

Package ‘DmelSGI’

August 1, 2024

Type Package

Title Experimental data and documented source code for the paper ``A Map of Directional Genetic Interactions in a Metazoan Cell''

Version 1.36.0

Description The package contains the experimental data and documented source code of the manuscript ``Fischer et al., A Map of Directional Genetic Interactions in a Metazoan Cell, eLife, 2015, in Press.". The vignette code generates all figures in the paper.

License Artistic-2.0

LazyLoad true

Imports grid, TSP, limma, rhdf5, knitr, abind, gplots, igraph, grDevices, graphics, stats

Depends R (>= 3.0)

VignetteBuilder knitr

Suggests BiocStyle, EBImage, RColorBrewer, RNAinteractMAPK, hwriter, xtable, beeswarm

NeedsCompilation no

biocViews MicrotitrePlateAssayData, CellCulture, HighthroughputImagingData, Drosophila_melanogaster_Data, ExperimentData, HighThroughputImagingData, ReproducibleResearch

git_url <https://git.bioconductor.org/packages/DmelSGI>

git_branch RELEASE_3_19

git_last_commit ade0469

git_last_commit_date 2024-04-30

Repository Bioconductor 3.19

Date/Publication 2024-08-01

Author Bernd Fischer [aut],
Wolfgang Huber [ctb],
Mike Smith [cre]

Maintainer Mike Smith <mike.smith@embl.de>

Contents

DmelSGI-package	3
applyDimensionReduction	4
callInteractions	5
datamatrix	6
DPiM	7
estimatePairwiseInteractions	8
FBgn2anno	9
Features	10
fitepistasis	11
getBaseDir	12
grid.spider	12
hrNames	13
Interactions	14
Intogen	15
learnCoComplexFct	16
mainEffects	17
myHeatmap	19
mymedpolish	20
orderDim	21
orderSpiderAxis	22
pimatrix	23
plot2Phenotypes	24
plotHairballLabels	25
qualityControlFeature	26
qualityControlGene	26
RohnEtAl	27
selectByStability	29
SelectedClusters	30
SelectedClustersComplexes	30
SKDdata	31
stabilitySelection	32
subSampleForStabilitySelection	33
subSampleForStabilitySelectionFct	34
TID2HUGO	35
toMatrix	35
toRaster	36
trsf	37

Index

38

DmelSGI-package	<i>DmelSGI.</i>
-----------------	-----------------

Description

The package contains the data and the source code to reproduce the results and figures from the paper *title TBC*.

Details

See `vignette("DmelSGI")` for details.

Package content

See `vignette("DmelSGI")` 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 following **datasets** are provided with this package:

- **Features and quality control**
 - `Features` Description of the extracted features.
 - `qualityControlFeature` Correlation of features between replicates
 - `qualityControlGene` Correlation of interaction profiles between independent dsRNA designs
- **Stability selection**
 - `subSampleForStabilitySelection` A subsampled dataset used to select features.
 - `stabilitySelection` The features selected by stability.
- **Pairwise interaction matrix**
 - `datamatrix` Pairwise perturbation screen data
 - `pimatrix` Pairwise genetic interaction scores per experiment (no summary per gene pair)
 - `Interactions` Pairwise genetic interaction scores and p-values (summary per gene pairs)
 - `mainEffects` Main effects (single knock down effects) estimated from the combinatorial data
 - `SKDdata` Single knock down screen

Functions in this package:

- **stability selection**
 - `subSampleForStabilitySelectionFct`
 - `stabilitySelection`
 - `applyDimensionReduction`
- **pairwise interactions**
 - `estimatePairwiseInteractions`
 - `mymedpolish`
 - `callInteractions`

Author(s)

Bernd Fischer

Maintainer: Bernd Fischer <bernd.fischer@embl.de>

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.

Examples

```
data(datamatrix, package="DmelsGI")
```

```
applyDimensionReduction
```

Subsets the features in a genetic interaction dataset in HDF5 format.

Description

Subsets the features in a genetic interaction dataset in HDF5 format. The features are selected by [selectByStability](#) beforehand.

Usage

```
applyDimensionReduction(fileMatrixData, fileNew, selected,  
                        verbose = TRUE, overwrite = FALSE)
```

Arguments

fileMatrixData	(Input) data matrix in HDF5 format.
fileNew	(Output) data matrix in HDF5 format with a subset of features.
selected	The names of the selected features that will be subsetted.
verbose	Prints more output on screen.
overwrite	If TRUE, overwrite an existing output file (fileNew), otherwise stops.

Value

NULL is returned. As a side effect the HDF5 file 'fileNew' is created and data matrices with subsetted features will be written to it.

Author(s)

Bernd Fischer

See Also

[selectByStability](#), [stabilitySelection](#), [DmelsGI-package](#)

Examples

```
print(applyDimensionReduction)
```

callInteractions	<i>A statistical test to call pairwise interactions from interaction scores.</i>
------------------	--

Description

Using the four replicates per gene pair (from the two-by-two dsRNA designs) the null hypothesis the the interaction score is zero is tested by a moderated t-test (R-package limma). The p-values are adjusted by the method of Benjamini-Hochberg. The adjusted p-values and the pairwise interaction scores are stored in an HDF5 file.

Usage

```
callInteractions(filePI, fileInteractions, verbose = TRUE, overwrite = FALSE)
```

Arguments

filePI	Filename of the HDF5 file with interaction scores. (input)
fileInteractions	Filename of the HDF5 file with adjusted p-values and interaction scores per gene pair and feature (output).
verbose	Prints more output on screen.
overwrite	If TRUE, overwrite an existing output file (fileInteractions), otherwise stops.

Value

NULL is returned. As a side effect the HDF5 file 'fileInteractions' is created and adjusted p-values and pairwise interaction scores are saved to this file.

Author(s)

Bernd Fischer

References

Horn T, Sandmann T, Fischer B, Axelsson E, Huber W, Boutros M (2011). *Mapping of signaling networks through synthetic genetic interaction analysis by RNAi*. Nature Methods 8: 341-346.

See Also

[DmelSGI-package](#)

Examples

```
print(callInteractions)
```

datamatrix

Pairwise perturbation screen data

Description

An data array (D) with the feature data of the pairwise perturbation screen. A contains the annotation of genes and features

Usage

```
data(datamatrix)
```

Format

The format is:

List of 2

\$ D : num [1:1293, 1:2, 1:72, 1:2, 1:21] 14.8 14.9 14.9 14.9 14.9 ...

