# Package 'RNAinteractMAPK'

October 17, 2024

Type Package

**Title** Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi

Version 1.42.0

## Description

This package includes all data used in the paper -Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi- by Horn, Sandmann, Fischer et al.., Nat. Methods, 2011. The package vignette shows the R code to reproduce all figures in the paper.

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LazyLoad yes

**Imports** grid, gdata, MASS, genefilter, methods, fields, utils, lattice, Biobase

**Depends** R (>= 2.12.0), sparseLDA, RNAinteract

Suggests qvalue

**biocViews** ExperimentData, MicrotitrePlateAssayData, Drosophila\_melanogaster\_Data, CellCulture

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18

# **Contents**

RNAinteractMAPK-package
cellTiterGlo
Dmel2PPMAPK
dsRNAiDilutionSeries
ElpB1phenotype
MAPK.cv.classifier
MAPK.estimate.TPS
MAPK.plot.heatmap.raster
MAPK.report.gene.lists.paper
MAPK.smooth.scatter
mRNAdoubleKDefficiency
mRNAsingleKDefficiency
Networks
pathwayMembership
PhysicalInteractions
singleKDphenotype

RNAinteractMAPK-package

Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi.

# **Description**

Index

The package contains the data and the source code to reproduce the results and figures from the paper

T. Horn, T. Sandmann, B. Fischer, W. Huber, M. Boutros. Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi. Nature Methods, 2011.

## **Details**

See vignette("RNAinteractMAPK") for details.

## Package content

See vignette("RNAinteractMAPK") for more detail on how to obtain the data used for specific figures. In addition this vignette contains the complete analysis and the generation of all figures.

The main screen can be loaded by data("Dmel2PPMAPK", package="RNAinteractMAPK"). Access to the pairwise interaction data is done via the getData function from the RNAinteract-package. See example below.

The following datasets are provided with this package:

Dme12PPMAPK

interaction data of main screen. See example below.

ElpB1phenotype
mRNAdoubleKDefficiency
mRNAsingleKDefficiency
singleKDphenotype
dsRNAiDilutionSeries
Networks
pathwayMembership
PhysicalInteractions
cellTiterGlo

in vivo experiment on ectopic wing vein formation (used in Figure 5f) mRNA level after double gene knockdown (used in Figure S1) mRNA level after single gene knockdown (used in Figure S2) single knockdown phenotypes (used in Figure S3) dsRNA dilution series (used in Figure S5) known interactions from DroID (used in Figure S12) pathway membership of genes (used in Figure S13) Known physical interactions (used in Figure S13) cellTiterGlo viability data (used in Figure S15)

Within this package a number of specialized functions is defined written for the analysis of the MAPK interaction screen and additional experiments shown in the paper. These functions are not intended to be general purpose analysis functions, but should document the steps of analysis of the paper. Therefore, these functions are not exported. A list of functions is given below. A general purpose package for the analysis of genetic interaction screens is the package RNAinteract.

The following functions are provided within this package.

Functions used for the classification: MAPK.predict.classification, MAPK.cv.classifier, MAPK.getCV, MAPK.ternary.plot, MAPK.getXY, MAPK.plot.classification.

Functions for the analysis of the interaction surfaces: MAPK.plot.TPS.single, MAPK.plot.TPS.all, MAPK.estimate.TPS, MAPK.cv.TPS, MAPK.screen.as.array.

A function to write the hit list: MAPK.report.gene.lists.paper.

A function to plot a heatmap: MAPK.plot.heatmap.raster.

A function to plot smaooth scatters: MAPK.smooth.scatter.

#### Author(s)

Bernd Fischer

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#### References

T. Horn, T. Sandmann, B. Fischer, W. Huber, M. Boutros. Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi. Nature Methods, 2011.

#### See Also

RNAinteractMAPK-package

4 cellTiterGlo

Commission of interesting and middle and MT's a Classical like
Comparison of interaction experiment with an cellTiterGlo viability
assay

# Description

For ten gene pairs genetic interactions are measured. The experiment contains 24 different conditions. These are repeated in each row of the three 384 mutli well plates. The data frame contains the plate annotation as well as the viability readout for the three plates.

