p}_h^*\) is the density estimator computed using \(X^*\);
3. Repeat the previous step \(B\) times to obtain \(\theta_1^*, \ldots, \theta_B^*\);
4. Compute \(q_\alpha = \inf \left\{ q : \frac{1}{B} \sum_{j=1}^{B} I(\theta_j^* \geq q) \leq \alpha \right\} ;
5. The \((1 - \alpha)\) confidence band for \(E[\hat{p}_h]\) is \(\left[ \hat{p}_h - \frac{q_\alpha}{\sqrt{n}}, \hat{p}_h + \frac{q_\alpha}{\sqrt{n}} \right]\).

*Fasy et al. (2014)* and *Chazal et al. (2014a)* prove the validity of the bootstrap algorithm for kernel density estimators, distance to measure, and kernel distance, and use it in the framework of persistent homology. The bootstrap algorithm is implemented in the function *bootstrapBand*, which provides the option of parallelizing the algorithm (\texttt{parallel} = \texttt{TRUE}) using the package *parallel*. The following code computes a 90\% confidence band for \(E[\hat{p}_h]\), showed in Figure 2.
> band <- bootstrapBand(X = X, FUN = kde, Grid = Grid, B = 100,
+     parallel = FALSE, alpha = 0.1, h = h)

Figure 2: the 90% confidence band for $\mathbb{E}[\hat{p}_h]$ has the form $[\ell, u] = [\hat{p}_h - q_\alpha / \sqrt{n}, \hat{p}_h + q_\alpha / \sqrt{n}]$. The plot on the right shows a section of the functions: the red surface is the KDE $\hat{p}_h$; the pink surfaces are $\ell$ and $u$. 
3. Persistent Homology

We provide an informal description of the implemented methods of persistent homology. We assume the reader is familiar with the basic concepts and, for a rigorous exposition, we refer to the textbook Edelsbrunner and Harer (2010).

3.1. Persistent Homology Over a Grid

In this section, we describe how to use the `gridDiag` function to compute the persistent homology of sublevel (and superlevel) sets of the functions described in Section 2. The function `gridDiag` evaluates a given real valued function over a triangulated grid, constructs a filtration of simplices using the values of the function, and computes the persistent homology of the filtration. From version 1.2, `gridDiag` works in arbitrary dimension. The core of the function is written in C++ and the user can choose to compute persistence diagrams using either the C++ library GUDHI, Dionysus, or PHAT.

The following code computes the persistent homology of the superlevel sets (sublevel = FALSE) of the kernel density estimator (FUN = kde, h = 0.3) using the point cloud stored in the matrix X from the previous example. The same code would work for the other functions defined in Section 2 (it is sufficient to replace kde and its smoothing parameter h with another function and the corresponding parameter). The function `gridDiag` returns an object of the class "diagram". The other inputs are the features of the grid over which the kde is evaluated (lim and by), the smoothing parameter h, and a logical variable that indicates whether a progress bar should be printed (printProgress).

```r
> DiagGrid <- gridDiag(  
+   X = X, FUN = kde, h = 0.3, lim = cbind(Xlim, Ylim), by = by,  
+   sublevel = FALSE, library = "Dionysus", location = TRUE,  
+   printProgress = FALSE)
```

We plot the data and the diagram, using the function `plot`, implemented as a standard S3 method for objects of the class "diagram". The following command produces the third plot in Figure 3.

```r
> plot(DiagGrid["diagram"], band = 2 * band["width"],  
+      main = "KDE Diagram")
```

The option (band = 2 * band["width"]) produces a pink confidence band for the persistence diagram, using the confidence band constructed for the corresponding kernel density estimator in the previous section. The features above the band can be interpreted as representing significant homological features, while points in the band are not significantly different from noise. The validity of the bootstrap confidence band for persistence diagrams of KDE, DTM, and Kernel Distance derive from the Stability Theorem (Chazal, de Silva, Glisse, and Oudot 2012) and is discussed in detail in Fasy et al. (2014) and Chazal et al. (2014a).

