# Package 'jackstraw'

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Type Package

Title Statistical Inference for Unsupervised Learning

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Description Test for association between the observed data

and their systematic patterns of variations.

The jackstraw enables statistical testing for association between observed variables and latent variables, as captured by principal component analysis (PCA), factor analysis (FA), or other estimates. Similarly, unsupervised clustering, such as K-means clustering, partition around medoids (PAM), and others, finds subpopulations among the observed variables. The jackstraw estimates statistical significance of cluster membership, including unsupervised evaluation of cell identities in single cell RNA-seq. P-values and posterior probabilities allows one to rigorously evaluate the strength of cluster membership assignments. See the GitHub repository for the latest developments and further helps.

#### LazyData true

**Depends** R (>= 3.0.0)

irlba, rsvd

#### biocViews

Imports corpcor, cluster, methods, ClusterR, qvalue, lfa, stats,

Suggests parallel, knitr, rmarkdown

URL https://github.com/ncchung/jackstraw

BugReports https://github.com/ncchung/jackstraw/issues

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dev.R

Compute Deviance for Logistic Factors

# Description

This function computes deviance between the full model and the null (intercept-only) model. It uses built-in R functions, namely glm; slow but no C++ dependencies. Make sure that LFr1 and LFr0 do not have intercept terms.

# Usage

dev.R(dat, LFr1, LFr0 = NULL, p = FALSE)

# Arguments

dat	a matrix with m rows and n columns.
LFr1	alternative logistic factors (an output from lfa or lfa.corpcor)
LFr0	null logistic factors (an output from lfa or lfa.corpcor)
р	estimate p-values (by default, 'FALSE')

# Value

When p=FALSE (by default), dev.R returns a vector of m deviances.

When p=TRUE, a list consisting of

dev	the m deviances
p.value	the m p-values based on a chisq distribution

#### find\_k

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

find\_k

Find a number of clusters or principal components

#### Description

There are a wide range of algorithms and visual techniques to identify a number of clusters or principal components embeddd in the observed data.

#### Usage

find\_k()

#### Details

It is critical to explore the eigenvalues, cluster stability, and visualization. See R packages bootcluster, EMCluster, and nFactors.

Please see the R package SC3, which provides estkTW() function to find the number of significant eigenvalues according to the Tracy-Widom test.

ADPclust package includes adpclust() function that runs the algorithm on a range of K values. It helps you to identify the most suitable number of clusters.

This package also provides an alternative methods in permutationPA. Through a resampling-based Parallel Analysis, it finds a number of significant components.

jackstraw

jackstraw: Statistical Inference for Unsupervised Learning

#### Description

Test for association between the observed data and their systematic patterns of variations. The jackstraw enables statistical testing for association between observed variables and latent variables, as captured by principal component analysis (PCA), factor analysis (FA), or other estimates. Similarly, unsupervised clustering, such as K-means clustering, partition around medoids (PAM), and others, finds subpopulations among the observed variables. The jackstraw estimates statistical significance of cluster membership, including unsupervised evaluation of cell identities in single cell RNA-seq. P-values and posterior probabilities allows one to rigorously evaluate the strength of cluster membership assignments. See the GitHub repository for the latest developments and further helps.

#### Details

The jackstraw package provides a resampling strategy and testing scheme to estimate statistical significance of association between the observed data and their latent variables. Depending on the data type and the analysis aim, the latent variables may be estimated by principal component analysis, K-means clustering, and related algorithms. The jackstraw methods learn over-fitting characteristics inherent in this circular analysis, where the observed data are used to estimate the latent variables and to again test against the estimated latent variables.

The jackstraw tests enable us to identify the data features (i.e., variables or observations) that are driving systematic variation, in a completely unsupervised manner. Using jackstraw\_pca, we can find statistically significant features with regard to the top r principal components. Alternatively, jackstraw\_kmeans can identify the data features that are statistically significant members of the data-dependent clusters. Furthermore, this package includes more general algorithms such as jackstraw\_subspace for the dimension reduction techniques and jackstraw\_cluster for the clustering algorithms.