..- attr(*, "dimnames")=List of 5

.. ..\$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...

.. ..\$ targetDesign: chr [1:2] "1" "2"

.. ..\$ query : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...

.. ..\$ queryDesign : chr [1:2] "1" "2"

.. ..\$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...

\$ Anno:List of 5

..\$ target : 'data.frame': 1293 obs. of 6 variables:

.. ..\$ TID : chr [1:1293(1d)] "FBgn0002441" "FBgn0038760" ...

.. ..\$ TargetPlate : int [1:1293(1d)] 1 1 1 1 1 1 1 1 1 1 ...

.. ..\$ group : chr [1:1293(1d)] "sample" "sample" "sample" ...

.. ..\$ Symbol : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" ...

.. ..\$ Name : chr [1:1293(1d)] "malignant brain tumor" ...

..\$ targetDesign: 'data.frame': 2 obs. of 1 variable:

.. ..\$ design: int [1:2(1d)] 1 2

..\$ query : 'data.frame': 72 obs. of 5 variables:

.. ..\$ TID : chr [1:72(1d)] "FBgn0051156" "FBgn0041111" ...

.. ..\$ Batch : int [1:72(1d)] 1 1 1 1 1 1 2 2 2 2 ...

.. ..\$ Symbol : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...

.. ..\$ Name : chr [1:72(1d)] "-" "lilliputian" "Smg1" "Axin" ...

..\$ queryDesign : 'data.frame': 2 obs. of 1 variable:

.. ..\$ design: int [1:2(1d)] 1 2

..\$ phenotype : 'data.frame': 21 obs. of 1 variable:

.. ..\$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...

Value

An array with the phenotypic data.

See Also[DmelSGI-package](#)**Examples**

```
data(datamatrix)
str(datamatrix)
```

DPiM

*Drosophila Protein Interaction Map (DPiM)***Description**

The Drosophila Protein Interaction Map (DPiM) dataset contains of two parts: (1.) The experimentally identified protein interaction partners and (2.) inferred protein complexes.

Usage

```
data("DPiM")
```

Format

The format is: List of 2 \$ interactions: 'data.frame': 10969 obs. of 5 variables: ..\$ Interactor_1 : chr [1:10969] "FBgn0036918" "FBgn0031143" "FBgn0030086" "FBgn0015019"\$ Interactor_2 : chr [1:10969] "FBgn0037893" "FBgn0035102" "FBgn0033342" "FBgn0037632"\$ HGSCore : num [1:10969] 742 737 733 730 726\$ Evidence.in.DroID: chr [1:10969] "human_orthology; yeast_orthology" "" "human_orthology" "human_orthology; yeast_orthology"\$ evidence : logi [1:10969] TRUE FALSE TRUE TRUE TRUE TRUE TRUE ... \$ complexes :List of 556 ..\$: chr [1:40] "FBgn0002284" "FBgn0002787" "FBgn0004066" "FBgn0010590"\$: chr [1:43] "FBgn0000212" "FBgn0000499" "FBgn0001276" "FBgn0001324"\$: chr [1:28] "FBgn0003660" "FBgn0011288" "FBgn0011708" "FBgn0013343"\$: chr [1:11] "FBgn0022023" "FBgn0025582" "FBgn0029629" "FBgn0033902" [list output truncated]

Value

The mass spec data and the inferred protein complexes of the DPiM dataset.

References

Guruharsha, K. G., et al. "A protein complex network of Drosophila melanogaster." Cell 147.3 (2011): 690-703.

See Also[DmelSGI-package](#)**Examples**

```
data(DPiM)
```

`estimatePairwiseInteractions`*Estimates pairwise interaction scores.*

Description

Estimates pairwise interaction scores for a large, multi-dimensional combinatorial screen.

Usage

```
estimatePairwiseInteractions(fileMatrixData,  
                             filePI,  
                             verbose = TRUE,  
                             overwrite = FALSE,  
                             useSKD = TRUE)
```

Arguments

<code>fileMatrixData</code>	Filename of HDF5 file containing the combinatorial screening data (input).
<code>filePI</code>	Filename of HDF5 file to save the pairwise interaction matrix (output).
<code>verbose</code>	Prints more output on screen.
<code>overwrite</code>	If TRUE, overwrite an existing output file (<code>filePI</code>), otherwise stops.
<code>useSKD</code>	If TRUE, the negative controls are used to estimate the overall effect.

Details

Estimates the pairwise interaction scores for each feature and each batch by calling the function [mymedpolish](#).

Value

Returns TRUE. As a sideeffect, the array with interaction scores is stored in the HDF5 file 'filePI'.

Author(s)

Bernd Fischer

References

Horn T, Sandmann T, Fischer B, Axelsson E, Huber W, Boutros M (2011). *Mapping of signaling networks through synthetic genetic interaction analysis by RNAi*. Nature Methods 8: 341-346.

See Also

[mymedpolish](#), [DmelSGI-package](#)

Examples

```
print(estimatePairwiseInteractions)
```

FBgn2anno	<i>GO annotation of flybase genes</i>
-----------	---------------------------------------

Description

GO annotation of the 1367 genes targeted in this screen.

Usage

```
data(FBgn2anno)
```

Format

A data frame with 118863 observations on the following 4 variables.

source a character vector
gene_id a character vector
Category a character vector
Name a character vector

Value

GO annotation.

See Also

[DmelSGI-package](#)

Examples

```
data(FBgn2anno)
```

Features	<i>Description of the extracted features.</i>
----------	---

Description

The data.frame describing all features extracted from the images. Beside the name of the feature given as the row.name of the data.frame, 7 columns describe each extracted feature:

- **mag** is the magnification of the image that is used (either 4x or 10x)
- **summary** Features are extracted for each single cell. Features are summarized by mean, standard deviation, quantiles, and histograms.
- **mask** The segmentation mask used to extract the features, e.g. the nuclei, the nuclei in the pH3 channel, or the cell body extracted from the a-tubulin channel
- **channel** The channel used to extract the feature
- **set** The feature category
- **type** The type of feature (area, intensity, ...)
- **param** Additional parameters of the feature extraction (quantiles, histogram bins, ...)