The function `plot` for the class "diagram" provide the options of rotating the diagram (rotated = TRUE), drawing the barcode in place of the diagram (barcode = TRUE), as well as other standard graphical options. See Figure 4.
3.2. Rips Diagrams

The Vietoris-Rips complex $R(X, \varepsilon)$ consists of simplices with vertices in $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^d$ and diameter at most $\varepsilon$. In other words, a simplex $\sigma$ is included in the complex if each pair of vertices in $\sigma$ is at most $\varepsilon$ apart. The sequence of Rips complexes obtained by gradually increasing the radius $\varepsilon$ creates a filtration.

The \texttt{ripsDiag} function computes the persistence diagram of the Rips filtration built on top of a point cloud. The user can choose to compute the Rips filtration using either the C++ library \texttt{GUDHI} or \texttt{Dionysus}. Then for computing the persistence diagram from the Rips filtration, the user can use either the C++ library \texttt{GUDHI}, \texttt{Dionysus}, or \texttt{PHAT}.

The following code generates 60 points from two circles:

\begin{equation}
\end{equation}

Figure 3: The plot on the right shows the persistence diagram of the superlevel sets of the KDE. Black points represent connected components and red triangles represent loops. The features are born at high levels of the density and die at lower levels. The pink 90\% confidence band separates significant features from noise.
> par(mfrow = c(1, 2), mai = c(0.8, 0.8, 0.3, 0.1))
> plot(DiagGrid[['diagram']], rotated = TRUE, band = band[['width']],
+       main = "Rotated Diagram")
> plot(DiagGrid[['diagram']], barcode = TRUE, main = "Barcode")

![Rotated Diagram and Barcode](image)

Figure 4: Rotated Persistence Diagram and Barcode

> Circle1 <- circleUnif(60)
> Circle2 <- circleUnif(60, r = 2) + 3
> Circles <- rbind(Circle1, Circle2)

We specify the limit of the Rips filtration and the max dimension of the homological features we are interested in (0 for components, 1 for loops, 2 for voids, etc.):

> maxscale <- 5  # limit of the filtration
> maxdimension <- 1  # components and loops

and we generate the persistence diagram:

> DiagRips <- ripsDiag(X = Circles, maxdimension, maxscale,
+                       library = c("GUDHI", "Dionysus"), location = TRUE, printProgress = FALSE)

Alternatively, using the option (dist = "arbitrary") in ripsDiag(), the input X can be an n × n matrix of distances. This option is useful when the user wants to consider a Rips filtration constructed using an arbitrary distance and is currently only available for the option (library = "Dionysus").

Finally we plot the data and the diagram, as in Figure 5:

### 3.3. Alpha Complex Persistence Diagram

For a finite set of points X ⊂ ℜ^d, the Alpha complex Alpha(X, s) is a simplicial subcomplex of the Delaunay complex of X consisting of simplices of circumradius less than or equal to √s.
For each $u \in X$, let $V_u$ be its Voronoi cell, i.e. $V_u = \{ x \in \mathbb{R}^d : d(x, u) \leq d(x, v) \text{ for all } v \in X \}$, and $B_u(r)$ be the closed ball with center $u$ and radius $r$. Let $R_u(r)$ consists of be the intersection of each ball of radius $r$ with the Voronoi cell of $u$, i.e. $R_u(r) = B_u(r) \cap V_u$. Then $\text{Alpha}(X, s)$ is defined as

$$
\text{Alpha}(X, r) = \left\{ \sigma \subset X : \bigcap_{u \in \sigma} R_u(\sqrt{s}) \neq \emptyset \right\}.
$$

See (Edelsbrunner and Harer 2010, Section 3.4) and (Rouvreau 2015). The sequence of Alpha complexes obtained by gradually increasing the parameter $s$ creates an Alpha complex filtration.

The \texttt{alphaComplexDiag} function computes the Alpha complex filtration built on top of a point cloud, using the C++ library \texttt{GUDHI}. Then for computing the persistence diagram from the Alpha complex filtration, the user can use either the C++ library \texttt{GUDHI}, \texttt{Dionysus}, or \texttt{PHAT}.