Overall, it computes m p-values of association between the m data features and their corresponding latent variables. From m p-values, pip computes posterior inclusion probabilities, that are useful for feature selection and visualization.

## Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### See Also

jackstraw\_pca jackstraw\_subspace jackstraw\_kmeans jackstraw\_cluster

jackstraw\_cluster Jackstraw for the User-Defined Clustering Algorithm

#### Description

Test the cluster membership using a user-defined clustering algorithm

## jackstraw\_cluster

# Usage

```
jackstraw_cluster(
    dat,
    k,
    cluster = NULL,
    centers = NULL,
    algorithm = function(x, centers) kmeans(x, centers, ...),
    s = 1,
    B = 1000,
    center = TRUE,
    noise = NULL,
    covariate = NULL,
    verbose = FALSE,
    seed = NULL,
    ...
)
```

#### Arguments

dat	a data matrix with m rows as variables and n columns as observations.
k	a number of clusters.
cluster	a vector of cluster assignments.
centers	a matrix of all cluster centers.
algorithm	a clustering algorithm to use, where an output must include 'cluster' and 'cen- ters'. For exact specification, see kmeans.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations.
center	a logical specifying to center the rows. By default, TRUE.
noise	specify a parametric distribution to generate a noise term. If NULL, a non-parametric jackstraw test is performed.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
verbose	a logical specifying to print the computational progress. By default, FALSE.
seed	a seed for the random number generator.
	optional arguments to control the clustering algorithm.

# Details

The clustering algorithms assign m rows into K clusters. This function enable statistical evaluation if the cluster membership is correctly assigned. Each of m p-values refers to the statistical test of that row with regard to its assigned cluster. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of clusters from the observed data and protects against an anti-conservative bias.

The user is expected to explore the data with a given clustering algorithm and determine the number of clusters k. Furthermore, provide cluster and centers as given by applying algorithm onto dat. The rows of centers correspond to k clusters, as well as available levels in cluster. This function allows you to specify a parametric distribution of a noise term. It is an experimental feature.

#### Value

jackstraw\_cluster returns a list consisting of

F.obs	m observed F statistics between variables and cluster centers.
F.null	F null statistics between null variables and cluster centers, from the jackstraw method.
p.F	m p-values of membership.

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities <a href="https://academic.oup.com/bioinformatics/article/36/10/3107/5788523">https://academic.oup.com/bioinformatics/article/36/10/3107/5788523</a>

jackstraw_irlba	Non-Parametric Jackstraw for Principal Component Analysis (PCA)
	using the augmented implicitly restarted Lanczos bidiagonalization al-
	gorithm (IRLBA)

#### Description

Test association between the observed variables and their latent variables captured by principal components (PCs). PCs are computed using the augmented implicitly restarted Lanczos bidiagonalization algorithm (IRLBA; see irlba).

## Usage

```
jackstraw_irlba(
    dat,
    r1 = NULL,
    r = NULL,
    s = NULL,
    B = NULL,
    covariate = NULL,
    verbose = TRUE,
```

seed = NULL,
...

#### Arguments

)

r1 a numeric vector of principal components of interest. Choose a subset of significant PCs to be used.	ſr
r a number (a positive integer) of significant principal components. See permutionPA and other methods.	ta-
s a number (a positive integer) of "synthetic" null variables. Out of m variables variables are independently permuted.	, S
B a number (a positive integer) of resampling iterations. There will be a total s*B null statistics.	of
covariate a data matrix of covariates with corresponding n observations (do not include intercept term).	an
verbose a logical specifying to print the computational progress.	
seed a numeric seed for the random number generator.	
additional arguments to irlba.	

#### Details

This function computes m p-values of linear association between m variables and their PCs. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of PCs from the observed data and protects against an anti-conservative bias.

Provide the data matrix, with m variables as rows and n observations as columns. Given that there are r significant PCs, this function tests for linear association between m variables and their r PCs.

