Tutorial on the R package TDA

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Abstract

This tutorial gives an introduction to the R package **TDA**, which provides some tools for Topological Data Analysis. The salient topological features of data can be quantified with persistent homology. The R package **TDA** provide an R interface for the efficient algorithms of the C++ libraries **GUDHI**, **Dionysus**, and **PHAT**. Specifically, The R package **TDA** includes functions for computing the persistent homology of the Rips complex, alpha complex, and alpha shape complex, and a function for the persistent homology of sublevel sets (or superlevel sets) of arbitrary functions evaluated over a grid of points. The R package **TDA** also provides a function for computing the confidence band that determines the significance of the features in the resulting persistence diagrams.

Keywords: Topological Data Analysis, Persistent Homology.

1. Introduction

R(http://cran.r-project.org/) is a programming language for statistical computing and graphics.

R has several good properties: R has many packages for statistical computing. Also, R is easy to make (interactive) plots. R is a script language, and it is easy to use. But, R is slow. C or C++ stands on the opposite end: C or C++ also has many packages(or libraries). But, C or C++ is difficult to make plots. C or C++ is a compiler language, and is difficult to use. But, C or C++ is fast. In short, R has short development time but long execution time, and C or C++ has long development time but short execution time.

Several libraries are developed for Topological Data Analysis: for example, GUDHI(https://project.inria.fr/gudhi/software/), Dionysus(http://www.mrzv.org/software/dionysus/), and PHAT(https://code.google.com/p/phat/). They are all written in C++, since Topological Data Analysis is computationally heavy and R is not fast enough.

R package TDA(http://cran.r-project.org/web/packages/TDA/index.html) bridges between C++ libraries(GUDHI, Dionysus, PHAT) and R. TDA package provides an R interface for the efficient algorithms of the C++ libraries GUDHI, Dionysus and PHAT. So by using TDA package, short development time and short execution time can be both achieved.

R package **TDA** provides tools for Topological Data Analysis. You can compute several different things with **TDA** package: you can compute common distance functions and density estimators, the persistent homology of the Rips filtration, the persistent homology of sublevel sets of a function over a grid, the confidence band for the persistence diagram, and the cluster density trees for density clustering.

2. Installation

First, you should download R. R of version at least 3.1.0 is required:

http://cran.r-project.org/bin/windows/base/ (for Windows)

http://cran.r-project.org/bin/macosx/ (for (Mac) OS X)

R is part of many Linux distributions, so you should check with your Linux package management system.

You can use whatever IDE that you would like to use(Rstudio, Eclipse, Emacs, Vim...). R itself also provides basic GUI or CUI. I personally use Rstudio:

http://www.rstudio.com/products/rstudio/download/

For Windows and Mac, you can install R package **TDA** as in the following code (or pushing 'Install R packages' button if you use Rstudio).

Loading required package: TDA

If you are using Linux, you should install R package **TDA** from the source. To do this, you need to install two libraries in advance: gmp (https://gmplib.org/) and mpfr (http://www.mpfr.org/). Installation of these packages may differ by your Linux distributions. Once those libraries are installed, you need to install four R packages: **parallel**, **FNN**, **igraph**, and **scales**. **parallel** is included when you install R, so you need to install **FNN**, **igraph**, and **scales** by yourself. You can install them by following code (or pushing 'Install R packages' button if you use Rstudio).

```
# installing required packages
if (!require(package = "FNN")) {
 install.packages(pkgs = "FNN")
}
## Loading required package:
                       FNN
if (!require(package = "igraph")) {
 install.packages(pkgs = "igraph")
}
## Loading required package:
                       igraph
##
## Attaching package:
                  'igraph'
## The following object is masked from 'package:FNN':
##
##
    knn
## The following objects are masked from 'package:stats':
##
##
    decompose, spectrum
## The following object is masked from 'package:base':
##
##
    union
```

```
if (!require(package = "scales")) {
    install.packages(pkgs = "scales")
}
## Loading required package: scales
```

Then you can install the ${\sf R}$ package ${\bf TDA}$ as in Windows or Mac:

Once installation is done, R package **TDA** should be loaded as in the following code, before using the package functions.

3. Sample on manifolds, Distance Functions, and Density Estimators

3.1. Uniform Sample on manifolds

A set of n points $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^d$ has been sampled from some distribution P.