We first generate 30 points from a circle:

```r
> X <- circleUnif(n = 30)
```

and the following code compute the persistence diagram of the alpha complex filtration using the point cloud $X$, with printing its progress (\texttt{printProgress = FALSE}). The function \texttt{alphaComplexDiag} returns an object of the class "diagram".

```r
> # persistence diagram of alpha complex
> DiagAlphaCmplx <- alphaComplexDiag(
+   X = X, library = c("GUDHI", "Dionysus"), location = TRUE,
+   printProgress = TRUE)
>
# Generated complex of size: 115
0% 10 20 30 40 50 60 70 80 90 100%
|----|----|----|----|----|----|----|----|----|----|
# Persistence timer: Elapsed time [ 0.000000 ] seconds

And we plot the diagram:

```r
> # plot
> par(mfrow = c(1, 2))
> plot(DiagAlphaCmplx[["diagram"]], main = "Alpha complex persistence diagram")
> one <- which(DiagAlphaCmplx[["diagram"]][, 1] == 1)
> one <- one[which.max(
+   DiagAlphaCmplx[["diagram"]][one, 3] - DiagAlphaCmplx[["diagram"]][one, 2])]
> plot(X, col = 1, main = "Representative loop")
> for (i in seq(along = one)) {
+   for (j in seq_len(dim(DiagAlphaCmplx[["cycleLocation"]]][[one[i]]][1]))) {
+     lines(DiagAlphaCmplx[["cycleLocation"]]][[one[i]]][j, , ], pch = 19,
+     cex = 1, col = i + 1)
+   }
+ }
> par(mfrow = c(1, 1))
```

Figure 6: Persistence diagram of Alpha complex. Black points represent connected components and red triangles represent loops.

### 3.4. Persistence Diagram of Alpha Shape

The **Alpha shape complex** $S(X, \alpha)$ is the polytope with its boundary consisting of $\alpha$-exposed simplices, where a simplex $\sigma$ is $\alpha$-exposed if there is an open ball $b$ of radius $\alpha$ such that $b \cap X = \emptyset$ and $\partial b \cap X = \sigma$. Suppose $\mathbb{R}^d$ is filled with ice cream, then consider scooping out the ice cream with sphere-shaped spoon of radius $\alpha$ without touching the points $X$. $S(X, \alpha)$ is the remaining polytope with straightening round surfaces. See (Fischer 2005) and (Edelsbrunner and Mücke 1994). The sequence of Alpha shape complexes obtained by gradually increasing the parameter $\alpha$ creates an Alpha shape complex filtration.

The `alphaShapeDiag` function computes the persistence diagram of the Alpha shape filtration built on top of a point cloud in 3 dimension, using the C++ library GUDHI. Then for
computing the persistence diagram from the Alpha shape filtration, the user can use either the C++ library GUDHI, Dionysus, or PHAT. Currently the point data cloud should lie in 3 dimension.

We first generate 500 points from a sphere:

> XX <- sphereUnif(n = 500, d = 2)

and the following code compute the persistence diagram of the alpha shape filtration using the point cloud XX, with printing its progress (printProgress = FALSE). The function alphaShapeDiag returns an object of the class "diagram".

> DiagAlphaShape <- alphaShapeDiag(X = XX, printProgress = FALSE)

And we plot the diagram:

> plot(DiagAlphaShape["diagram"], diagLim = c(0, 1))

Figure 7: Persistence diagram of Alpha shape. Black points represent connected components, red triangles represent loops, and blue diamonds represent voids.

3.5. Persistence Diagrams from Filtration

Rather than computing persistence diagrams from built-in function, it is also possible to compute persistence diagrams from a user-defined filtration. A filtration consists of simplicial complex and the filtration values on each simplex. The functionsripsDiag, alphaComplexDiag, alphaShapeDiag have their counterparts for computing corresponding filtrations instead of persistence diagrams: namely,ripsFiltration corresponds to the Rips filtration built on top of a point cloud, alphaComplexFiltration to the alpha complex filtration, and alphaShapeFiltration to the alpha shape filtration.

We first generate 30 points from a circle:

> X <- circleUnif(n = 100)
Then, after specifying the limit of the Rips filtration and the max dimension of the homological features, the following code compute the Rips filtration using the point cloud $X$.

