

Package ‘attract’

June 23, 2025

Type Package

Title Methods to Find the Gene Expression Modules that Represent the Drivers of Kauffman's Attractor Landscape

Version 1.61.0

Date 2018-06-29

Author Jessica Mar

Maintainer Samuel Zimmerman <samuel.e.zimmerman@gmail.com>

Description This package contains the functions to find the gene expression modules that represent the drivers of Kauffman's attractor landscape. The modules are the core attractor pathways that discriminate between different cell types of groups of interest. Each pathway has a set of synexpression groups, which show transcriptionally-coordinated changes in gene expression.

License LGPL (>= 2.0)

Collate classdef.R findAttractorsStep.R removeFlatGenesStep.R
findSynexprsStep.R findFuncEnrichAndCorr.R

Depends R (>= 3.4.0), AnnotationDbi

Imports Biobase, limma, cluster, GOSTats, graphics, stats,
reactome.db, KEGGREST, org.Hs.eg.db, utils, methods

Suggests illuminaHumanv1.db

biocViews ImmunoOncology, KEGG, Reactome, GeneExpression, Pathways,
GeneSetEnrichment, Microarray, RNASeq

NeedsCompilation no

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

git_branch devel

git_last_commit 19c3524

git_last_commit_date 2025-04-15

Repository Bioconductor 3.22

Date/Publication 2025-06-23

Contents

attract-package	2
AttractorModuleSet-class	3
buildCorMatrix	4
buildCustomIncidenceMatrix	5
buildKeggIncidenceMatrix	6
calcFuncSynexprs	7
calcInform	8
calcModfstat	9
calcRss	10
exprs.dat	11
filterDataSet	12
findAttractors	12
findCorrPartners	14
findOnepwaySynexprs	15
findSynexprs	16
flagPwayExists	17
getCustomGenes	18
getPwayGenes	19
loring.eset	20
plotsynexprs	20
removeFlatGenes	21
samp.info	22
subset.loring.eset	23
SynExpressionSet-class	24
Index	25

attract-package	<i>Methods to find the Gene Expression Modules that Represent the Drivers of Kauffman’s Attractor Landscape</i>
-----------------	---

Description

This package contains functions used to determine the gene expression modules that represent the drivers of Kauffman’s attractor landscape.

Details

Package: attract
Type: Package
Version: 1.33.2
Date: 2018-06-29
License:
LazyLoad: yes

The method can be summarized in the following key steps: (1) Determine core KEGG or reactome pathways that discriminate the most strongly between celltypes or experimental groups of interest (see findAttractors)). (2) Find the different synexpression groups that are present within a core

attractor pathway (see `findSynexprs`). (3) Find sets of genes that show highly similar profiles to the synexpression groups within an attractor pathway module (see `findCorrPartners`). (4) Test for functional enrichment for each of the synexpression groups to detect any potentially shared biological themes (see `calcFuncSynexprs`).

Author(s)

Jessica Mar <jess@jimmy.harvard.edu>

References

Kauffman S. 2004. A proposal for using the ensemble approach to understand genetic regulatory networks. *J Theor Biol.* 230:581. Mar JC, Wells CA, Quackenbush J. 2010. Identifying Gene Expression Modules that Represent the Drivers of Kauffman's Attractor Landscape. To Appear. Müller F et al. 2008. Regulatory networks define phenotypic classes of human stem cell lines. *Nature.* 455(7211): 401. Mar JC, Wells CA, Quackenbush J. 2010. Defining an Informativeness Metric for Clustering Gene Expression Data. To Appear.

Examples

```
## Not run:
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db",
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, remove.these.genes)
mapk.cor <- findCorrPartners(mapk.syn, subset.loring.eset, remove.these.genes)
mapk.func <- calcFuncSynexprs(mapk.syn, attractor.states, "CC", annotation="illuminaHumanv1.db")

## End(Not run)
```

AttractorModuleSet-class

Class AttractorModuleSet

Description

This is a class representation for storing the output of the `findAttractors` function.

Objects from the Class

Objects are output by the function `findAttractors`. Objects can also be created by using `new("AttractorModuleSet", ...)`.

Slots

eSet: ExpressionSet which primarily stores the expression data and the phenotype/sample data sets.

cellTypeTag: character string of the tag which stores the group membership information for the samples. Must be a column name of the data frame `pData(eset)`.

incidenceMatrix: incidence matrix used as input to GSEAlm.

rankedPathways: Data frame of significantly enriched pathways, ranked first by significance and then by size.