Usage

```
data(Features)
```

Format

The head of the data.frame is:

```
mag summary mask channel set type param
4x.count "4x" "nrNuclei" "nucleus" "DAPI" "M" "" ""
4x.countpH3 "4x" "nrNuclei" "mitoticNuclei" "pH3" "M" "" ""
4x.isMitotic "4x" "nrNuclei" "nucleus" "DAPI" "M" "" ""
4x.ratioMitotic "4x" "mitoticRatio" "nucleus" "DAPI" "M" "" ""
4x.areaNuc "4x" "mean" "nucleus" "DAPI" "M" "area" ""
4x.areaNucSD "4x" "stddev" "nucleus" "DAPI" "M" "area" ""
```

Value

A data.frame listing all extracted features.

See Also

[DmelSGI-package](#)

Examples

```
data(Features)
head(Features)
```

fitepistasis

*Fit of the pi-score vectors as a function of main effects***Description**

The output of the linear fit of the pi-score vectors as a function of main effects. The list contains four datasets:

- A 6-dimensional array of the original data with pi-scores and main effects,
- the coefficients of the linear fit,
- the p-values from Anova and
- the variance explained by the main effects.

Usage

```
data(fitepistasis)
```

Format

The format is:

List of 2

```
$ Coef: num [1:3, 1:1293, 1:2, 1:72, 1:2] -0.0464 0.3702 -0.0475 -0.0794 ...
..- attr(*, "dimnames")=List of 5
.. ..$: chr [1:3] "const" "xt" "xq"
.. ..$: chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
.. ..$: chr [1:2] "1" "2"
.. ..$: chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
.. ..$: chr [1:2] "1" "2"
$ Sq : num [1:3, 1:1293, 1:2, 1:72, 1:2] 0.18515 0.00207 0.45705 0.15047 ...
..- attr(*, "dimnames")=List of 5
.. ..$: chr [1:3] "xt" "xq" "res"
.. ..$: chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
.. ..$: chr [1:2] "1" "2"
.. ..$: chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
.. ..$: chr [1:2] "1" "2"
```

Value

The output of the epistasis estimation

See Also

[DmelSGI-package](#)

Examples

```
data(fitepistasis)
```

getBaseDir	<i>Returns the base directory for the vignette.</i>
------------	---

Description

Returns the base directory for the vignette. When knitr is applied on the main vignette, this function ensures that the subvignettes get knowledge of the base directory.

Usage

```
getBaseDir(default = ".")
```

Arguments

default	The default base directory.
---------	-----------------------------

Value

Returns a character with the directory name

Author(s)

Bernd Fischer

See Also

[DmelSGI-package](#)

Examples

```
getBaseDir()
```

grid.spider	<i>Spider plot</i>
-------------	--------------------

Description

A grid function to draw a spider plot.

Usage

```
grid.spider(v, col, col.arms = "black", dlim = NULL)
grid.spider.legend(vn, col.arms = "black", dlim = NULL)
```

Arguments

v	A vector of numbers to be presented in the spider plot.
vn	A vector of dimension names that are represented by the spider arms. Has the same length as v.
col	The color of the polygon area.
col.arms	The color of the background of the spider arms.
dlim	A vector with two values. Limits of the spider arm axis.

Details

These function draw a grid spider plot or a legend for the spider arms.

Value

Both functions return an invisible NULL, but they have an side-effect that draws a spider plot using grid.

Author(s)

Bernd Fischer

See Also

[orderSpiderAxis](#), [DmelSGI-package](#)

Examples

```
print(grid.spider)
print(grid.spider.legend)
```

hrNames

Human readable feature names

Description

Translate feature names to human readable feature names. Names not known to this function for conversion are returned unchanged.

Usage

```
hrNames(names)
```

Arguments

names	Original feature names.
-------	-------------------------

Value

A vector of translated feature names.

Author(s)

Bernd Fischer

See Also

[DmelSGI-package](#)

Examples

```
hrNames(c("4x.count", "4x.ratioMitotic"))
```

Interactions	<i>Pairwise genetic interaction scores and p-values (summary per gene pairs)</i>
--------------	--

Description

Two arrays are provided in this dataset: The pairwise interaction scores summarized per gene pair and the respective adjusted p-values. p-values are computed by a moderated t-test (limma) and corrected for multiple testing by the method of Benjamini-Hochberg. The list (Anno) contains the annotation of the target genes, query genes, and features. See [pimatrix](#) for interaction scores that are not yet summarized per gene pair.

Usage

```
data(Interactions)
```

Format

The format is:

List of 3

```
$ pscore: num [1:1293, 1:72, 1:21] -0.0866 -0.0924 -0.0707 -0.0878 -0.0587 ...
```

```
..- attr(*, "dimnames")=List of 3
```

```
.. ..$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
```

```
.. ..$ query : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
```

```
.. ..$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...
```

```
$ padj : num [1:1293, 1:72, 1:21] 0.232 0.249 0.286 0.281 0.369 ...
```

```
..- attr(*, "dimnames")=List of 3
```

```
.. ..$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
```

```
.. ..$ query : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
```

```
.. ..$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...
```

```
$ Anno :List of 3
```

```
..$ target :'data.frame': 1293 obs. of 6 variables:
```

```
.. ..$ TID : chr [1:1293(1d)] "FBgn0002441" "FBgn0038760" ...
.. ..$ TargetPlate : int [1:1293(1d)] 1 1 1 1 1 1 1 1 1 ...
.. ..$ group : chr [1:1293(1d)] "sample" "sample" "sample" ...
.. ..$ Symbol : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" ...
.. ..$ Name : chr [1:1293(1d)] "lethal (3) malignant brain tumor" ...
..$ query : 'data.frame': 72 obs. of 5 variables:
.. ..$ TID : chr [1:72(1d)] "FBgn0051156" "FBgn0041111" ...
.. ..$ Batch : int [1:72(1d)] 1 1 1 1 1 1 2 2 2 ...
.. ..$ Symbol : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
.. ..$ Name : chr [1:72(1d)] "-" "lilliputian" "Smg1" "Axin" ...
..$ phenotype: 'data.frame': 21 obs. of 1 variable:
.. ..$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...
```

Value

An object containing the pi-scores, the adjusted p-values and the annotation of the statistical genetic interactions.