```r
> maxscale <- 0.4 # limit of the filtration
> maxdimension <- 1 # components and loops
> FltRips <- ripsFiltration(X = X, maxdimension = maxdimension,
+ maxscale = maxscale, dist = "euclidean", library = "GUDHI",
+ printProgress = TRUE)
# Generated complex of size: 2476
```

One way of defining a user-defined filtration is to build a filtration from a simplicial complex and function values on the vertices. The function `funFiltration` takes function values (`FUNvalues`) and simplicial complex (`cmplx`) as input, and build a filtration, where a filtration value on a simplex is defined as the maximum of function values on the vertices of the simplex.

In the following example, the function `funFiltration` construct a filtration from a Rips complex and the DTM function values on data points.

```r
> m0 <- 0.1
> dtmValues <- dtm(X = X, Grid = X, m0 = m0)
> FltFun <- funFiltration(FUNvalues = dtmValues, cmplx = FltRips[["cmplx"]])
```

Once the filtration is computed, the function `filtrationDiag` computes the persistence diagram from the filtration. The user can choose to compute the persistence diagram using either the C++ library `GUDHI` or `PHAT`.

```r
> DiagFltFun <- filtrationDiag(filtration = FltFun, maxdimension = maxdimension,
+ library = "Dionysus", location = TRUE, printProgress = TRUE)
```

Then we plot the data and the diagram:

### 3.6. Bottleneck and Wasserstein Distances

Standard metrics for measuring the distance between two persistence diagrams are the bottleneck distance and the $p$th Wasserstein distance (Edelsbrunner and Harer 2010). The TDA package includes the functions `bottleneck` and `wasserstein`, which are R wrappers of the functions “bottleneck_distance” and “wasserstein_distance” of the C++ library `Dionysus`.

We generate two persistence diagrams of the Rips filtrations built on top of the two (separate) circles of the previous example.

```r
> Diag1 <- ripsDiag(Circle1, maxdimension = 1, maxscale = 5)
> Diag2 <- ripsDiag(Circle2, maxdimension = 1, maxscale = 5)
```
and we compute the bottleneck distance and the 2nd Wasserstein distance between the two diagrams. In the following code, the option \texttt{dimension} = 1 specifies that the distances between diagrams are computed using only one dimensional features (loops).

\begin{verbatim}
> print(bottleneck(Diag1["diagram"], Diag2["diagram"],
+   dimension = 1))
[1] 1.161506
> print(wasserstein(Diag1["diagram"], Diag2["diagram"], p = 2,
+   dimension = 1))
[1] 1.770667
\end{verbatim}

3.7. Landscapes and Silhouettes

Persistence landscapes and silhouettes are real-valued functions that further summarize the information contained in a persistence diagram. They have been introduced and studied in Bubenik (2012), Chazal et al. (2014c), and Chazal, Fasy, Lecci, Michel, Rinaldo, and Wasserman (2014b). We briefly introduce the two functions.

\textbf{Landscape.} The persistence landscape is a collection of continuous, piecewise linear functions \( \lambda : \mathbb{Z}^+ \times \mathbb{R} \to \mathbb{R} \) that summarizes a persistence diagram. To define the landscape, consider the set of functions created by tenting each each point \( p = (x, y) = (b+\frac{d}{2}, d-b) \) representing a birth-death pair \((b, d)\) in the persistence diagram \( D \) as follows:

\[
\Lambda_p(t) = \begin{cases} 
  t - x + y & t \in [x - y, x] \\
  x + y - t & t \in [x, x + y] \\
  0 & \text{otherwise}
\end{cases}
\]

(1)
We obtain an arrangement of piecewise linear curves by overlaying the graphs of the functions $\{\Lambda_p\}_p$; see Figure 9 (left). The persistence landscape of $D$ is a summary of this arrangement. Formally, the persistence landscape of $D$ is the collection of functions

$$\lambda(k, t) = k\max_p \Lambda_p(t), \quad t \in [0, T], k \in \mathbb{N},$$

where $k\max$ is the $k$th largest value in the set; in particular, $1\max$ is the usual maximum function. see Figure 9 (middle).

**Silhouette.** Consider a persistence diagram with $N$ off diagonal points $\{(b_j, d_j)\}_{j=1}^N$. For every $0 < p < \infty$ we define the power-weighted silhouette

$$\phi^{(p)}(t) = \frac{\sum_{j=1}^N |d_j - b_j|^p \Lambda_j(t)}{\sum_{j=1}^N |d_j - b_j|^p}.$$

The value $p$ can be thought of as a trade-off parameter between uniformly treating all pairs in the persistence diagram and considering only the most persistent pairs. Specifically, when $p$ is small, $\phi^{(p)}(t)$ is dominated by the effect of low persistence features. Conversely, when $p$ is large, $\phi^{(p)}(t)$ is dominated by the most persistent features; see Figure 9 (right).

Figure 9: Left: we use the rotated axes to represent a persistence diagram $D$. A feature $(b, d) \in D$ is represented by the point $(\frac{b+d}{2}, \frac{d-b}{2})$ (pink). In words, the $x$-coordinate is the average parameter value over which the feature exists, and the $y$-coordinate is the half-life of the feature. Middle: the blue curve is the landscape $\lambda(1, \cdot)$. Right: the blue curve is the silhouette $\phi^{(1)}(\cdot)$.

The landscape and silhouette functions can be evaluated over a one-dimensional grid of points $tseq$ using the functions `landscape` and `silhouette`. In the following code, we use the persistence diagram from Figure 5 to construct the corresponding landscape and silhouette for one-dimensional features ($\text{dimension} = 1$). The option ($\text{KK} = 1$) specifies that we are interested in the 1st landscape function, and ($\text{p} = 1$) is the power of the weights in the definition of the silhouette function.

