

# Package ‘metaLINCS’

July 11, 2022

**Title** Meta-level analysis between experimental gene expression profiles and LINCS L1000 connectivity map signatures

**Version** 0.9.0

**Description** 'metaLINCS' calculates and visualises correlation between experimental gene expression profiles with perturbation signatures from the LINCS L1000 project that has collected gene expression profiles for thousands of perturbagens at a variety of time points, doses, and cell lines.

**License** GPL (>= 3)

**URL** <https://github.com/bigomics/metaLINCS>

**BugReports** <https://github.com/bigomics/metaLINCS/issues>

**Depends** R (>= 3.5.0)

**Imports** corrplot,  
fgsea,  
gplots,  
graphics,  
Matrix,  
qlcMatrix,  
stats,  
utils,  
uwot

**Suggests** knitr,  
rmarkdown,  
testthat (>= 3.0.0)

**VignetteBuilder** knitr

**Config/testthat/edition** 3

**Encoding** UTF-8

**LazyData** true

**LazyDataCompression** bzip2

**Roxygen** list(markdown = TRUE)

**RoxygenNote** 7.1.2

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computeConnectivityEnrichment
<i>Compute Connectivity Enrichment</i>

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Description

Compute Connectivity Enrichment

Usage

```
computeConnectivityEnrichment(  
  mFC,  
  names = NULL,  
  mDrugEnrich = SpaceLINCS::mDrugEnrich,  
  nmin = 15,  
  nprune = 250  
)
```

Arguments

mFC	A matrix of differential gene expression fold-change of own experiment, the rows name of the matrix must be the genes names
names	Names in case mFC is a vector
mDrugEnrich	a large matrix represents the gene expression fold change in the presence of different perturbations, either various drugs concentrations or other genetic engineering techniques as gene-knock-in etc..
nmin	the minimum number of experiments in the drug set
nprune	takes only the (nprune) top matching drugs for each comparison to reduce the size of the matrix. default is 0 to take the full matrix

Value

list of drugs enrichment statistics and annotations

### Examples

```
# from the data-sets provided as examples within the package load the .rda files
# DrugsAnnot, mDrugEnrich, mFC
res <- computeConnectivityEnrichment(mFC, mDrugEnrich, nmin=15, nprune = 250)
```

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computeMoaEnrichment	<i>Get the mechanism of action object</i>
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### Description

Get the mechanism of action object

### Usage

```
computeMoaEnrichment(res, annot = SpaceLINCS::DrugsAnnot)
```

### Arguments

res	is the output object from computeConnectivityEnrichment()
annot	is a drug annotation table with moa and target columns

### Value

MoA Mechanism of Action object

### Examples

```
# from the data-sets provided as examples within the package load the .rda files
# DrugsAnnot, mDrugEnrich, mFC
res <- computeConnectivityEnrichment(mFC)
moa <- computeMoaEnrichment(res)
```

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DrugsAnnot	<i>Annotation of a set of drugs</i>
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### Description

This data set is a matrix represents the drugs annotation that contains drugs' names, targets, mechanism of action , clinical phase, disease area, and indication.

### Usage

```
data("DrugsAnnot")
```

**Format**

A data frame with 6125 observations on the following 6 variables.

pert\_iname a character vector  
 clinical\_phase a character vector  
 moa a character vector  
 target a character vector  
 disease\_area a character vector  
 indication a character vector

**Details**

L1000 repurposing drugs . For research use only. Do not use the Repurposing Hub to make clinical treatment decisions. BROAD DOES NOT GUARANTEE OR WARRANT THE ACCURACY OF THE DATA WITHIN THE REPURPOSING HUB.

**Source**

"The Drug Repurposing Hub, Broad Institute" <http://www.broadinstitute.org/repurposing>

**References**

Corsello SM, et al. Nature Medicine. 2017 Apr 7;23(4):405-408. doi: 10.1038/nm.4306

**Examples**

```
head(SpaceLINCS::DrugsAnnot)
## str(DrugsAnnot)
```

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mDrugEnrich	<i>The drugs activity on 1001 genes</i>
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**Description**

Large matrix representing the gene expression fold change of 1001 genes in the presence of 20220 different drugs concentrations and working time.

**Usage**

```
SpaceLINCS::mDrugEnrich
```

**Format**

A gene expression matrix (log2FC) with 1001 rows and 20220 columns corresponding to 1001 genes and 20220 drug perturbation experiments.

## Details

The LINCS L1000 project has collected gene expression profiles for thousands of perturbagens at a variety of time points, doses, and cell lines. A full list of the chemical and genetic perturbations used can be found on the CLUE website along with their descriptions.

## Source

<https://lincsproject.org/LINCS/tools/workflows/find-the-best-place-to-obtain-the-lincs-l1000-data>

## References

Duan Q et al. LINCS Canvas Browser: interactive web app to query, browse and interrogate LINCS L1000 gene expression signatures. Nucleic Acids Res. 2014 Jul;42 (Web Server issue):W449-60.