You could specify a subset of significant PCs that you are interested in (PC). If PC is given, then this function computes statistical significance of association between m variables and PC, while adjusting for other PCs (i.e., significant PCs that are not your interest). For example, if you want to identify variables associated with 1st and 2nd PCs, when your data contains three significant PCs, set r=3 and PC=c(1,2).

Please take a careful look at your data and use appropriate graphical and statistical criteria to determine a number of significant PCs, r. The number of significant PCs depends on the data structure and the context. In a case when you fail to specify r, it will be estimated from a permutation test (Buja and Eyuboglu, 1992) using a function permutationPA.

If s is not supplied, s is set to about 10% of m variables. If B is not supplied, B is set to m\*10/s.

#### Value

jackstraw\_irlba returns a list consisting of

p.value	${\tt m} p\mbox{-values}$ of association tests between variables and their principal components
obs.stat	m observed F-test statistics
null.stat	s*B null F-test statistics

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### See Also

jackstraw jackstraw\_subspace permutationPA

#### Examples

```
set.seed(1234)
## simulate data from a latent variable model: Y = BL + E
B = c(rep(1,50),rep(-1,50), rep(0,900))
L = rnorm(20)
E = matrix(rnorm(1000*20), nrow=1000)
dat = B %*% t(L) + E
dat = t(scale(t(dat), center=TRUE, scale=TRUE))
## apply the jackstraw
out = jackstraw_irlba(dat, r=1)
## Use optional arguments
## For example, set s and B for a balance between speed of the algorithm and accuracy of p-values
## Not run:
## out = jackstraw_irlba(dat, r=1, s=10, B=1000, seed=5678)
## End(Not run)
```

jackstraw\_kmeans Non-Parametric Jackstraw for K-means Clustering

# Description

Test the cluster membership for K-means clustering

#### Usage

```
jackstraw_kmeans(
   dat,
   kmeans.dat,
   s = NULL,
```

## jackstraw\_kmeans

```
B = NULL,
center = TRUE,
covariate = NULL,
verbose = FALSE,
pool = TRUE,
seed = NULL,
...
```

#### Arguments

)

dat	a matrix with m rows as variables and n columns as observations.
kmeans.dat	an output from applying kmeans() onto dat.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations.
center	a logical specifying to center the rows. By default, TRUE.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
verbose	a logical specifying to print the computational progress. By default, FALSE.
pool	a logical specifying to pool the null statistics across all clusters. By default, TRUE.
seed	a seed for the random number generator.
	optional arguments to control the k-means clustering algorithm (refers to kmeans).

# Details

K-means clustering assign m rows into K clusters. This function enable statistical evaluation if the cluster membership is correctly assigned. Each of m p-values refers to the statistical test of that row with regard to its assigned cluster. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of clusters from the observed data and protects against an anti-conservative bias.

The input data (dat) must be of a class 'matrix'.

#### Value

jackstraw\_kmeans returns a list consisting of

F.obs	m observed F statistics between variables and cluster centers.
F.null	F null statistics between null variables and cluster centers, from the jackstraw method.
p.F	m p-values of membership.

# Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities <a href="https://academic.oup.com/bioinformatics/article/36/10/3107/5788523">https://academic.oup.com/bioinformatics/article/36/10/3107/5788523</a>

#### Examples

```
## Not run:
set.seed(1234)
dat = t(scale(t(Jurkat293T), center=TRUE, scale=FALSE))
kmeans.dat <- kmeans(dat, centers=2, nstart = 10, iter.max = 100)
jackstraw.out <- jackstraw_kmeans(dat, kmeans.dat)</pre>
```

## End(Not run)

jackstraw_kmeanspp	Non-Parametric Jackstraw for K-means Clustering using RcppA	1 <i>r</i> -
	madillo	

#### Description

Test the cluster membership for K-means clustering, using K-means++ initialization

#### Usage

```
jackstraw_kmeanspp(
  dat,
  kmeans.dat,
  s = NULL,
  B = NULL,
  center = TRUE,
  covariate = NULL,
  verbose = FALSE,
  pool = TRUE,
  seed = NULL,
  ...
```

# )

# Arguments

dat	a matrix with m rows as variables and n columns as observations.
kmeans.dat	an output from applying ClusterR::KMeans_rcpp onto dat.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
	pendentry permuted.