See Also

[pimatrix,DmelSGI-package](#)

Examples

```
data(Interactions)
str(Interactions)
```

Intogen	<i>Intogen: Interactive Onco Genomics</i>
---------	---

Description

A list of recurrently mutated genes from Intogen.

Usage

```
data("Intogen")
```

Format

A data frame with 2933 observations on the following 10 variables.

- gene a character vector
- symbol a character vector
- project.name a character vector
- mut.freq a numeric vector
- MuSiC a character vector

oncodriveFM a character vector
oncodriveCLUST a character vector
ActiveDriver a character vector
MutSig a character vector
driver.category a character vector

Value

The Intogen dataset.

References

Tamborero, David, et al. "Comprehensive identification of mutational cancer driver genes across 12 tumor types." Scientific reports 3 (2013).

See Also

[DmelSGI-package](#)

Examples

data(Intogen)

learnCoComplexFct	<i>Learning the co-complex function</i>
-------------------	---

Description

Learning the co-complex function from a correlation matrix.

Usage

learnCoComplexFct(C, ProteinComplexes)
convertCorrelations(C, coComplexFct)

Arguments

- | | |
|------------------|--|
| C | A genes-by-genes correlation matrix (or matrix containing any other pairwise scores). |
| ProteinComplexes | A list of protein complexes. Each element of the list is one protein complex and contains a data.frame with at least one column gene_id. |
| coComplexFct | coComplexFct is an object containing the co-complex function as returned by learnCoComplexFct. |

Details

An empirical density function of the values in C is computed once for the gene pairs that are co-member of at least one protein complex and once for all other gene pairs. The definition of protein complexes that is used to learn the co-complex function is taken from ProteinComplexes.

Value

learnCoComplexFct will provide an object that contains the empirical density function of correlation.

convertCorrelations will provide a matrix of co-complex scores.

Author(s)

Bernd Fischer

References

A similar approach is used in Ryan, C.J., et al. (2012). Hierarchical modularity and the evolution of genetic interactomes across species. Molecular cell 46, 691-704.

See Also

[DmelSGI-package](#)

Examples

```
print(learnCoComplexFct)
print(convertCorrelations)
```

mainEffects	<i>Main effects (single knock down effects) estimated from the combinatorial data</i>
-------------	---

Description

The overall effect (effect of the negative control experiments), and the estimated main effects (single knock down effects) for template and query genes. Overall effects and template main effects are estimated separately for each batch (1:12). Query main effects are estimated separately for each template plate (1:4)

Usage

```
data(mainEffects)
```

Format

The format is:

List of 4

\$ overall: num [1:2, 1:12, 1:21] 14.6 14.6 14 14 14.7 ...

.. attr(*, "dimnames")=List of 3

.. ..\$ targetDesign: chr [1:2] "1" "2"

.. ..\$ batch : chr [1:12] "1" "2" "3" "4" ...

.. ..\$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...

\$ query : num [1:72, 1:2, 1:4, 1:21] 0.0409 0.0192 -0.587 -0.0636 -0.4709 ...

.. attr(*, "dimnames")=List of 4

.. ..\$ query : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...

.. ..\$ queryDesign: chr [1:2] "1" "2"

.. ..\$ templatePlate: chr [1:4] "1" "2" "3" "4"

.. ..\$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...

\$ target : num [1:1293, 1:2, 1:12, 1:21] 0.0921 0.3444 0.1603 0.2252 0.1293 ...

.. attr(*, "dimnames")=List of 4

.. ..\$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...

.. ..\$ targetDesign: chr [1:2] "1" "2"

.. ..\$ batch : chr [1:12] "1" "2" "3" "4" ...

.. ..\$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...

\$ Anno :List of 7

..\$ target : 'data.frame': 1293 obs. of 6 variables:

.. ..\$ TID : chr [1:1293(1d)] "FBgn0002441" "FBgn0038760" ...

.. ..\$ TargetPlate : int [1:1293(1d)] 1 1 1 1 1 1 1 1 1 1 ...

.. ..\$ group : chr [1:1293(1d)] "sample" "sample" "sample" ...

.. ..\$ Symbol : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" ...

.. ..\$ Name : chr [1:1293(1d)] "lethal (3) malignant brain tumor"...

..\$ targetDesign : 'data.frame': 2 obs. of 1 variable:

.. ..\$ design: int [1:2(1d)] 1 2

..\$ query : 'data.frame': 72 obs. of 5 variables:

.. ..\$ TID : chr [1:72(1d)] "FBgn0051156" "FBgn0041111" ...

.. ..\$ Batch : int [1:72(1d)] 1 1 1 1 1 1 2 2 2 2 ...

.. ..\$ Symbol : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...

.. ..\$ Name : chr [1:72(1d)] "-" "lilliputian" "Smg1" "Axi" ...

..\$ queryDesign : 'data.frame': 2 obs. of 1 variable:

.. ..\$ design: int [1:2(1d)] 1 2

..\$ phenotype : 'data.frame': 21 obs. of 1 variable:

.. ..\$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...

..\$ batch : 'data.frame': 12 obs. of 1 variable:

.. ..\$ batch: int [1:12] 1 2 3 4 5 6 7 8 9 10 ...

..\$ templatePlate : 'data.frame': 4 obs. of 1 variable:

.. ..\$ templatePlate: int [1:4] 1 2 3 4

Value

An object containing the query and target main effects.

See Also[DmelSGI-package](#)**Examples**

```
data(mainEffects)
str(mainEffects)
```

myHeatmap	<i>Draws a heatmap for a three dimensional array</i>
-----------	--

Description

Draws a heatmap for a three dimensional array as e.g. the three dimensional genetic interaction cube.

Usage

```
myHeatmap(x, cuts, col, fontsize = 18, colnames = TRUE, rownames = FALSE)
```

Arguments

x	A three dimensional array.
cuts	break points for mapping the values in x to col. The length of cuts is one larger than the length of col.
col	A color bar as returned by colorRampPalette .
fontsize	The size of the text labels.
colnames	Logical. If TRUE, the column names are printed.
rownames	Logical. If TRUE, the row names are printed.

Details

This function is used to draw the heatmap of the three dimensional genetic interaction cube.