# Usage

```
data(cellTiterGlo)
```

## **Format**

A data frame with 384 observations on the following 6 variables.

```
Well a character vector
dsRNA_1 a character vector
dsRNA_2 a character vector
plate1 a numeric vector
plate2 a numeric vector
plate3 a numeric vector
```

## **Source**

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S15.

```
data(cellTiterGlo)
head(cellTiterGlo)
```

Dmel2PPMAPK 5

Dme12PPMAPK

The interaction data of the main screen

## **Description**

Dmel2PPMAPK is an object of class RNAinteract. It contains the raw data, the computed main effects, pairwise interaction scores, p-values and q-values computed by a t-test. The package vignette vontains the complete code and documentation for the statistical analysis.

#### Usage

```
data(Dmel2PPMAPK)
```

#### **Format**

An object of class RNAinteract.

#### Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, main interaction screen.

# **Examples**

dsRNAiDilutionSeries dsRNA dilution series

## **Description**

A dilution series for 6 x 6 gene. For each gene pair all combinations of 8 different concentrations of dsRNA reagent are screened. Three readout channels (nrCells, area, intensity) are available in the data.frame dsRNAiDilutionSeries. The plate annotation is given in the data.frame dsRNAiDilutionSeriesAnno and precomputed degrees of freedom for thin plate splines are available in the matrix dsRNAiDilutionSeriesDF.

# Usage

```
data(dsRNAiDilutionSeries)
```

6 ElpB1phenotype

#### **Format**

• dsRNAiDilutionSeries: A data frame with 2688 observations on the following 3 variables.

nrCells a numeric vector representing the number of cells readout.

area a numeric vector representing the are readout.

intensity a numeric vector representing the intensity readout.

dsRNAiDilutionSeriesAnno: A data frame with 2688 observations on the following 7 variables.

plate a numeric vector representing the plate number.

well a numeric vector representing the well.

row a numeric vector representing the row on plate.

col a numeric vector representing the column on plate.

targetID1 a numeric vector representing the target number of the first reagent (see targets).

targetID2 a numeric vector representing the target number of the second reagent (see targets).

targets a data.frame representing the 49 target reagents For each target reagent the gene name and the dsRNAi concentration is given.

• dsRNAiDilutionSeriesDF: A 6 x 6 matrix with degrees of freedom for thin-plate spline regression.

#### **Source**

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure 1abc and Figure S5.

# **Examples**

data(dsRNAiDilutionSeries)
head(dsRNAiDilutionSeries)
head(dsRNAiDilutionSeriesAnno)
head(dsRNAiDilutionSeriesDF)

ElpB1phenotype

Ectopic wing vein formation phenotype caused by the EgfrElpB1.

# **Description**

Partial suppression of ectopic wing vein formation (in vivo fly phenotpye) caused by the EgfrElpB1 in Cka heterozygous mutant backgrounds. The wing vein formation phenotype is classified as strong and notstrong. It is tested for three fly mutants.

## Usage

data(ElpB1phenotype)

MAPK.cv.classifier 7

#### **Format**

A data frame with 3 observations on the following 2 variables.

```
strong a numeric vector notstrong a numeric vector
```

#### Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure 5f.

## **Examples**

```
data(ElpB1phenotype)
ElpB1phenotype
```

MAPK.cv.classifier

A classifier for genetic interaction data.

## **Description**

These functions implement a classifier to classify three classes of pathway membership of the RasMAPK and JNK pathway. For each sample and each channel a sparse linear discriminat classifier is trained. The posterior probabilities are averaged over all single classifiers. The classification posterior probabilities of three classes are plotted as a ternary plot (ternary plot adapted from CRAN package vcd).

## Usage

# **Arguments**

sgi An object of class RNAinteract

traingroups A list of gene names for the training examples. For each class there should be a

vector of gene names.

posterior A matrix of posterior probabilities. Each row represents one gene, each column

represents one class.

8 MAPK.cv.classifier

classes The three classes to be displayed on the ternary plot.

classnames The class names to be displayed.

col The color used for the text labels.

y A factor representing the class label for each gene in posterior.

classcol The color used for the three classes.

main The title of the plot.

pop If TRUE, all viewports are popped before finishing the function.

threshText A threshold for the posterior probability of the three classes. Only genes that are

assigned with a larger probability to the three classes are shown.

textToLeft These text labels will be shown on the left hand side.

textToRight These text labels will be shown on the right hand side.

#### **Details**

The code for the ternary plot (used by MAPK.plot.classification) is adapted from the function ternaryplot in the CRAN package vcd Author of the original code is David Meyer (David.Meyer@R-project.org). References: M. Friendly (2000), Visualizing Categorical Data. SAS Institute, Cary, NC. This code is specialized for the publication "Mapping Signalling Networks by RNAi ..." in Nat. Methods. It is highly recommended to use the original code by David Meyer.

