# Package 'TDAstats'

October 12, 2022

```
Title Pipeline for Topological Data Analysis
Version 0.4.1
Description A comprehensive toolset for any
      useR conducting topological data analysis, specifically via the
      calculation of persistent homology in a Vietoris-Rips complex.
      The tools this package currently provides can be conveniently split
      into three main sections: (1) calculating persistent homology; (2)
      conducting statistical inference on persistent homology calculations;
      (3) visualizing persistent homology and statistical inference.
      The published form of TDAstats can be found in Wadhwa et al. (2018)
      <doi:10.21105/joss.00860>.
      For a general background on computing persistent homology for
      topological data analysis, see Otter et al. (2017)
      <doi:10.1140/epjds/s13688-017-0109-5>.
      To learn more about how the permutation test is used for
      nonparametric statistical inference in topological data analysis,
      read Robinson & Turner (2017) <doi:10.1007/s41468-017-0008-7>.
      To learn more about how TDAstats calculates persistent homology,
      you can visit the GitHub repository for Ripser, the software that
      works behind the scenes at <a href="https://github.com/Ripser/ripser">https://github.com/Ripser/ripser</a>.
      This package has been published as Wadhwa et al. (2018)
      <doi:10.21105/joss.00860>.
License GPL-3
Encoding UTF-8
LazyData true
Depends R (>= 3.3)
Imports ggplot2 (>= 2.2.1), Rcpp (>= 0.12.15)
URL https://github.com/rrrlw/TDAstats
BugReports https://github.com/rrrlw/TDAstats/issues
LinkingTo Rcpp
RoxygenNote 6.1.0
```

Type Package

2 calculate\_homology

# SystemRequirements C++11 Suggests testthat (>= 2.0.0), knitr, rmarkdown, covr VignetteBuilder knitr NeedsCompilation yes Author Raoul Wadhwa [aut, cre], Andrew Dhawan [aut], Drew Williamson [aut], Jacob Scott [aut], Jason Cory Brunson [ctb], Shota Ochi [ctb]

Maintainer Raoul Wadhwa < raoulwadhwa@gmail.com>

Repository CRAN

**Date/Publication** 2019-12-12 22:30:03 UTC

# **R** topics documented:

	calculate_homology	2
	circle2d	3
	id_significant	4
	permutation_test	5
	phom.dist	6
	plot_barcode	6
	plot_persist	7
	sphere3d	
	TDAstats	9
	unif2d	9
	unif3d	9
Index		11

calculate\_homology

Calculate Persistent Homology of a Point Cloud

# **Description**

Calculates the persistent homology of a point cloud, as represented by a Vietoris-Rips complex. This function is an R wrapper for Ulrich Bauer's Ripser C++ library for calculating persistent homology. For more information on the C++ library, see <a href="https://github.com/Ripser/ripser">https://github.com/Ripser/ripser</a>.

# Usage

```
calculate_homology(mat, dim = 1, threshold = -1, format = "cloud",
   standardize = FALSE, return_df = FALSE)
```

circle2d 3

# **Arguments**

mat numeric matrix containing point cloud or distance matrix

dim maximum dimension of features to calculate

threshold maximum diameter for computation of Vietoris-Rips complexes

format format of 'mat', either "cloud" for point cloud or "distmat" for distance matrix

standardize boolean determining whether point cloud size should be standardized return\_df defaults to 'FALSE', returning a matrix; if 'TRUE', returns a data frame

# **Details**

The 'mat' parameter should be a numeric matrix with each row corresponding to a single point, and each column corresponding to a single dimension. Thus, if 'mat' has 50 rows and 5 columns, it represents a point cloud with 50 points in 5 dimensions. The 'dim' parameter should be a positive integer. Alternatively, the 'mat' parameter could be a distance matrix (upper triangular half is ignored); note: 'format' should be specified as "ldm".