```R
> maxscale <- 5
> tseq <- seq(0, maxscale, length = 1000)  # domain
> Land <- landscape(DiagRips[["diagram"]], dimension = 1, KK = 1, tseq)
> Sil <- silhouette(DiagRips[["diagram"]], p = 1, dimension = 1, tseq)
```
The functions `landscape` and `silhouette` return real valued vectors, which can be simply plotted with `plot(tseq, Land, type = "l"); plot(tseq, Sil, type = "l"). See Figure 10.

![Figure 10: Landscape and Silhouette of the one-dimensional features of the diagram of Figure 5.](image)

3.8. Confidence Bands for Landscapes and Silhouettes

Recent results in Chazal et al. (2014c) and Chazal et al. (2014b) show how to construct confidence bands for landscapes and silhouettes, using a bootstrap algorithm (multiplier bootstrap). This strategy is useful in the following scenario. We have a very large dataset with \( N \) points. There is a diagram \( D \) and landscape \( \lambda \) corresponding to some filtration built on the data. When \( N \) is large, computing \( D \) is prohibitive. Instead, we draw \( n \) subsamples, each of size \( m \). We compute a diagram and a landscape for each subsample yielding landscapes \( \lambda_1, \ldots, \lambda_n \). (Assuming \( m \) is much smaller than \( N \), these subsamples are essentially independent and identically distributed.) Then we compute \( \frac{1}{n} \sum_i \lambda_i \), an estimate of \( E(\lambda_i) \), which can be regarded as an approximation of \( \lambda \). The function `multipBootstrap` uses the landscapes \( \lambda_1, \ldots, \lambda_n \) to construct a confidence band for \( E(\lambda_i) \). The same strategy is valid for silhouette functions. We illustrate the method with a simple example.

First we sample \( N \) points from two circles:

```r
> N <- 4000
> XX1 <- circleUnif(N / 2)
> XX2 <- circleUnif(N / 2, r = 2) + 3
> X <- rbind(XX1, XX2)
```

Then we specify the number of subsamples \( n \), the subsample size \( m \), and we create the objects that will store the \( n \) diagrams and landscapes:

```r
> m <- 80   # subsample size
> n <- 10   # we will compute n landscapes using subsamples of size m
> tseq <- seq(0, maxscale, length = 500)   #domain of landscapes
> #here we store n Rips diags
> Diags <- list()
> #here we store n landscapes
> Lands <- matrix(0, nrow = n, ncol = length(tseq))
```
For $n$ times, we subsample from the large point cloud, compute $n$ Rips diagrams and the corresponding 1st landscape functions ($\text{KK} = 1$), using 1 dimensional features ($\text{dimension} = 1$):

```r
> for (i in seq_len(n)) {
+   subX <- X[sample(seq_len(N), m), ]
+   Diags[[i]] <- ripsDiag(subX, maxdimension = 1, maxscale = 5)
+   Lands[i, ] <- landscape(Diags[[i]]["diagram"], dimension = 1,
+                             KK = 1, tseq)
+ }
```

Finally we use the $n$ landscapes to construct a 95% confidence band for the mean landscape

```r
> bootLand <- multipBootstrap(Lands, B = 100, alpha = 0.05,
+                             parallel = FALSE)
```

which is plotted by the following code. See Figure 11.