## Value

MAPK.cv.classifier returns a list with the cross validated class assignment probability, as well as the results of the single classifiers.

MAPK.predict.classifier returns the predicted posterior probabilities of new genes as well as the classification results of the single classifier.

MAPK.plot.classifier returns nothing.

## Author(s)

Bernd Fischer

#### References

function ternaryplot, CRAN package vdc. M. Friendly (2000), Visualizing Categorical Data. SAS Institute, Cary, NC.

## See Also

RNAinteract, RNAinteractMAPK

MAPK.estimate.TPS 9

MAPK.estimate.TPS	genetic interaction surfaces
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# Description

Genetic interaction surfaces are estimated from a dilution experiment. Cells are treated with two RNAi's. The concentration of the RNAi reagent is changed in 8 steps. All 8 x 8 combinations of concentrations are tested for 6 x 6 gene pairs.

# Usage

# Arguments

data, Anno	
data	A data.frame containing the read.out of the dilution screen. Each row is one well. Each column one feature.
Anno	A data frame containing the plate configuration. For each row in data there should a row in Anno.
A	An array of dimension concentration $x$ concentration $x$ genes $x$ genes $x$ channel as returned by MAPK.screen.as.array.
DF	A 6 x 6 matrix of degrees of freedom for the thin spline plate regression.
n.out	number of points for sampling from the regression function.
channel	The channel used.
range.df	The range of degrees of freedom that is considered for cross validation.
range	The range of pairwise interaction scores that is shown by the colorbar.
gene1, gene2	The genes for which the interaction surface is plotted.
TPSmodel	The TPS model estimated by MAPK.estimate.TPS
fill	The range of colors used for the color code.

#### **Details**

The screen readout can be reshaped as an array with dimensions concentration x concentration x genes x genes x channel by the function MAPK.screen.as.array. Then the function MAPK.estimate.TPS fits a regression in the 8 x 8 pairwise dilution series. The degrees of freedom for the regression can be estimated automatically by cross validation with the function MAPK.cv.TPS. Finally one can plot the interaction surface for a single gene or an overview of interaction surfaces for all genes with the functions MAPK.plot.TPS.single or MAPK.plot.TPS.all.

#### Value

- MAPK. screen. as. array returns an array of dimensions concentration x concentration x genes x genes x channel with the screen data.
- MAPK.estimate.TPS returns a regression model estimated by thin plate splines for each pair of genes and subsampled matrices.
- MAPK.cv.TPS.all returns
  - DF a matrix with degrees of freedom.
  - CVerror The prediction error estimated by cross validation.
  - CVerrorSD The standard deviation of the prediction error estimated by cross validation.
- MAPK.plot.TPS.single,MAPK.plot.TPS.all: An object of class "trellis". See levelplot for details.

## Author(s)

Bernd Fischer

## See Also

RNAinteract-package, RNAinteractMAPK-package

```
MAPK.plot.heatmap.raster
```

Plots a heatmap using grid.raster

#### **Description**

This functions provides a grid plot that displays the raster image of a heatmap without any axis or label. This function is adapted from the function grid.sgiHeatmap from the package RNAinteract-package. It is highly recommended to use the original function grid.sgiHeatmap.

# Usage

## **Arguments**

X	A matrix of pairwise interaction scores.
subset	A subset of genes that are displayed in the rows.
hc.row	A helust object.
hc.col	A helust object.
pi.max	The maximum interaction score of the colorbar. All interaction scores larger

than this value will be displayed in the same color.

## Value

Nothing is returned.

# Author(s)

Bernd Fischer

## See Also

RNAinteract-package, RNAinteractMAPK-package

```
MAPK.report.gene.lists.paper

A hitlist report.
```

# **Description**

Reports the hitlist of genetic interactions, with p-values from a t-test with pooled variance estimate, from limma, and from Hotelling T^2 test.

# Usage

```
MAPK.report.gene.lists.paper(sgi, sgilimma, sgi3T2, screen = "mean")
```

# **Arguments**

sgi	An object of class RNAinteract containing p-values from a t-test with pooled variance estimate.
sgilimma	An object of class RNAinteract containing p-values from limma.
sgi3T2	An object of class RNAinteract containing p-values from a Hotelling T^2 test.
screen	The screen name for which the report should be written.