В	a number of resampling iterations.
center	a logical specifying to center the rows. By default, TRUE.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
verbose	a logical specifying to print the computational progress. By default, FALSE.
pool	a logical specifying to pool the null statistics across all clusters. By default, TRUE.
seed	a seed for the random number generator.
	$optional \ arguments \ to \ control \ the \ k-means \ clustering \ algorithm \ (refers \ to \ Cluster R:: KMeans\_rcpp).$

#### Details

K-means clustering assign m rows into K clusters. This function enable statistical evaluation if the cluster membership is correctly assigned. Each of m p-values refers to the statistical test of that row with regard to its assigned cluster. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of clusters from the observed data and protects against an anti-conservative bias.

Generally, it functions identical to jackstraw\_kmeans, but this uses ClusterR::KMeans\_rcpp instead of stats::kmeans. A speed improvement is gained by K-means++ initialization and RcppArmadillo. If the input data is still too large, consider using jackstraw\_MiniBatchKmeans.

The input data (dat) must be of a class 'matrix'.

#### Value

jackstraw\_kmeanspp returns a list consisting of

F.obs	m observed F statistics between variables and cluster centers.
F.null	F null statistics between null variables and cluster centers, from the jackstraw method.
p.F	m p-values of membership.

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

# Examples

```
## Not run:
set.seed(1234)
library(ClusterR)
dat = t(scale(t(Jurkat293T), center=TRUE, scale=FALSE))
kmeans.dat <- KMeans_rcpp(dat, clusters = 10, num_init = 1,
max_iters = 100, initializer = 'kmeans++')
jackstraw.out <- jackstraw_kmeanspp(dat, kmeans.dat)
## End(Not run)
```

jackstraw\_lfa Non-Parametric Jackstraw for Logistic Factor Analysis

#### Description

Test association between the observed variables and their latent variables captured by logistic factors (LFs).

#### Usage

```
jackstraw_lfa(
  dat,
  FUN = function(x) lfa(x, r)[, , drop = FALSE],
  devR = FALSE,
  r = NULL,
  r1 = NULL,
  s = NULL,
  B = NULL,
  covariate = NULL,
  verbose = TRUE,
  seed = NULL
)
```

#### Arguments

dat	a genotype matrix with m rows as variables and n columns as observations.
FUN	a function to use for LFA (by default, it uses the lfagen package)
devR	use a R function to compute deviance. By default, FALSE (uses C++).
r	a number of significant LFs.
r1	a numeric vector of LFs of interest (implying you are not interested in all r LFs).
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations. There will be a total of s*B null statistics.
covariate	a data matrix of covariates with corresponding n observations (do not include an intercept term).

#### jackstraw\_lfa

verbose	a logical specifying to print the computational progress.
seed	a seed for the random number generator.

#### Details

This function uses logistic factor analysis (LFA) from Wei et al. (2014). Particularly, a deviation dev in logistic regression (the full model with r LFs vs. the intercept-only model) is used to assess association.

# Value

jackstraw\_lfa returns a list consisting of

p.value	m p-values of association tests between variables and their LFs
obs.stat	m observed devs
null.stat	s*B null devs

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### See Also

jackstraw\_pca jackstraw jackstraw\_subspace

#### Examples

```
set.seed(1234)
## Not run:
## simulate genotype data from a logistic factor model: drawing rbinom from logit(BL)
m=5000; n=100; pi0=.9
m0 = round(m*pi0)
m1 = m-round(m*pi0)
B = matrix(0, nrow=m, ncol=1)
B[1:m1,] = matrix(runif(m1*n, min=-.5, max=.5), nrow=m1, ncol=1)
L = matrix(rnorm(n), nrow=1, ncol=n)
BL = B %*% L
prob = exp(BL)/(1+exp(BL))
dat = matrix(rbinom(m*n, 2, as.numeric(prob)), m, n)
## apply the jackstraw_lfa
```

```
out = jackstraw_lfa(dat, 2)
## apply the jackstraw_lfa using self-contained R functions
out = jackstraw_lfa(dat, FUN = function(x) lfa.corpcor(x, 2)[, , drop = FALSE], r = 2, devR = TRUE)
## End(Not run)
```