```r
> plot(tseq, bootLand["mean"], main = "Mean Landscape with 95% band")
> polygon(c(tseq, rev(tseq)),
+         c(bootLand["band"], 1, rev(bootLand["band"], 2)),
+         col = "pink")
> lines(tseq, bootLand["mean"], lwd = 2, col = 2)
```

![Figure 11: 95% confidence band for the mean landscape function.](image)

### 3.9. Selection of Smoothing Parameters

An unsolved problem in topological inference is how to choose the smoothing parameters, for example $h$ for KDE and $m_0$ for DTM.

Chazal et al. (2014a) suggest the following method, that we describe here for the kernel density estimator, but works also for the kernel distance and the distance to measure.
Let $\ell_1(h), \ell_2(h), \ldots$, be the lifetimes of the features of a persistence diagram at scale $h$. Let $q_\alpha(h)/\sqrt{n}$ be the width of the confidence band for the kernel density estimator at scale $h$, as described in Section 2.1. We define two quantities that measure the amount of significant information at level $h$:

- The number of significant features, $N(h) = \#\{i : \ell(i) > 2q_\alpha(h)/\sqrt{n}\}$;
- The total significant persistence, $S(h) = \sum_i [\ell_i - 2q_\alpha(h)/\sqrt{n}]_+$. 

These measures are small when $h$ is small since $q_\alpha(h)$ is large. On the other hand, they are small when $h$ is large since then all the features of the KDE are smoothed out. Thus we have a kind of topological bias-variance tradeoff. We choose $h$ to maximize $N(h)$ or $S(h)$.

The method is implemented in the function `maxPersistence`, as shown in the following toy example. First, we sample 1600 point from two circles (plus some clutter noise) and we specify the limits of the grid over which the KDE is evaluated:

```r
> XX1 <- circleUnif(600)
> XX2 <- circleUnif(1000, r = 1.5) + 2.5
> noise <- cbind(runif(80, -2, 5), runif(80, -2, 5))
> X <- rbind(XX1, XX2, noise)
> # Grid limits
> Xlim <- c(-2, 5)
> Ylim <- c(-2, 5)
> by <- 0.2
```

Then we specify a sequence of smoothing parameters among which we will select the optimal one, the number of bootstrap iterations and the level of the confidence bands to be computed:

```r
> parametersKDE <- seq(0.1, 0.6, by = 0.05)
> B <- 50  # number of bootstrap iterations. Should be large.
> alpha <- 0.1  # level of the confidence bands
```

The function `maxPersistence` can be parallelized (`parallel = TRUE`) and a progress bar can be printed (`printProgress = TRUE`):

```r
> maxKDE <- maxPersistence(kde, parametersKDE, X,
+     lim = cbind(Xlim, Ylim), by = by, sublevel = FALSE,
+     B = B, alpha = alpha, parallel = TRUE,
+     printProgress = TRUE, bandFUN = "bootstrapBand")
```

Then we specify a sequence of smoothing parameters among which we will select the optimal one, the number of bootstrap iterations and the level of the confidence bands to be computed:

```r
> parametersKDE <- seq(0.1, 0.6, by = 0.05)
> B <- 50  # number of bootstrap iterations. Should be large.
> alpha <- 0.1  # level of the confidence bands
```

The function `maxPersistence` can be parallelized (`parallel = TRUE`) and a progress bar can be printed (`printProgress = TRUE`):