Value

Nothing is returned, but the function plots a heatmap as a site-effect.

Author(s)

Bernd Fischer and Wolfgang Huber

See Also[DmelSGI-package](#)**Examples**

```
print(myHeatmap)
```

mymedpolish

A variant of the medpolish function.

Description

A variant of the R-implementation of [medpolish](#). Fits an additive model using Tukey's median polish procedure. The variant uses negative controls to estimate the overall effect and it estimates column effects separately for each batch of rows indicated by TP (for template plate). **It is highly recommended to use the original function from the R-package stats.** The function is used by [estimatePairwiseInteractions](#).

Usage

```
mymedpolish(x, TP, TemplateNeg, QueryNeg, eps = 1e-04, maxiter = 100, na.rm = TRUE)
```

Arguments

x	a numeric matrix.
TP	integer. Specifying the plate number for each row element (template genes). Counting starts with 1. Column effects (query genes) will be estimated for each template plate separately
TemplateNeg	Index of negative controls on template plate
QueryNeg	Index of negative controls in the columns (queries)
eps	real number greater than 0. A tolerance for convergence: see medpolish .
maxiter	the maximum number of iterations
na.rm	logical. Should missing values be removed?

Value

neg	the fitted constant term.or overall effect representing the effect of negative controls
templateMainEffect	the fitted row effects representing the single knock down effects of the template genes
queryMainEffect	the fitted column effects representing the single knock down effects of the query genes. It is a matrix with dimensions query genes x template plates
pi	the residuals which are the pairwise interaction scores

Author(s)

Original implementation in the R package stats (See [medpolish](#)). Changes for estimating pairwise interaction scores by Bernd Fischer.

References

Tukey, J. W. (1977). *Exploratory Data Analysis*, Reading Massachusetts: Addison-Wesley.
Horn T, Sandmann T, Fischer B, Axelsson E, Huber W, Boutros M (2011). *Mapping of signaling networks through synthetic genetic interaction analysis by RNAi*. Nature Methods 8: 341-346.

See Also

[medpolish](#), [estimatePairwiseInteractions](#), [DmelSGI-package](#)

Examples

```
print(mymedpolish)
```

orderDim	<i>Orders a three dimensional array along one dimension</i>
----------	---

Description

Orders a three dimensional array along one dimension.

Usage

```
orderDim(x, i)
```

Arguments

x	A three dimensional array.
i	The array is ordered along the i-th dimension.

Details

The three dimensional array is ordered along the i-th dimension.

Value

An integer vector with the ordering.

Author(s)

Bernd Fischer

See Also

[DmelSGI-package](#)

Examples

```
print(orderDim)
```

orderSpiderAxis	<i>Orders the axis of a spider plot.</i>
-----------------	--

Description

Solves a traveling salesperson problem to optimally order the arms of a spider plot.

Usage

```
orderSpiderAxis(X)
```

Arguments

X	A d x n matrix of values for n instances that are plotted in the spider plots with d arms.
---	--

Details

The arms are ordered such that two neighboring spider arms are similar to each other.

Value

An integer vector with an optimal order of the spider arms.

Author(s)

Bernd Fischer

See Also

[grid.spider](#), [DmeISGI-package](#)

Examples

```
print(orderSpiderAxis)
```

pimatrix	<i>Pairwise genetic interaction scores per experiment (no summary per gene pair)</i>
----------	--

Description

An array (D) with pairwise interaction scores. The interaction scores are given per experiment are not yet summarized per gene pair. See [Interactions](#) for interactions summarized per gene pair. A contains the annotation of genes and features.

Usage

```
data(pimatrix)
```

Format

The format is:

List of 2

```
$ D : num [1:1293, 1:2, 1:72, 1:2, 1:21] -0.1086 -0.1799 -0.045 -0.0664 ...
```

```
..- attr(*, "dimnames")=List of 5
```

```
.. ..$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
```

```
.. ..$ targetDesign: chr [1:2] "1" "2"
```

```
.. ..$ query : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
```

```
.. ..$ queryDesign : chr [1:2] "1" "2"
```

```
.. ..$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...
```

```
$ Anno:List of 5
```

```
..$ target : 'data.frame': 1293 obs. of 6 variables:
```

```
.. ..$ TID : chr [1:1293(1d)] "FBgn0002441" "FBgn0038760" ...
```

```
.. ..$ TargetPlate : int [1:1293(1d)] 1 1 1 1 1 1 1 1 1 1 ...
```

```
.. ..$ group : chr [1:1293(1d)] "sample" "sample" "sample" ...
```

```
.. ..$ Symbol : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" ...
```

```
.. ..$ Name : chr [1:1293(1d)] "lethal (3) malignant brain tumor"...
```

```
..$ targetDesign: 'data.frame': 2 obs. of 1 variable:
```

```
.. ..$ design: int [1:2(1d)] 1 2
```

```
..$ query : 'data.frame': 72 obs. of 5 variables:
```

```
.. ..$ TID : chr [1:72(1d)] "FBgn0051156" "FBgn0041111" ...
```

```
.. ..$ Batch : int [1:72(1d)] 1 1 1 1 1 1 2 2 2 2 ...
```

```
.. ..$ Symbol : chr [1:72(1d)] "CG31156" "lilli" "Smg1" "Axn" ...
```

```
.. ..$ Name : chr [1:72(1d)] "-" "lilliputian" "Smg1" "Axin" ...
```

```
..$ queryDesign : 'data.frame': 2 obs. of 1 variable:
```

```
.. ..$ design: int [1:2(1d)] 1 2
```

```
..$ phenotype : 'data.frame': 21 obs. of 1 variable:
```

```
.. ..$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...
```

Value

The matrix of pairwise interaction scores per RNAi design.

See Also

[Interactions,DmeISGI-package](#)

Examples

```
data(pimatrix)
str(pimatrix)
```

plot2Phenotypes	<i>Plot directed epistatic interactions.</i>
-----------------	--

Description

Plots the data to estimate a directed epistatic interactions.

Usage

```
plot2Phenotypes(X, gt, gq, f1, f2, length = 1, ...)
plotPIdata(X, gt, gq, show = "summary", ...)
```

Arguments

X	A 6-dimensional array (phenotype x [xt, xq, pi] x target genes x targetDesigns x query genes x queryDesigns).
gt	The target gene name.
gq	The query gene name.
f1	The first phenotypic feature.
f2	The second phenotypic feature.
length	Length of arrow head.
show	Either show='summary' to show the mean over all dsRNA designs or it is a vector of length 2 that specifies the two dsRNAs to show.
...	Other arguments passed to plot.