# **Details**

Writes tab-separated lists for each single test as well as a joint table with all three tests.

12 MAPK.smooth.scatter

# Value

Nothing is returned.

# Author(s)

Bernd Fischer

# See Also

RNAinteract-package, RNAinteractMAPK-package

MAPK.smooth.scatter smooth scatter using grid raster

# Description

This function is a reimplementation of smoothScatter. For nicer graphics output the background image is written by grid.raster. It is recommended to use the smoothScatter function from the graphics package.

# Usage

```
MAPK.smooth.scatter(x, y, n = 75,
    nrpoints = 100, col = "blue",
    pch = 20, size = unit(0.3, "char"), cex = 1.2,
    colramp = colorRampPalette(c("white", "blue", "green", "yellow", "red"))(256),
    xlab = "", ylab = "", respect = FALSE)
```

# Arguments

X	x-values.
У	y-values. Has to be the same length as x.
n	nr of bins used for the kernel density estimation.
nrpoints	nrpoints points in the lowest density region will be plotted. This allows the identification of outliers.
col	color of points.
pch	symbol to plot points.
size	The size of the points.
cex	The size of the label text.
colramp	color ramp for the density plot.
xlab, ylab	axis labels.
respect	A logical value indicating if the height and width of the axis scales should respect each other.

#### **Details**

Plots a density plot with grid graphics.

#### Author(s)

Bernd Fischer

#### See Also

RNAinteractMAPK-package, smoothScatter

mRNAdoubleKDefficiency

mRNA levels for double knock downs

# **Description**

qPCR measurements for the mRNA level after a double gene knock down (ratio relative to wild type control). The experiments tests the knock down efficiency in the presence of a second gene knock down.

## Usage

data(mRNAdoubleKDefficiency)

#### **Format**

A data frame with 320 observations on the following 5 variables.

```
template an ordered factor with levels Fluc < CG10417 < CG13197 < CG9391 < egr < lic < PRL-1 < Rho1 < Tak1
```

query a factor with levels CG10417 [query dsRNA] CG13197 [query dsRNA] CG9391 [query dsRNA] egr [query dsRNA] lic [query dsRNA] PRL-1 [query dsRNA] Rho1 [query dsRNA] Tak1 [query dsRNA]

qPCR.target a factor with levels CG10417 CG13197 CG9391 egr lic PRL-1 Rho1 Tak1 passage a factor with levels passage 4 passage 42 RNAi a numeric vector

#### Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S2.

```
data(mRNAdoubleKDefficiency)
head(mRNAdoubleKDefficiency)
```

14 Networks

```
mRNAsingleKDefficiency
```

mRNA levels for single gene knock downs

# **Description**

qPCR measurements for the mRNA level after a single gene knock down (ratio relative to wild type control). The experiments is done for two independent designs of RNAi reagents.

## Usage

```
data(mRNAsingleKDefficiency)
```

#### **Format**

A data frame with 89 observations on the following 5 variables.

```
Symbol a character vector

MeanDesign1 a numeric vector

StderrDesign1 a numeric vector

MeanDesign2 a numeric vector

StderrDesign2 a numeric vector
```

#### Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S1.

## **Examples**

```
data(mRNAsingleKDefficiency)
head(mRNAsingleKDefficiency)
```

Networks

Knock interaction networks

# **Description**

This dataset is a subset of the DroID database. It contains the known (genetic) interactions between the genes regarded in the main screen.

## Usage

```
data(Networks)
```

pathwayMembership 15

## **Format**

A data frame with 402 observations on the following 5 variables.

```
gene1 a character vector
gene2 a character vector
correlation a numeric vector
genetic a numeric vector
human a numeric vector
```

#### Source

Data as used in Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S12.

The data is a subset from the drosophila interactions database (DroID), http://www.droidb.org, Data version 2010\_10 updated 20 October 2010.

Murali T, Pacifico S, Yu J, Guest S, Roberts GG 3rd, Finley RL Jr . DroID 2011: a comprehensive, integrated resource for protein, transcription factor, RNA and gene interactions for Drosophila. Nucleic Acids Res. 2010 Oct 29.

## **Examples**

```
data(Networks)
head(Networks)
```

pathwayMembership

Pathway membership of genes.

# **Description**

The membership of the tested genes in the four pathways JAK/STAT, RasMAPK, JNK, and p38.

