# Package 'SMVar'

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Title Structural Model for Variances
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<b>Depends</b> R (>= $2.6.0$ )
<b>Description</b> Implementation of the structural model for variances in order to detect differentially expressed genes from gene expression data.
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NeedsCompilation no
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SMVar-package Structural Model for Variances

## Description

Type Package

Package containing moderated t-tests to detect differentially expressed genes for paired and unpaired data

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## **Details**

Package: SMVar
Type: Package
Version: 1.3.3
Date: 2011-08-03
License: GPL

SMVar.unpaired and SMVar.paired are the most important functions.

## Author(s)

Guillemette Marot <guillemette.marot@inria.fr>

#### References

F. Jaffrezic, Marot, G., Degrelle, S., Hue, I. and Foulley, J. L. (2007) A structural mixed model for variances in differential gene expression studies. Genetical Research (89) 19:25

#### **Examples**

```
library(SMVar)
data(ApoAIdata)
attach(ApoAIdata)
SMVar.unpaired(ApoAIGeneId,list(ApoAICond1,ApoAICond2))
```

ApoAIdata

ApoAIdata

## **Description**

Example dataset for unpaired data

## Usage

data(ApoAIdata)

#### **Format**

ApoAIdata is a list with 3 elements

ApoAIGeneId vector of fictive gene names)

ApoAICond1 matrix with 6226 rows and 8 columns with normalized normal mice measurements

ApoAICond2 matrix with 6226 rows and 8 columns with normalized KO mice measurements

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#### **Source**

Similar to the example dataset used in the package Varmixt

#### References

M.J. Callow, S. Dudoit, E.L. Gong, T.P. Speed, and E.M. Rubin. Microarray expression profiling identifies genes with altered expression in hdl-deficien mice. Genome Res., 10(12): 2022-9, 2000

## **Examples**

```
data(ApoAIdata)
attach(ApoAIdata)
```

SMVar.paired

Structural model for variances with paired data

## Description

Function to detect differentially expressed genes when data are paired

## Usage

#### **Arguments**

geneNumbers	Vector with gene names or dataframe which contains all information about spots on the chip
logratio	matrix with one row by gene and one column by replicate giving the logratio
fileexport	file to export the list of differentially expressed genes
minrep	minimum number of replicates to take a gene into account, minrep must be higher than $\boldsymbol{2}$
method	method of multiple tests adjustment for p.values
threshold	threshold of False Discovery Rate

## **Details**

This function implements the structural model for variances described in (Jaffrezic et al., 2007). Data must be normalized before calling the function. Matrix geneNumbers must have one of the following formats: "matrix", "data.frame", "vector", "character", "numeric", "integer".

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#### Value

Only the number of differentially expressed genes is printed. If asked, the file giving the list of differentially expressed genes is created

If the user creates an object when calling the function (for example "Stat=SMVar.paired(...)") then Stat contains the information for all genes, is sorted by ascending p-values and

Stat\$TestStat gives the test statistics as described in the paper

Stat\$StudentPValue

gives the raw p-values

Stat\$DegOfFreedom

gives the number of degrees of freedom for the Student distribution for the test

statistics

Stat\$LogRatio gives the logratios

Stat\$AdjPValue gives the adjusted p-values

#### Note

If the first column of the file geneNumbers contains identical names for two different spots, these two spots are only counted once if they are both differentially expressed. By default, the correction for multiple testing is Benjamini Hochberg with a threshold of False Discovery Rate (FDR) of 5%. The FDR threshold can be changed, and it is also possible to choose the multiple test correction method ("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"). To see the references for these methods, use the R-help ?p.adjust.

## Author(s)

Guillemette Marot with contributions from Anne de la Foye

#### References

F. Jaffrezic, Marot, G., Degrelle, S., Hue, I. and Foulley, J. L. (2007) A structural mixed model for variances in differential gene expression studies. Genetical Research (89) 19:25

## **Examples**

library(SMVar)
data(Spleendata)
attach(Spleendata)
SMVar.paired(SpleenGeneId,SpleenLogRatio)

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SMVar.unpaired	Structural model for variances with unpaired data	

## **Description**

Function to detect differentially expressed genes when data are unpaired

#### Usage

#### **Arguments**

geneNumbers Vector with gene names or dataframe which contains all information about spots on the chip

list cond list of the different conditions to be compared fileexport file to export the list of differentially expressed genes

minrep minimum number of replicates to take a gene into account, minrep must be

higher than 2

method method of multiple tests adjustment for p.values

threshold threshold of False Discovery Rate

#### Details

This function implements the structural model for variances described in (Jaffrezic et al., 2007). Data must be normalized before calling the function. Matrix geneNumbers must have one of the following formats: "matrix", "data.frame", "vector", "character", "numeric", "integer".

#### Value

Only the number of differentially expressed genes is printed. If asked, the file giving the list of differentially expressed genes is created.

If the user creates an object when calling the function (for example "Stat=SMVar.paired(...)") then Stat contains the information for all genes, is sorted by ascending p-values and

Stat\$TestStat gives the test statistics as described in the paper Stat\$StudentPValue

gives the raw p-values

Stat\$DegOfFreedom

gives the number of degrees of freedom for the Student distribution for the test

statistics

Stat\$Cond1 gives the first condition considered in the log-ratio
Stat\$Cond2 gives the second condition considered in the log-ratio
Stat\$LogRatio gives the logratios (listcond[[Cond2]]-listcond[[Cond1]])

Stat\$AdjPValue gives the adjusted p-values

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#### Note

If the first column of the file geneNumbers contains identical names for two different spots, these two spots are only counted once if they are both differentially expressed. By default, the correction for multiple testing is Benjamini Hochberg with a threshold of False Discovery Rate (FDR) of 5%. The FDR threshold can be changed, and it is also possible to choose the multiple test correction method ("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none"). To see the references for these methods, use the R-help ?p.adjust.

#### Author(s)

Guillemette Marot with contributions from Anne de la Foye

#### References

F. Jaffrezic, Marot, G., Degrelle, S., Hue, I. and Foulley, J. L. (2007) A structural mixed model for variances in differential gene expression studies. Genetical Research (89) 19:25

#### **Examples**

```
library(SMVar)
data(ApoAIdata)
attach(ApoAIdata)
SMVar.unpaired(ApoAIGeneId,list(ApoAICond1,ApoAICond2))
```

Spleendata

Spleendata

## Description

Example dataset for paired data

#### Usage

```
data(Spleendata)
```

#### **Format**

Spleendata is a list with 2 elements

```
SpleenGeneId Gene names)
```

SpleenLogRatio Matrix with 4360 rows and 6 columns with normalized log-ratio

#### Source

Similar to the example dataset used in the package Varmixt

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## References

P. Delmar, Robin, S., Tronik-Le Roux S. and Daudin J.-J. (2005) Mixture model on the variance for the differential analysis of gene expression data, JRSS series C, 54(1), 31:50

## Examples

data(Spleendata)
attach(Spleendata)

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