jackstraw\_MiniBatchKmeans

Non-Parametric Jackstraw for Mini Batch K-means Clustering

#### Description

Test the cluster membership for K-means clustering

#### Usage

```
jackstraw_MiniBatchKmeans(
    dat,
    MiniBatchKmeans.output = NULL,
    s = NULL,
    B = NULL,
    center = TRUE,
    covariate = NULL,
    verbose = FALSE,
    seed = NULL,
    batch_size = floor(nrow(dat)/100),
    initializer = "kmeans++",
    pool = TRUE,
    ...
)
```

# Arguments

dat MiniBatchKmeans	a data matrix with m rows as variables and n columns as observations. s.output an output from applying ClusterR::MiniBatchKmeans() onto dat. This pro- vides more controls over the algorithm and subsequently the initial centroids used.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations.
center	a logical specifying to center the rows. By default, TRUE.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
verbose	a logical specifying to print the computational progress. By default, FALSE.

seed	a seed for the random number generator.
batch_size	the size of the mini batches.
initializer	the method of initialization. By default, kmeans++.
pool	a logical specifying to pool the null statistics across all clusters. By default, TRUE.
	optional arguments to control the Mini Batch K-means clustering algorithm (refers to ClusterR::MiniBatchKmeans).

#### Details

K-means clustering assign m rows into K clusters. This function enable statistical evaluation if the cluster membership is correctly assigned. Each of m p-values refers to the statistical test of that row with regard to its assigned cluster. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of clusters from the observed data and protects against an anti-conservative bias.

#### Value

jackstraw\_MiniBatchKmeans returns a list consisting of

F.obs	m observed F statistics between variables and cluster centers.
F.null	F null statistics between null variables and cluster centers, from the jackstraw method.
p.F	m p-values of membership.

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### Examples

```
## Not run:
library(ClusterR)
set.seed(1234)
dat = t(scale(t(Jurkat293T), center=TRUE, scale=FALSE))
MiniBatchKmeans.output <- MiniBatchKmeans(data=dat, clusters = 2, batch_size = 300,
initializer = "kmeans++")
jackstraw.output <- jackstraw_MiniBatchKmeans(dat,
MiniBatchKmeans.output = MiniBatchKmeans.output)
```

## End(Not run)

jackstraw\_pam

#### Description

Test the cluster membership for Partitioning Around Medoids (PAM)

#### Usage

```
jackstraw_pam(
  dat,
  pam.dat,
  s = NULL,
  B = NULL,
  center = TRUE,
  covariate = NULL,
  verbose = FALSE,
  pool = TRUE,
  seed = NULL,
  ...
)
```

#### Arguments

dat	a matrix with m rows as variables and n columns as observations.
pam.dat	an output from applying cluster::pam() on dat.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations.
center	a logical specifying to center the rows. By default, TRUE.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
verbose	a logical specifying to print the computational progress. By default, FALSE.
pool	a logical specifying to pool the null statistics across all clusters. By default, TRUE.
seed	a seed for the random number generator.
	optional arguments to control the k-means clustering algorithm (refers to kmeans).

#### Details

PAM assigns m rows into K clusters. This function enable statistical evaluation if the cluster membership is correctly assigned. Each of m p-values refers to the statistical test of that row with regard to its assigned cluster. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of clusters from the observed data and protects against an anti-conservative bias. For a large dataset, PAM could be too slow. Consider using cluster::clara and jackstraw::jackstraw\_clara. The input data (dat) must be of a class 'matrix'.