# Usage

```
data(pathwayMembership)
```

## **Format**

The format is: chr "pathwayMembership"

## Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S13.

```
data(pathwayMembership)
head(pathwayMembership)
```

16 singleKDphenotype

PhysicalInteractions Known physical interactions

## **Description**

This dataset contains a collection of known physical interactions assembled from the literature. It contains the known pathway structure of the RasMAPK and the JNK pathway.

## Usage

```
data(PhysicalInteractions)
```

#### **Format**

A data frame with 29 observations on the following 2 variables.

V1 a character vector

V2 a character vector

#### Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S13.

## **Examples**

```
data(PhysicalInteractions)
head(PhysicalInteractions)
```

singleKDphenotype

Single knockdown phenotype

## **Description**

This data.frame singleKDphenotype contains a screen assessing the single knock down phenotypes (nrCells, intensity, and area) of the tested genes. singleKDphenotypeAnno is a data.frame describing the plate annotation.

# Usage

```
data(singleKDphenotype)
```

# **Format**

The format is: chr "singleKDphenotype" chr "singleKDphenotypeAnno"

singleKDphenotype 17

# Source

Horn, Sandmann, Fischer, Huber, Boutros, Mapping of Signalling Networks through Synthetic Genetic Interaction Analysis by RNAi, Nature Methods, 2011, Figure S3.

# **Examples**

data(singleKDphenotype)
head(singleKDphenotype)
head(singleKDphenotypeAnno)

# **Index**

* ~~ Optionally other standard keywords,	graphics, 12
one per line, from file	grid, <i>13</i>
KEYWORDS in the R ~~	grid.raster, 12
RNAinteractMAPK-package, 2	grid.sgiHeatmap, <i>10</i>
* ~~ documentation directory ~~	
RNAinteractMAPK-package, 2	levelplot, <i>10</i>
* datasets	
cellTiterGlo,4	MAPK.cv.classifier,7
Dme12PPMAPK, 5	MAPK.cv.TPS (MAPK.estimate.TPS), 9
dsRNAiDilutionSeries, 5	MAPK.estimate.TPS, 9
ElpB1phenotype, 6	MAPK.plot.classification
mRNAdoubleKDefficiency, 13	(MAPK.cv.classifier), 7
mRNAsingleKDefficiency, 14	MAPK.plot.heatmap.raster, 10
Networks, 14	MAPK.plot.TPS.all(MAPK.estimate.TPS), 9
pathwayMembership, 15	MAPK.plot.TPS.single
PhysicalInteractions, 16	(MAPK.estimate.TPS), 9
singleKDphenotype, 16	MAPK.predict.classification
* misc	(MAPK.cv.classifier), 7
MAPK.cv.classifier, 7	MAPK.report.gene.lists.paper,11
MAPK.estimate.TPS, 9	MAPK.screen.as.array
MAPK.plot.heatmap.raster, 10	(MAPK.estimate.TPS), 9
MAPK.report.gene.lists.paper, 11	MAPK.smooth.scatter, 12
MAPK.smooth.scatter, 12	mRNAdoubleKDefficiency, 3, 13
* package	mRNAsingleKDefficiency, $3$ , $14$
RNAinteractMAPK-package, 2	
*	Networks, $3$ , $14$
RNAinteractMAPK-package, 2	3 15
MAINTEN activity package, 2	pathwayMembership, 3, 15
cellTiterGlo, 3, 4	PhysicalInteractions, $3$ , $16$
CCIIIICCI G10, 3, 1	RNAinteract, 2, 3, 5, 7, 8, 11
Dmel2PPMAPK, 2, 5	RNAInteract, 2, 3, 3, 7, 8, 11
dsRNAiDilutionSeries, 3, 5	RNAInteractMAPK
dsRNAiDilutionSeriesAnno	(RNAinteractMAPK-package), 2
(dsRNAiDilutionSeries), 5	RNAInteractMAPK-package, 2
dsRNAiDilutionSeriesDF	KNATITUEL activary-package, 2
(dsRNAiDilutionSeries), 5	singleKDphenotype, 3, 16
(dominibilitationoci 103), 5	singleKDphenotypeAnno
ElpB1phenotype, 3, 6	(singleKDphenotype), 16
21p3 (pilotio 0)pc, 0, 0	smoothScatter, 12, 13
getData, 2	5.1100 5110 64 5 5 6 7 5 7 5 7 5 7 5 7 5 7 5 7 5 7 5 7