• *n* sample from the uniform distribution on the circle in \mathbb{R}^2 with radius *r*.



3.2. Distance Functions, and Density Estimators

We compute distance functions and density estimators over a grid of points. Suppose a set of points $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^d$ has been sampled from some distribution P. 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.

lim <- c(-1.7, 1.7)
by <- 0.05
margin <- seq(from = lim[1], to = lim[2], by = by)
Grid <- expand.grid(margin, margin)</pre>

• The distance function is defined for each $y \in \mathbb{R}^d$ as $\Delta(y) = \inf_{x \in X} ||x - y||_2$.



• The Gaussian Kernel Density Estimator (KDE), for each $y \in \mathbb{R}^d$, is defined as

$$\hat{p}_h(y) = \frac{1}{n(\sqrt{2\pi}h)^d} \sum_{i=1}^n \exp\left(\frac{-\|y-x_i\|_2^2}{2h^2}\right)$$

where h is a smoothing parameter.



4. Persistent Homology

4.1. Persistent Homology Over a Grid

gridDiag function computes the persistent homology of sublevel (and superlevel) sets of the functions. The function gridDiag evaluates a given real valued function over a triangulated grid (in arbitrary dimension), constructs a filtration of simplices using the values of the function, and computes the persistent homology of the filtration. The user can choose to compute persistence diagrams using either the C++ library GUDHI (library = "GUDHI"), Dionysus (library = "Dionysus"), or PHAT (library = "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 other inputs are the features of the grid over which the kde is evaluated (lim and by), and a logical variable that indicates whether a progress bar should be printed (printProgress).

The function plot plots persistence diagram for objects of the class "diagram".

```
xlab = "", ylab = "", zlab = "", theta = -20, phi = 35, scale = FALSE,
expand = 3, col = "red", border = NA, ltheta = 50, shade = 0.9,
main = "KDE")
plot(x = DiagGrid[["diagram"]], main = "KDE Diagram")
```



4.2. Rips Persistent Homology

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 ε . In other words, a simplex σ is included in the complex if each pair of vertices in σ is at most ε apart. The sequence of Rips complexes obtained by gradually increasing the radius ε creates a filtration.

The 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 **GUDHI** or **Dionysus**. Then for computing the persistence diagram from the Rips filtration, the user can use either the C++ library **GUDHI**, **Dionysus**, or **PHAT**.

The following code computes the persistent homology of the Rips filtratio using the point cloud stored in the matrix X from the previous example, and the plot the data and the diagram.

```
par(mfrow = c(1,2))
plot(X, xlab = "", ylab = "", main = "Sample X", pch = 20)
plot(x = DiagRips[["diagram"]], main = "Rips Diagram")
```



4.3. Persistent Homology 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 functions **ripsDiag** has their counterparts for computing corresponding filtrations instead of persistence diagrams: namely, **ripsFiltration** corresponds to the Rips filtration built on top of a point cloud.

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.

```
FltRips <- ripsFiltration(X = X, maxdimension = 1, maxscale = 0.5,
library = "GUDHI")
```

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 kernel density estimates on data points.

```
h <- 0.3
KDEx <- kde(X = X, Grid = X, h = h)
FltFun <- funFiltration(FUNvalues = KDEx, cmplx = FltRips[["cmplx"]],
    sublevel = FALSE)</pre>
```

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 **Dionysus**.



5. Statistical Inference on Persistent Homology

 $(1 - \alpha)$ confidence band can be computed 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 $q_{\alpha} = \inf \left\{ q : \frac{1}{B} \sum_{j=1}^{B} I(\theta_{j}^{*} \ge q) \le \alpha \right\};$
- 5. The (1α) confidence band for $\mathbb{E}[\hat{p}_h]$ is $\left[\hat{p}_h \frac{q_\alpha}{\sqrt{n}}, \hat{p}_h + \frac{q_\alpha}{\sqrt{n}}\right]$.

bootstrapBand computes $(1 - \alpha)$ bootstrap confidence band, with the option of parallelizing the algorithm (parallel=TRUE). The following code computes a 90% confidence band for $\mathbb{E}[\hat{p}_h]$.

Then such confidence band for $\mathbb{E}[\hat{p}_h]$ can be used as the confidence band for the persistent homology.



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