#### jackstraw\_pca

#### Value

jackstraw\_pam returns a list consisting of

F.obs	m observed F statistics between variables and cluster medoids.
F.null	F null statistics between null variables and cluster medoids, from the jackstraw method.
p.F	m p-values of membership.

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities <a href="https://academic.oup.com/bioinformatics/article/36/10/3107/5788523">https://academic.oup.com/bioinformatics/article/36/10/3107/5788523</a>

## Examples

```
## Not run:
library(cluster)
set.seed(1234)
dat = t(scale(t(Jurkat293T), center=TRUE, scale=FALSE))
pam.dat <- pam(dat, k=2)
jackstraw.out <- jackstraw_pam(dat, pam.dat = pam.dat)</pre>
```

## End(Not run)

jackstraw\_pca Non-Parametric Jackstraw for Principal Component Analysis (PCA)

#### Description

Test association between the observed variables and their latent variables captured by principal components (PCs).

#### Usage

jackstraw\_pca(
 dat,
 r1 = NULL,
 r = NULL,
 s = NULL,
 B = NULL,

```
covariate = NULL,
verbose = TRUE,
seed = NULL
)
```

#### Arguments

dat	a data matrix with m rows as variables and n columns as observations.
r1	a numeric vector of principal components of interest. Choose a subset of r significant PCs to be used.
r	a number (a positive integer) of significant principal components. See permuta- tionPA and other methods.
S	a number (a positive integer) of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number (a positive integer) of resampling iterations. There will be a total of s*B null statistics.
covariate	a data matrix of covariates with corresponding n observations (do not include an intercept term).
verbose	a logical specifying to print the computational progress.
seed	a numeric seed for the random number generator.

#### Details

This function computes m p-values of linear association between m variables and their PCs. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of PCs from the observed data and protects against an anti-conservative bias.

Provide the data matrix, with m variables as rows and n observations as columns. Given that there are r significant PCs, this function tests for linear association between m variables and their r PCs.

You could specify a subset of significant PCs that you are interested in (PC). If PC is given, then this function computes statistical significance of association between m variables and PC, while adjusting for other PCs (i.e., significant PCs that are not your interest). For example, if you want to identify variables associated with 1st and 2nd PCs, when your data contains three significant PCs, set r=3 and PC=c(1,2).

Please take a careful look at your data and use appropriate graphical and statistical criteria to determine a number of significant PCs, r. The number of significant PCs depends on the data structure and the context. In a case when you fail to specify r, it will be estimated from a permutation test (Buja and Eyuboglu, 1992) using a function permutationPA.

If s is not supplied, s is set to about 10% of m variables. If B is not supplied, B is set to m\*10/s.

#### Value

jackstraw\_pca returns a list consisting of

p.value	m p-values of association tests between variables and their principal components
obs.stat	m observed F-test statistics
null.stat	s*B null F-test statistics

#### jackstraw\_rpca

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities <a href="https://academic.oup.com/bioinformatics/article/36/10/3107/5788523">https://academic.oup.com/bioinformatics/article/36/10/3107/5788523</a>

#### See Also

jackstraw jackstraw\_subspace permutationPA

#### Examples

```
## Not run:
set.seed(1234)
## simulate data from a latent variable model: Y = BL + E
B = c(rep(1,50),rep(-1,50), rep(0,900))
L = rnorm(20)
E = matrix(rnorm(1000*20), nrow=1000)
dat = B %*% t(L) + E
dat = t(scale(t(dat), center=TRUE, scale=TRUE))
## apply the jackstraw
out = jackstraw_pca(dat, r=1)
## Use optional arguments
## For example, set s and B for a balance between speed of the algorithm and accuracy of p-values
## out = jackstraw_pca(dat, r=1, s=10, B=1000, seed=5678)
```

```
## End(Not run)
```

jackstraw_rpca	Non-Parametric Jackstraw for Principal Component Analysis (PCA)
	using Randomized Singular Value Decomposition

# Description

Test association between the observed variables and their latent variables captured by principal components (PCs). PCs are computed by randomized Singular Value Decomposition (see rsvd).