## Examples

```
head(SpaceLINCS::mDrugEnrich)
## maybe str(mDrugEnrich) ; plot(mDrugEnrich) ...
```

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mFC	<i>Fold change matrix of differential gene expression from RNA-seq data of multiple myeloma</i>
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## Description

Example signature data as query input for the analysis. This is the matrix that you probably want to replace. There are 6 contrasts: glucocorticoid sensitive vs glucocorticoid resistant ("Resistant.vs.Sensitive"); Treated with Withaferin-A vs untreated ("WithaferinA.vs.Untreated"); untreated glucocorticoid resistant cells vs untreated glucocorticoid sensitive cells ("UTR.vs.UTS"); Treated with Withaferin-A glucocorticoid resistant cells vs Treated with Withaferin-A glucocorticoid sensitive cells ("WAR.vs.WAS"); Treated Resistant vs Untreated Resistant ("WAR.vs.UTR"); Treated sensitive vs Untreated sensitive ("WAS.vs.UTS")

## Usage

```
dim(mFC)
head(mFC)
```

## Format

A data frame with 5332 observations on the following 6 variables.

```
Resistant.vs.Sensitive a numeric vector
WithaferinA.vs.Untreated a numeric vector
UTR.vs.UTS a numeric vector
WAR.vs.WAS a numeric vector
WAR.vs.UTR a numeric vector
WAS.vs.UTS a numeric vector
```

## Details

The RNAseq data from (Logie et al 2021) study. They compared the therapeutic efficacy of the phytochemical kinase inhibitor withaferin A with the clinically approved BTK inhibitor ibrutinib to target hyperactivated tyrosine kinase signaling in glucocorticoid-resistant multiple myeloma cells. The results demonstrate that withaferin-A induced cell death of glucocorticoid-resistant MM1R cells involves covalent cysteine targeting of multiple Hinge-6 domain type tyrosine kinases of the kinase cysteinome classification, including BTK.

## Source

GEO accession number GSE162475

## References

Logie E, Chirumamilla CS, Perez-Novo C, Shaw P et al. Covalent Cysteine Targeting of Bruton's Tyrosine Kinase (BTK) Family by Withaferin-A Reduces Survival of Glucocorticoid-Resistant Multiple Myeloma MM1 Cell. Cancers (Basel) 2021 Mar 31;13(7). PMID: 33807411

## Examples

```
data(mFC)
## maybe str(mFC) ; plot(mFC) ...
```

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plotActivationMap	<i>Plot drug Activity Map</i>
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## Description

Plot drug Activity Map

## Usage

```
plotActivationMap(res, nterms = 60, nfc = 20, rot = FALSE)
```

## Arguments

nterms	integer
nfc	integer
dsea	drugs set enrichment analysis object as output of getActiveDSEA
drugs	a list of drugs enrichment stats and annotations the output of computePeturbEnrichment()
method	the computation methods of Enrichment, either using GSEA algorithm or the rank correlation.
contr	is a character string represent the two compared conditions(contrast) as it is provided from the fold-change matrix

**Value**

plot of drug activity map

**Examples**

```
# from the data-sets provided as examples within the package load the .rda files
#DrugsAnnot, mDrugEnrich, mFC
res <- computeConnectivityEnrichment(mFC)
plotActivationMap(res, nterms = 50, nfc=20, rot=TRUE)
```

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plotDrugConnectivity    *Plot Drug Connectivity*

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**Description**

Plot Drug Connectivity

**Usage**

```
plotDrugConnectivity(res, contr, drugs = NULL, nplots = 25)
```

**Arguments**

res	is a result object from the output of computeConnectivityEnrichment()
contr	is a index of the selected contrast
drugs	is a vector of selected drugs to show
nplots	is the number of plots to show

**Value**

a plot of drug connectivity

**Examples**

```
# from the data-sets provided as examples within the package load the .rda files
# DrugsAnnot, mDrugEnrich, mFC
res <- computeConnectivityEnrichment(mFC)
plotDrugConnectivity(res, contr=1, nplots=16)
```

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plotMOA	<i>Plot mechanism of action</i>
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**Description**

Plot mechanism of action

**Usage**

```
plotMOA(moa, contr = NULL, type = c("drugClass", "targetGene"), ntop = 20)
```

**Arguments**

ntop	the number of the top enteries (genes or drug), ntop = 16 as a default value
dsea	is dsea object which is the output of the function getActiveDSEA()

**Value**

plot of mechanism of action

**Examples**

```
# from the data-sets provided as examples within the package load the .rda files
# DrugsAnnot, mDrugEnrich, mFC
res <- computeConnectivityEnrichment(mFC)
moa <- computeMoaEnrichment(res)
plotMOA(moa, contr=1, type="drugClass", ntop=25)
```

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selectResult	<i>Select result</i>
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**Description**

Select result

**Usage**

```
selectResult(res, contr)
```

**Arguments**

res	is the result from computeConnectivityEnrichment()
contr	contrast selection

**Value**

selected result for selected contrast



**Examples**

```
res1 <- selectResult(res,1)
```

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SpaceLINCS

*SpaceLINCS: is a new method to visualise correlations between experimental gene expression profiles and drug connectivity map profiles*

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**Description**

The SpaceLINCS package provides three categories of important functions: computePeturbEnrichment, computeComboEnrichment and several plotting functions

**SpaceLINCS functions**

computeConnectivityEnrichment: Compute Connectivity Enrichment based on database and annotation provided as input parameter  
USAGE: computeConnectivityEnrichment(mFC = mFC, mDrugEnrich = mDrugEnrich, nprune = 250, contrast = NULL)

plotActivationMap: Plot drug Activity Map. USAGE: plotActivationMap(res, nterms = 50, nfc = 20)

computeMoaEnrichment: Compute mechanism-of-action enrichment  
USAGE: computeMoaEnrichment(dsea)

plotDrugConnectivity: Plot Drug Connectivity  
USAGE: plotDrugConnectivity(dsea)

plotMOA: Plot mechanism of action bargraph  
USAGE: plotMOA(dsea, ntop = 16)

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