# Value

3-column matrix or data frame, with each row representing a TDA feature

# **Examples**

```
# create a 2-d point cloud of a circle (100 points)
num.pts <- 100
rand.angle <- runif(num.pts, 0, 2*pi)
pt.cloud <- cbind(cos(rand.angle), sin(rand.angle))
# calculate persistent homology (num.pts by 3 numeric matrix)
pers.hom <- calculate_homology(pt.cloud)</pre>
```

circle2d

2-dimensional point cloud of a unit circle

# **Description**

A dataset containing the Cartesian coordinates of 100 points uniformly distributed on the circumference of a unit circle.

# Usage

circle2d

# Format

A matrix with 100 rows and 2 columns: the x- and y-coordinates

id\_significant

### Source

https://github.com/rrrlw/TDAstats/blob/master/data-raw/circle2d.R

id\_significant

Identify Significant Features in Persistent Homology

# **Description**

An empirical method (bootstrap) to differentiate between features that constitute signal versus noise based on the magnitude of their persistence relative to one another. Note: you must have at least 5 features of a given dimension to use this function.

# Usage

```
id_significant(features, dim = 1, reps = 100, cutoff = 0.975)
```

# **Arguments**

features 3xn data frame of features; the first column must be dimension, the second birth,

and the third death

dim dimension of features of interest

reps number of replicates

cutoff percentile cutoff past which features are considered significant

# **Examples**

permutation\_test 5

permutation_test	Statistical Inference for Topological Data Analysis	
------------------	---	--

# Description

Conducts a permutation test for nonparametric statistical inference of persistent homology in topological data analysis.

# Usage

```
permutation_test(data1, data2, iterations, exponent = 1, update = 0,
    ...)
```

# **Arguments**

data1	first dataset
data2	second dataset
iterations	number of iterations for distribution in permutation test
exponent	parameter 'p' that returns Wasserstein-p metric
update	if greater than zero, will print a message every 'update' iterations
• • •	arguments for 'calculate_homology' used for each permutation; this includes the 'format', 'dim', and 'threshold' parameters

### **Details**

The persistent homology of two point clouds are compared with the Wasserstein metric (where Wasserstein-1 is also known as the Earth Mover's Distance). However, the magnitude of the metric for a single pair of point clouds is meaningless without a reference distribution. This function uses a permutation test (permuting the points between the two clouds) as a nonparametric hypothesis test for statistical inference.

For more details on permutation tests for statistical inference in topological data analysis, see Robinson A, Turner K. Hypothesis testing for topological data analysis. J Appl Comput Topology. 2017; 1(2): 241-261.<a href="https://doi.org/10.1007/s41468-017-0008-7">doi:10.1007/s41468-017-0008-7</a>>

### Value

list containing results of permutation test

6 plot\_barcode

phom.dist Calculate Distance between Homology Matrices
--

# Description

Calculates the distance between two matrices containing persistent homology features, usually as returned by the 'calculate\_homology' function.

# Usage

```
phom.dist(phom1, phom2, limit.num = 0)
```

# **Arguments**

phom1	3-by-n numeric matrix containing persistent homology for first dataset
phom2	3-by-n numeric matrix containing persistent homology for second dataset
limit.num	limit comparison to only top 'limit.num' features in each dimension

# **Details**

Note that the absolute value of this measure of distance is not meaningful without a null distribution or at least another value for relative comparison (e.g. finding most similar pair within a triplet).

# Value

distance vector (1 element per dimension) between 'phom1' and 'phom2'

plot_barcode Plot Persistent Homology as Topological Barcode	plot_barcode	Plot Persistent Homology as Topological Barcode	
--	--------------	---	--

# **Description**

Plots a feature matrix as a topological barcode. See 'plot\_persist' for an alternate visualization method of persistent homology.

# Usage

```
plot_barcode(feature.matrix)
```

# **Arguments**

feature.matrix nx3 matrix representing persistent homology features

plot\_persist 7

### **Details**

The 'feature.matrix' parameter should be a numeric matrix with each row corresponding to a single feature. It should have 3 columns corresponding to feature dimension (col 1), feature birth (col 2), and feature death (col 3). The first column should be filled with integers, and the next two columns should be filled with numeric values. The output from the 'calculate\_homology' function in this package will be a valid value for the 'feature.matrix' parameter.

This function uses the ggplot2 framework to generate persistence diagrams. For details, see: Wickham H (2009). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag: New York, NY.