#### Usage

```
jackstraw_rpca(
  dat,
  r1 = NULL,
  r = NULL,
  s = NULL,
  B = NULL,
  covariate = NULL,
  verbose = TRUE,
  seed = NULL,
  ...
)
```

#### Arguments

dat	a data matrix with m rows as variables and n columns as observations.
r1	a numeric vector of principal components of interest. Choose a subset of r significant PCs to be used.
r	a number (a positive integer) of significant principal components. See permuta- tionPA and other methods.
S	a number (a positive integer) of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number (a positive integer) of resampling iterations. There will be a total of s*B null statistics.
covariate	a data matrix of covariates with corresponding n observations (do not include an intercept term).
verbose	a logical specifying to print the computational progress.
seed	a numeric seed for the random number generator.
	additional arguments to rpca.

#### Details

This function computes m p-values of linear association between m variables and their PCs. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of PCs from the observed data and protects against an anti-conservative bias.

Provide the data matrix, with m variables as rows and n observations as columns. Given that there are r significant PCs, this function tests for linear association between m variables and their r PCs.

You could specify a subset of significant PCs that you are interested in (PC). If PC is given, then this function computes statistical significance of association between m variables and PC, while adjusting for other PCs (i.e., significant PCs that are not your interest). For example, if you want to identify variables associated with 1st and 2nd PCs, when your data contains three significant PCs, set r=3 and PC=c(1,2).

Please take a careful look at your data and use appropriate graphical and statistical criteria to determine a number of significant PCs, r. The number of significant PCs depends on the data structure

#### jackstraw\_rpca

and the context. In a case when you fail to specify r, it will be estimated from a permutation test (Buja and Eyuboglu, 1992) using a function permutationPA.

If s is not supplied, s is set to about 10% of m variables. If B is not supplied, B is set to m\*10/s.

#### Value

jackstraw\_rpca returns a list consisting of

p.value	m p-values of association tests between variables and their principal components
obs.stat	m observed F-test statistics
null.stat	s*B null F-test statistics

## References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### See Also

jackstraw jackstraw\_subspace permutationPA

#### Examples

```
set.seed(1234)
## simulate data from a latent variable model: Y = BL + E
B = c(rep(1,50),rep(-1,50), rep(0,900))
L = rnorm(20)
E = matrix(rnorm(1000*20), nrow=1000)
dat = B %*% t(L) + E
dat = t(scale(t(dat), center=TRUE, scale=TRUE))
## apply the jackstraw
out = jackstraw_rpca(dat, r=1)
## Use optional arguments
## For example, set s and B for a balance between speed of the algorithm and accuracy of p-values
## out = jackstraw_rpca(dat, r=1, s=10, B=1000, seed=5678)
```

jackstraw\_subspace Jackstraw for the User-Defined Dimension Reduction Methods

# Description

Test association between the observed variables and their latent variables, captured by a user-defined dimension reduction method.

# Usage

```
jackstraw_subspace(
  dat,
  FUN,
  r = NULL,
  r1 = NULL,
  s = NULL,
  B = NULL,
  covariate = NULL,
  verbose = TRUE,
  seed = NULL
)
```

## Arguments

dat	a data matrix with m rows as variables and n columns as observations.
FUN	optionally, provide a specifc function to estimate LVs. Must output r estimated LVs in a n $\star$ r matrix.
r	a number of significant latent variables.
r1	a numeric vector of latent variables of interest.
S	a number of "synthetic" null variables. Out of m variables, s variables are independently permuted.
В	a number of resampling iterations.
covariate	a model matrix of covariates with n observations. Must include an intercept in the first column.
noise	specify a parametric distribution to generate a noise term. If NULL, a non-parametric jackstraw test is performed.
verbose	a logical specifying to print the computational progress.
seed	a seed for the random number generator.

#### Details

This function computes m p-values of linear association between m variables and their latent variables, captured by a user-defined dimension reduction method. Its resampling strategy accounts for the over-fitting characteristics due to direct computation of PCs from the observed data and protects against an anti-conservative bias.