Details

plot2Phenotypes shows a plot showing the two phenotypes on the axis. 2 x 2 arrows for the single gene effects of the two dsRNA designs for the two genes are shown in green and purple, the expected double knock-down effects under the non-interacting model for the 4 combinations of dsRNA designs in gray, and the four measured double knock-down effects in black. The black arrows are the genetic interactions.

plotPIdata shows two scatter plots for the fit of the vector of pairwise interaction scores across all phenotypes as a function of the single gene effects. Each dot represents one phenotype.

Value

Nothing is returned, but the function draws a plot as a site-effect.

Author(s)

Bernd Fischer

See Also

[DmelSGI-package](#)

Examples

```
print(plot2Phenotypes)
print(plotPIdata)
```

plotHairballLabels	<i>Adds the cluster labels to a graph</i>
--------------------	---

Description

Adds the cluster labels to a graph

Usage

```
plotHairballLabels(g, co, Labels, Col)
```

Arguments

g	An igraph object for the graph.
co	A n x 2 matrix of layout coordinates as returned by the igraph layout algorithms.
Labels	A list with a vector of gene names per cluster as they appear as vertexes in the igraph objects. The list element names are printed to the graph plot.
Col	The colors of the cluster names.

Value

Nothing is returned, but as a side-effect, the labels for the hairball are added to a plot.

Author(s)

Bernd Fischer

See Also

[DmelSGI-package](#)

Examples

```
print(plotHairballLabels)
```

qualityControlFeature *Correlation of features between replicates*

Description

The quality control of features is described by three vectors: The correlation between two replicates, the fraction of finite values, and a logical vector indicating which feature passed the quality control. The features are described in the dataset [Features](#).

Usage

```
data(qualityControlFeature)
```

Format

List of 3 \$ correlation : num [1:328] 0.933 0.927 0.927 0.922 0.97 ... \$ ratioFiniteValues: num [1:328] 0.999 0.999 0.999 0.999 0.999 ... \$ passed : logi [1:328] TRUE TRUE TRUE TRUE TRUE TRUE ...

Value

A data.frame containing the output of the feature quality control.

See Also

[DmelSGI-package](#)

Examples

```
data(qualityControlFeature, package="DmelSGI")
str(qualityControlFeature)
```

qualityControlGene *Correlation of interaction profiles between independent dsRNA designs*

Description

The quality control of dsRNA designs of the template genes is described by the correlation of the interaction profile between two independent dsRNA designs. In addition the annotation of the genes and a logical vector indicating which gene passed the quality control is given.

Usage

```
data(qualityControlGene)
```

Format

The format is:

List of 3

\$ correlation: num [1:1463] 0.876 0.877 0.897 0.889 0.865 ...

\$ Annotation : 'data.frame': 1463 obs. of 6 variables:

..\$ TID : chr [1:1463(1d)] "FBgn0002441" "FBgn0038760" ...

..\$ TemplatePlate : int [1:1463(1d)] 1 1 1 1 1 1 1 1 1 ...

..\$ group : chr [1:1463(1d)] "sample" "sample" "sample" ...

..\$ Symbol : chr [1:1463(1d)] "l(3)mbt" "MED25" "CG31156" ...

..\$ Name : chr [1:1463(1d)] "lethal (3) malignant brain tumor"...

\$ passed : logi [1:1463] TRUE TRUE TRUE TRUE TRUE TRUE ...

Value

A dataset with the output of the gene quality control.

See Also

[DmelSGI-package](#)

Examples

```
data(qualityControlGene, package="DmelSGI")
str(qualityControlGene)
```

RohnEtAl

Fly RNAi phenotype data

Description

This dataset is a list of RNAi Phenotypes captures in Drosophila S2R+ cells.

Usage

```
data("RohnEtAl")
```

Format

A data frame with 556 observations on the following 29 variables.

Primer a character vector

Computed.Target a character vector

Symbol a character vector

Decreased.cell.size a numeric vector

Increased.cell.size a numeric vector

Cell.shape.variable a numeric vector

Cell.shape.round.or.non.adherent a numeric vector
 Cell.shape.processes.or.spiky.or.stretchy a numeric vector
 Disorganised.peripheral.actin a numeric vector
 Increased.number.of.actin.stress.fibres a numeric vector
 Increased.number.of.actin.puncta.or.dots a numeric vector
 Asymmetric.lamellae a numeric vector
 Decreased.level.of.actin a numeric vector
 Increased.level.of.actin a numeric vector
 Increased.cytoplasmic.actin a numeric vector
 Decreased.peripheral.actin a numeric vector
 Increased.peripheral.actin a numeric vector
 Increased.nuclear.actin a numeric vector
 Microtubule.clumps a numeric vector
 Microtubules.disorganised a numeric vector
 Microtubule.processes a numeric vector
 Decreased.level.of.microtubules a numeric vector
 Increased.level.of.microtubules a numeric vector
 Increased.number.of.multinucleate.cells a numeric vector
 Increased.DNA.area a numeric vector
 No.cells a numeric vector
 Decreased.cell.number a numeric vector
 Loss.of.cell.monolayer a numeric vector
 Multiple.layers.of.cells a numeric vector

Value

A list of phenotypes. The genes showing the respective phenotype are listed in the vector for each phenotype.

References

Rohn, Jennifer L., et al. "Comparative RNAi screening identifies a conserved core metazoan actinome by phenotype." *The Journal of cell biology* 194.5 (2011): 789-805.

See Also

[DmelSGI-package](#)

Examples

```
data(RohnEtAl)
```

selectByStability	Select features by stability.
-------------------	-------------------------------

Description

Features are selected in a greedy manner. A linear model is fitted to estimate the remove the contribution of each features that can already be explained by previously selected features. The next features is selected such, that the residuals are maximal correlated between replicates.