### Value

ggplot instance representing topological barcode

# **Examples**

```
# create a 2-d point cloud of a circle (100 points)
num.pts <- 100
rand.angle <- runif(num.pts, 0, 2*pi)
pt.cloud <- cbind(cos(rand.angle), sin(rand.angle))
# calculate persistent homology (num.pts by 3 numeric matrix)
pers.hom <- calculate_homology(pt.cloud)
# plot calculated homology features as persistence diagram
plot_barcode(pers.hom)</pre>
```

plot\_persist

Plot Persistent Homology as Persistence Diagram

# **Description**

Plots a feature matrix as a persistence diagram. See 'plot\_barcode' for an alternate visualization method of persistent homology.

# Usage

```
plot_persist(feature.matrix, flat = FALSE, cutoff = 0)
```

# Arguments

```
feature.matrix nx3 matrix representing persistent homology features

flat default FALSE; if TRUE, plots flat persistent homology instead

cutoff threshold for significant features; line added as marker on plot
```

8 sphere3d

### **Details**

The 'feature.matrix' parameter should be a numeric matrix with each row corresponding to a single feature. It should have 3 columns corresponding to feature dimension (col 1), feature birth (col 2), and feature death (col 3). The first column should be filled with integers, and the next two columns should be filled with numeric values. The output from the 'calculate\_homology' function in this package will be a valid value for the 'feature.matrix' parameter.

This function uses the ggplot2 framework to generate persistence diagrams. For details, see: Wickham H (2009, ISBN:9780387981413). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag: New York, NY.

# Value

ggplot instance representing persistence diagram

# **Examples**

```
# create a 2-d point cloud of a circle (100 points)
num.pts <- 100
rand.angle <- runif(num.pts, 0, 2*pi)
pt.cloud <- cbind(cos(rand.angle), sin(rand.angle))
# calculate persistent homology (num.pts by 3 numeric matrix)
pers.hom <- calculate_homology(pt.cloud)
# plot calculated homology features as persistence diagram
plot_persist(pers.hom)</pre>
```

sphere3d

3-dimensional point cloud of a unit sphere

# Description

A dataset containing the Cartesian coordinates of 100 points uniformly distributed on the surface of a unit sphere.

### Usage

sphere3d

### **Format**

A matrix with 100 rows and 3 columns: the x-, y-, and z-coordinates

### Source

https://github.com/rrrlw/TDAstats/blob/master/data-raw/sphere3d.R

TDAstats 9

TDAstats	Statistical Inference for Persistent Homology in Topological Data Analysis

# **Description**

This package aims to be a comprehensive toolset for any useR conducting topological data analysis, specifically via the calculation of persistent homology in a Vietoris-Rips complex. The tools this package currently provides can be conveniently split into three main sections: (1) calculating persistent homology; (2) conducting statistical inference on persistent homology calculations; (3) visualizing persistent homology and statistical inference.

unif2d

2-dimensional point cloud of a unit square

# **Description**

A dataset containing the Cartesian coordinates of 100 points uniformly distributed within a unit square.

# Usage

unif2d

# **Format**

A matrix with 100 rows and 2 columns: the x- and y-coordinates

# Source

https://github.com/rrrlw/TDAstats/blob/master/data-raw/unif2d.R

unif3d

3-dimensional point cloud of a unit cube

# **Description**

A dataset containing the Cartesian coordinates of 100 points uniformly distributed within a unit cube.

# Usage

unif3d

10 unif3d

# **Format**

A matrix with 100 rows and 3 columns: the x-, y-, and z-coordinates

# Source

https://github.com/rrrlw/TDAstats/blob/master/data-raw/unif3d.R

# **Index**

```
\ast datasets
    circle2d, 3
    sphere3d, 8
    unif2d, 9
    unif3d, 9
calculate_homology, 2
circle2d, 3
id_significant, 4
permutation_test, 5
phom.dist, 6
\verb|plot_barcode|, 6
plot\_persist, 7
sphere3d, 8
\mathsf{TDAstats}, 9
TDAstats-package (TDAstats), 9
unif2d, 9
unif3d, 9
```