This function allows you to specify a parametric distribution of a noise term. It is an experimental feature. Then, a small number s of observed variables are replaced by synthetic null variables generated from a specified distribution.

#### Value

jackstraw\_subspace returns a list consisting of

p.value	m p-values of association tests between variables and their principal components
obs.stat	m observed statistics
null.stat	s*B null statistics

#### Author(s)

Neo Christopher Chung <nchchung@gmail.com>

#### References

Chung and Storey (2015) Statistical significance of variables driving systematic variation in highdimensional data. Bioinformatics, 31(4): 545-554 https://academic.oup.com/bioinformatics/ article/31/4/545/2748186

Chung (2020) Statistical significance of cluster membership for unsupervised evaluation of cell identities https://academic.oup.com/bioinformatics/article/36/10/3107/5788523

#### See Also

jackstraw\_pca jackstraw

#### Examples

```
set.seed(1234)
## simulate data from a latent variable model: Y = BL + E
B = c(rep(1,50),rep(-1,50), rep(0,900))
L = rnorm(20)
E = matrix(rnorm(1000*20), nrow=1000)
dat = B %*% t(L) + E
dat = t(scale(t(dat), center=TRUE, scale=TRUE))
```

```
## apply the jackstraw with the svd as a function
out = jackstraw_subspace(dat, FUN = function(x) svd(x)$v[,1,drop=FALSE], r=1, s=100, B=50)
```

Jurkat293T

# Description

50

# Usage

Jurkat293T

#### Format

A data frame with 3381 rows correspoding to single cells and 10 columns corresponding to the top 10 principal components

#### Source

Supplementary Data 1 from Zheng et al. (2017) https://www.nature.com/articles/ncomms14049

#### References

Zheng et al. (2017) Massively parallel digital transcriptional profiling of single cells. Nature Communications. 8:14049. DOI: 10.1038/ncomms14049

lfa.corpcor

Logistic Factor Analysis without C++ Dependency

#### Description

Estimate populatoin structure in genome-wide genotype matrices.

#### Usage

lfa.corpcor(x, d, ltrace = FALSE)

#### Arguments

Х	a matrix with m loci (rows) and n observations (columns).
d	a number of logistic factors.
ltrace	a logical indicator as to whether to print the progress.

## permutationPA

# Details

It performs the logistic factor analysis, similar to 1fa function in the lfa package. This function works without C++ dependencies. However, it would be much slower, does not include any other LFA-related functions, checks, and warnings. This function remains here for backward compatability - please use the 1fa package.

# Value

lfa.corpcor returns a n\*d matrix of d logistic factors. The last column is always an intercept term.

## See Also

jackstraw\_lfa

permutationPA Permutation Parallel Analysis

# Description

Estimate a number of significant principal components from a permutation test.

#### Usage

```
permutationPA(dat, B = 100, threshold = 0.05, verbose = TRUE, seed = NULL)
```

#### Arguments

dat	a data matrix with m rows as variables and n columns as observations.
В	a number (a positive integer) of resampling iterations.
threshold	a numeric value between 0 and 1 to threshold p-values.
verbose	a logical indicator as to whether to print the progress.
seed	a seed for the random number generator.

# Details

Adopted from sva::num.sv, and based on Buja and Eyuboglu (1992)

#### Value

permutationPA returns

р	a list of p-values for significance of principal components
r	an estimated number of significant principal components based on thresholding
	p-values at threshold

# References

Buja A and Eyuboglu N. (1992) Remarks on parrallel analysis. Multivariate Behavioral Research, 27(4), 509-540

pip

Compute posterior inclusion probabilities (PIPs)

# Description

Computes posterior probabilities that a feature is a true member of an assigned cluster

# Usage

pip(pvalue, group = NULL, ...)

# Arguments

pvalue	a vector of p-values.
group	a vector of group indicators (optional).
	optional arguments to control a local FDR estimation.

# Value

pip returns a vector of posterior inclusion probabilities

# Author(s)

Neo Christopher Chung <nchchung@gmail.com>

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