Usage

```
selectByStability(subsample,
                  preselect = c("4x.count",
                                "4x.ratioMitotic",
                                "10x.meanNonmitotic.cell.0.s.area"),
                  Rdim = 40,
                  verbose = TRUE)
```

Arguments

subsample	A list with subsampled data as produced by subSampleForStabilitySelectionFct . <code>data(subSampleForStabilitySelection, package='Dm1SGI')</code> provides the dataset used in the paper.
preselect	The names of features that are preselected, e.g. the features '4x.count' and '4x.ratioMitotic' should be selected, because they have a special interpretation.
Rdim	The maximum number of selected features.
verbose	If TRUE, more output is provided.

Value

selected	The names of the selected features. Rdim features will be reported.
correlation	The correlation of the residual features.
ratioPositive	The fraction of positively correlated residual features in each step of the selection process. The features with ratioPositive > 0.5 should be selected.
correlationAll	The correlation of the all residual features in each step of the selection process.

Author(s)

Bernd Fischer

See Also

[Dm1SGI-package](#)

Examples

```
print(selectByStability)
```

SelectedClusters	<i>Selected processes displayed on the hairball</i>
------------------	---

Description

A list of gene sets that are displayed on the hairball.

Usage

```
data(SelectedClusters)
```

Format

The format is:

List of 24

\$ SWI/SNF : chr [1:6] "Bap60" "brm" "dalao" ...

\$ Condensin/Cohesin : chr [1:5] "SMC2" "Cap-D2" "glu" ...

\$ Cytokinesis : chr [1:12] "pav" "sqh" "rok" ...

...

Value

A list with selected clusters for visualization.

See Also

[DmelSGI-package](#)

Examples

```
data(SelectedClusters)
```

SelectedClustersComplexes	<i>Selected complexes</i>
---------------------------	---------------------------

Description

Manually curated protein complexes.

Usage

```
data(SelectedClustersComplexes)
```

Format

The format is:
List of 27
\$ DREAM complex : chr [1:5] "Caf1" "mip120" "mip130" "mip40" ...
\$ Condensin/Cohesin : chr [1:5] "SMC2" "Cap-D2" "glu" "eco" ...
\$ Apc/C : chr [1:9] "APC10" "Cdc23" "Cdc16" "shtd" ...
...

Value

A list of gene sets that represent well-defined protein complexes.

See Also

[DmelSGI-package](#)

Examples

```
data(SelectedClustersComplexes)
```

SKDdata	<i>Single knock down screen</i>
---------	---------------------------------

Description

D is the single knock down screen data for the 12 negative control query genes. The annotation of each dimension of D is provided is the list A.

Usage

```
data(SKDdata)
```

Format

The format is: List of 2 \$ D : num [1:1293, 1:2, 1:12, 1:21] 14.7 15 14.8 14.8 14.8 ...
..- attr(*, "dimnames")=List of 4
.. ..\$ target : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" "CG6833" ...
.. ..\$ targetDesign: chr [1:2] "1" "2"
.. ..\$ batch : chr [1:12] "1" "2" "3" "4" ...
.. ..\$ phenotype : chr [1:21] "4x.count" "4x.ratioMitotic" ...
\$ Anno:List of 4
..\$ target :'data.frame': 1293 obs. of 6 variables:
.. ..\$ TID : chr [1:1293(1d)] "FBgn0002441" "FBgn0038760" ...
.. ..\$ TargetPlate : int [1:1293(1d)] 1 1 1 1 1 1 1 1 1 1 ...
.. ..\$ group : chr [1:1293(1d)] "sample" "sample" "sample" ...
.. ..\$ Symbol : chr [1:1293(1d)] "l(3)mbt" "MED25" "CG31156" ...
.. ..\$ Name : chr [1:1293(1d)] "lethal (3) malignant brain tumor"...

```

..$ targetDesign:'data.frame': 2 obs. of 1 variable:
.. ..$ design: int [1:2(1d)] 1 2
..$ batch :'data.frame': 12 obs. of 1 variable:
.. ..$ batch: int [1:12] 1 2 3 4 5 6 7 8 9 10 ...
..$ phenotype :'data.frame': 21 obs. of 1 variable:
.. ..$ phenotype: chr [1:21] "4x.count" "4x.ratioMitotic" ...

```

Value

A dataset with the single knockdown data.

See Also

[DmelSGI-package](#)

Examples

```

data(SKDdata)
str(SKDdata)

```

stabilitySelection	<i>The features selected by stability.</i>
--------------------	--

Description

The features selected by stability used in the selection process are available in this dataset. Furthermore, it contains the correlation of the residual features and the fraction of positive correlated features that is used as a stop criterion.

Usage

```
data(stabilitySelection)
```

Format

The format is:
List of 4
\$ selected : chr [1:50] "4x.count" "4x.ratioMitotic" ...
\$ correlation : num [1:50] 0.912 0.868 0.559 ...
\$ ratioPositive : num [1:50] 1 0.957 0.95 ...
\$ correlationAll:List of 50
..\$: Named num [1:162] 0.912 0.946 0.948 ...
.. ..- attr(*, "names")= chr [1:162] "4x.count" "4x.countpH3" ...
..\$: Named num [1:161] 0.946 0.795 0.868 ...
.. ..- attr(*, "names")= chr [1:161] "4x.countpH3" "4x.isMitotic" ...
..\$: Named num [1:160] 0.946 0.129 0.927 ...
.. ..- attr(*, "names")= chr [1:160] "4x.countpH3" "4x.isMitotic" ...
...

Value

An object containing the output of the feature selection.

See Also

[DmelSGI-package](#)

Examples

```
data(stabilitySelection)
str(stabilitySelection)
```

```
subSampleForStabilitySelection
```

A subsampled dataset for use stability selection.

Description

This dataset contains a subsample of the interaction screen for use in the function [selectByStability](#). It contains the data matrix D with 3000 experiments x 2 replicates (dsRNA designs) x 162 features.

Usage

```
data(subSampleForStabilitySelection)
```

Format

The format is:
 List of 3
 \$ D : num [1:3000, 1:2, 1:162] -1.447 0.351 0.44 ...
 \$ Sample : int [1:3000] 77465 94252 95176 ...
 \$ phenotype: chr [1:162(1d)] "4x.count" "4x.countpH3" ...

Value

The dataset of samples used for feature selection.