```r
> maxKDE <- maxPersistence(kde, parametersKDE, X,
+     lim = cbind(Xlim, Ylim), by = by, sublevel = FALSE,
+     B = B, alpha = alpha, parallel = TRUE,
+     printProgress = TRUE, bandFUN = "bootstrapBand")
```

The `S3` methods `summary` and `plot` are implemented for the class "maxPersistence". We can display the values of the parameters that maximize the two criteria:
> print(summary(maxKDE))

Call:
maxPersistence(FUN = kde, parameters = parametersKDE, X = X, 
lim = cbind(Xlim, Ylim), by = by, sublevel = FALSE, B = B, 
alpha = alpha, bandFUN = "bootstrapBand", parallel = TRUE, 
printProgress = TRUE)

The number of significant features is maximized by
[1] 0.25 0.30

The total significant persistence is maximized by
[1] 0.3

and produce the summary plot of Figure 12.

> par(mfrow = c(1, 2), mai = c(0.8, 0.8, 0.35, 0.3))
> plot(X, pch = 16, cex = 0.5, main = "Two Circles")
> plot(maxKDE, main = "Max Persistence – KDE")

Figure 12: Max Persistence Method for the selection of smoothing parameters. For each value of the smoothing parameter we display the persistence of the corresponding homological features, along with a (pink) confidence band that separates the statistically significant features from the topological noise.

4. Density Clustering

The last example of this vignette illustrates the use of the function `clusterTree`, which is an implementation of Algorithm 1 in Kent, Rinaldo, and Verstynen (2013).

First, we briefly describe the task of density clustering; we defer the reader to Kent (2013) for a more rigorous and complete description. Let $f$ be the density of the probability distribution $P$
generating the observed sample \( X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^d \). For a threshold value \( \lambda > 0 \), the corresponding super level set of \( f \) is \( \mathcal{L}_f(\lambda) := \text{cl}\{x \in \mathbb{R}^d : f(x) > \lambda\} \), and its \( d \)-dimensional subsets are called high-density regions. The high-density clusters of \( P \) are the maximal connected subsets of \( \mathcal{L}_f(\lambda) \). By considering all the level sets simultaneously (from \( \lambda = 0 \) to \( \lambda = \infty \)), we can record the evolution and the hierarchy of the high-density clusters of \( P \). This naturally leads to the notion of the cluster density tree of \( P \) (see, e.g., Hartigan (1981)), defined as the collection of sets \( \mathcal{T} := \{\mathcal{L}_f(\lambda), \lambda \geq 0\} \), which satisfies the tree property: \( A, B \in \mathcal{T} \) implies that \( A \subset B \) or \( B \subset A \) or \( A \cap B = \emptyset \). We will refer to this construction as the \( \lambda \)-tree. Alternatively, Kent et al. (2013) introduced the \( \alpha \)-tree and \( \kappa \)-tree, which facilitate the interpretation of the tree by precisely encoding the probability content of each tree branch rather than the density level. Cluster trees are particularly useful for high dimensional data, whose spatial organization is difficult to represent.

We illustrate the strategy with a simple example. First we generate a 2D point cloud from three (not so well) separated clusters (see top left plot of Figure 13):

```r
> X1 <- cbind(rnorm(300, 1, .8), rnorm(300, 5, 0.8))
> X2 <- cbind(rnorm(300, 3.5, .8), rnorm(300, 5, 0.8))
> X3 <- cbind(rnorm(300, 6, 1), rnorm(300, 1, 1))
> XX <- rbind(X1, X2, X3)
```

Then we use the function `clusterTree` to compute cluster trees using the k Nearest Neighbors density estimator (\( k = 100 \) nearest neighbors) and the Gaussian kernel density estimator, with smoothing parameter \( h \).

```r
> Tree <- clusterTree(XX, k = 100, density = "knn",
+    printProgress = FALSE)
> TreeKDE <- clusterTree(XX, k = 100, h = 0.3, density = "kde",
+    printProgress = FALSE)
```

Note that, even when \( \text{kde} \) is used to estimate the density, we have to provide the option (\( k = 100 \)), so that the algorithm can compute the connected components at each level of the density using a \( k \) Nearest Neighbors graph.

The "clusterTree" objects `Tree` and `TreeKDE` contain information about the \( \lambda \)-tree, \( \alpha \)-tree and \( \kappa \)-tree. The function `plot` for objects of the class "clusterTree" produces the plots in Figure 13.

```r
> plot(Tree, type = "lambda", main = "lambda Tree (knn)")
> plot(Tree, type = "kappa", main = "kappa Tree (knn)")
> plot(TreeKDE, type = "lambda", main = "lambda Tree (kde)")
> plot(TreeKDE, type = "kappa", main = "kappa Tree (kde)")
```
Figure 13: The lambda trees and kappa trees of the k Nearest Neighbor density estimator and the kernel density estimator.

References


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