# dSVDsignif

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dSVDsignif

Function to obtain SVD-based gene significance from the input genesample matrix

## Description

dSVDsignif is supposed to obtain gene signficance from the given gene-sample matrix according to singular value decomposition (SVD)-based method. The method includes: 1) singular value decomposition of the input matrix; 2) determination of the eigens in consideration (if not given); 3) construction of the gene-specific project vector based on the considered eigens; 4) calculation of the distance statistic from the projection vector to zero point vector; and 5) based on distance statistic to obtain the gene significance.

## Usage

```
dSVDsignif(data, num.eigen = NULL, pval.eigen = 0.01, signif = c("fdr",
"pval"), orient.permutation = c("row", "column", "both"),
num.permutation = 100, fdr.procedure = c("stepup", "stepdown"),
verbose = T)
```

#### **Arguments**

data	an input gene-sample data matrix used for singular value decomposition
num.eigen	an integer specifying the number of eigens in consideration. If NULL, this num-

ber will be automatically decided on based on the observed relative eigenexpression against randomised relative eigenexpression calculated from a list (here

100) of permutated input matrix

pval.eigen p-value used to call those eigens as dominant. This parameter is used only

when parameter 'num.eigen' is NULL. Here, p-value is calcualted to assess how likely the observed relative eigenexpression are more than the maximum relative

eigenexpression calculated from permutated matrix

signif the singificance to return. It can be either "pval" for using the p-value as the

gene significance, or "fdr" for using the fdr as the gene significance

orient.permutation

the orientation of matrix being permutated. It can be either "row" to permutate values within each row, or "column" to permutate values within each column, or "both" to permutate values both within rows and columns. Notably, when using the p-value as the gene significance, it is always to permutate values within each row.

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num.permutation

an integer specifying how many permutations are used

fdr.procedure the procedure to adjust the fdr. To ensure that the high distance statistic the

more significance, the fdr should be adjusted either using "stepup" for step-up procedure (from the most significant to the least significant) or using "stepdown" for step-down procedure (from the least significant to the most significant)

verbose logical to indicate whether the messages will be displayed in the screen. By

default, it sets to true for display

#### Value

a vector storing gene significance

#### Note

none

### See Also

dFDRscore

# **Examples**

```
# 1) generate data with an iid matrix of 1000 x 9
data <- cbind(matrix(rnorm(1000*3,mean=0,sd=1), nrow=1000, ncol=3),
matrix(rnorm(1000*3,mean=0.5,sd=1), nrow=1000, ncol=3),
matrix(rnorm(1000*3,mean=-0.5,sd=1), nrow=1000, ncol=3))
# 2) calculate the significance according to SVD
# using "fdr" significance
fdr <- dSVDsignif(data, signif="fdr", num.permutation=10)
# using "pval" significance
pval <- dSVDsignif(data, signif="pval", num.permutation=10)</pre>
```