See Also

[selectByStability](#), [subSampleForStabilitySelectionFct](#), [DmelSGI-package](#)

Examples

```
data(subSampleForStabilitySelection)
str(subSampleForStabilitySelection)
```

`subSampleForStabilitySelectionFct`*Subsampling the data for stability selection.*

Description

The data is subsampled to reduce computation time and memory demand for stability selection. 10000 experiments are selected by chance to estimate the most stable directions.

Usage

```
subSampleForStabilitySelectionFct(fileMatrixData, N = 10000, random.seed = NULL)
```

Arguments

<code>fileMatrixData</code>	Filename of the HDF5 file containing the data array
<code>N</code>	Number of experiments used for stability selection
<code>random.seed</code>	If not NULL, the random.seed is set before sampling, to generate a reproducible analysis script

Details

For each dsRNA design, query gene, and feature, the median value is subtracted and the data are divided by the median deviation.

Value

<code>D</code>	An array of dimension $N \times 2 \times F$, where N is the given number of experiments used, 2 represents the two query dsRNA designs, and F is the number of features.
<code>Sample</code>	The index of the sampled elements.
<code>phenotype</code>	The name of the features.

Author(s)

Bernd Fischer

See Also

[selectByStability](#), [subSampleForStabilitySelection](#), [DmelsGI-package](#)

Examples

```
print(subSampleForStabilitySelectionFct)
```

TID2HUGO

*Mapping of flygene names to human.***Description**

Mapping of flybase gene identifier to their human orthologues. It is a one-to-many mapping.

Usage

```
data(TID2HUGO)
```

Format

The format is:

List of 1293

```
$ FBgn0002441: chr [1:3] "L3MBTL1" "L3MBTL3" "L3MBTL4"
```

```
$ FBgn0038760: chr "MED25"
```

```
$ FBgn0051156: chr "SRBD1"
```

```
... [list output truncated]
```

```
- attr(*, "dim")= int 1293
```

```
- attr(*, "dimnames")=List of 1
```

```
..$ : chr [1:1293] "FBgn0002441" "FBgn0038760" "FBgn0051156" "FBgn0036405" ...
```

Value

A list containing the conversion of gene identifier from fly to human.

See Also

[DmelSGI-package](#)

Examples

```
data(TID2HUGO)
```

toMatrix

*Flattens a three dimensional array to a two dimensional matrix***Description**

Flattens a three dimensional array to a two dimensional matrix.

Usage

```
toMatrix(x)
```

Arguments

x A three dimensional array.

Value

A matrix.

See Also

[DmeISGI-package](#)

Examples

```
print(toMatrix)
```

toRaster	<i>Converts a real valued matrix in a matrix of color codes printable by grid.raster</i>
----------	--

Description

A matrix of real values in a matrix are converted in a matrix of RGB values that can be printed by [grid.raster](#).

Usage

```
toRaster(x, cuts, col)
```

Arguments

x A real valued matrix.

cuts Break points for the color values. Length of cuts has to be the length of col plus one.

col A vector of color values, e.g. as produced by [colorRampPalette](#).

Value

Returns a matrix of RGB color values that can be printed by [grid.raster](#).

Author(s)

Bernd Fischer

See Also

[grid.raster](#), [colorRampPalette](#), [DmeISGI-package](#)

Examples

```
print(toRaster)
```

trsf*Transform a correlation to a distance*

Description

Transforms a correlation to a distance.

Usage

```
trsf(x)
```

Arguments

x A real valued vector with values in $[-1,1]$. It is intended to be a correlation.

Value

a real valued vector of distances.

See Also

[DmelSGI-package](#)

Examples

```
print(trsf)
```

Index

* datasets

- applyDimensionReduction, 4
- callInteractions, 5
- datamatrix, 6
- DPiM, 7
- estimatePairwiseInteractions, 8
- FBgn2anno, 9
- Features, 10
- fitepistasis, 11
- getBaseDir, 12
- grid.spider, 12
- hrNames, 13
- Interactions, 14
- Intogen, 15
- learnCoComplexFct, 16
- mainEffects, 17
- myHeatmap, 19
- mymedpolish, 20
- orderDim, 21
- orderSpiderAxis, 22
- pimatrix, 23
- plot2Phenotypes, 24
- plotHairballLabels, 25
- qualityControlFeature, 26
- qualityControlGene, 26
- RohnEtAl, 27
- selectByStability, 29
- SelectedClusters, 30
- SelectedClustersComplexes, 30
- SKDdata, 31
- stabilitySelection, 32
- subSampleForStabilitySelection, 33
- subSampleForStabilitySelectionFct, 34
- TID2HUGO, 35
- toMatrix, 35
- toRaster, 36
- trsf, 37

* package

DmelSGI-package, 3

applyDimensionReduction, 3, 4

callInteractions, 3, 5
colorRampPalette, 19, 36
convertCorrelations
(learnCoComplexFct), 16

datamatrix, 3, 6
DmelSGI (DmelSGI-package), 3
DmelSGI-package, 3, 4, 5, 7–17, 19, 21, 22,
24–37
DPiM, 7

estimatePairwiseInteractions, 3, 8, 20,
21

FBgn2anno, 9
Features, 3, 10, 26
fitepistasis, 11

getBaseDir, 12
grid.raster, 36
grid.spider, 12, 22

hrNames, 13

Interactions, 3, 14, 23, 24
Intogen, 15

learnCoComplexFct, 16

mainEffects, 3, 17
medpolish, 20, 21
myHeatmap, 19
mymedpolish, 3, 8, 20

orderDim, 21
orderSpiderAxis, 13, 22

pimatrix, 3, 14, 15, 23

plot2Phenotypes, [24](#)
plotHairballLabels, [25](#)
plotPIData (plot2Phenotypes), [24](#)

qualityControlFeature, [3](#), [26](#)
qualityControlGene, [3](#), [26](#)

RohnEtAl, [27](#)

selectByStability, [4](#), [29](#), [33](#), [34](#)
SelectedClusters, [30](#)
SelectedClustersComplexes, [30](#)
SKDdata, [3](#), [31](#)
stabilitySelection, [3](#), [4](#), [32](#)
subSampleForStabilitySelection, [3](#), [33](#),
[34](#)
subSampleForStabilitySelectionFct, [3](#),
[29](#), [33](#), [34](#)

TID2HUGO, [35](#)
toMatrix, [35](#)
toRaster, [36](#)
trsf, [37](#)