Methods

No methods have yet been defined with class "AttractorModuleSet" in the signature.

Note

This class is better describe in the vignette.

Author(s)

Jessica Mar <jess@jimmy.harvard.edu>

Examples

```
## Not run:
new.attractmodule <- new("AttractorModuleSet", eSet=new("ExpressionSet"), cellTypeTag=character(1), incidence
## End(Not run)
```

buildCorMatrix	<i>Internal function - builds the correlation matrix between an average transcriptional module expression profile and a set of other genes.</i>
----------------	---

Description

Internal function - builds the correlation matrix between an average transcriptional module expression profile and a set of other genes.

Usage

```
buildCorMatrix(dat.fr, module.genes, cor.cutoff)
```

Arguments

dat.fr	a matrix object of gene expression values.
module.genes	character vector specifying genes that belong in this pathway module.
cor.cutoff	numeric value specifying the correlation cut-off.

Details

This function is called internally by findCorrPartners which is easier for the user to call since findCorrPartners uses the SynExpressionSet and ExpressionSet class objects directly.

Value

A character vector of genes that meet the correlation cut-off.

Author(s)

Jessica Mar

Examples

```
## Not run:
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, remove.these.genes)
cormat <- buildCorMatrix(exprs(subset.loring.eset), mapk.syn, 0.95)

## End(Not run)
```

```
buildCustomIncidenceMatrix
```

This function builds an incidence matrix for custom gene sets.

Description

This function builds an incidence matrix for custom gene sets.

Usage

```
buildCustomIncidenceMatrix(geneSetFrame, geneNames, databaseGeneFormat, expressionSetGeneFormat, g
```

Arguments

geneSetFrame	a dataframe where rows are gene sets and columns are genes.
geneNames	a vector of all the genes in the geneSetFrame dataframe
databaseGeneFormat	a character string specifying the type of identifier for a gene in a database (KEGG, reactome, MsigDB) gene set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)
expressionSetGeneFormat	a character string specifying the type of identifier for a gene in your expression data set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)
geneSetNames	a vector of the name of the custom gene sets.

Details

This function creates an incidence matrix from a dataframe where the rows are the names of gene sets and the columns are genes.

Value

A matrix object with 0 and 1 entries where 1 denotes membership of a gene in a custom gene set, 0 denotes non-membership.

Author(s)

Jessica Mar

References

Mar, J., C. Wells, and J. Quackenbush, Identifying the Gene Expression Modules that Represent the Drivers of Kauffman's Attractor Landscape. to appear, 2010.

buildKeggIncidenceMatrix

Internal function - buildings the KEGG incidence matrix required by GSEAlm.

Description

Internal function - buildings the KEGG incidence matrix required by GSEAlm.

Usage

```
buildKeggIncidenceMatrix(kegg.ids, gene.ids, annotation, database, analysis, envPos, expressionSet)
```

Arguments

kegg.ids	character vector of KEGG pathway ids.
gene.ids	character vector of gene ids.
annotation	character string that denotes which annotation package to be used, eg. illuminaHumanv1.db.
database	a character string specifying what pathway database you would like to use.
analysis	a character string specifying what type of experiment you performed, microarray or RNAseq.
envPos	the position of the annotation package in the R search path.
expressionSetGeneFormat	a character string specifying the type of identifier for a gene in your expression data set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)

Details

This function is called internally by findAttractors.

Value

A matrix object with 0 and 1 entries where 1 denotes membership of a gene in a KEGG or reactome pathway, 0 denotes non-membership.

Author(s)

Jessica Mar

References

Kanehisa, M. and S. Goto, KEGG: Kyoto Encyclopedia of Genes and Genomes. . Nucleic Acids Res., 2000. 28: p. 27-30.

Examples

```
## Not run:
# this takes a long time!
require("illuminaHumanv2.db", character.only=TRUE)
loadNamespace("illuminaHumanv2.db")
envPos <- match(paste("package:", "illuminaHumanv2.db", sep=""), search())
kegg.ids <- ls(illuminaHumanv2PATH2PROBE)
gene.ids <- ls(illuminaHumanv2PATH)
database <- "KEGG"
analysis <- "microarray"
imat <- buildKeggIncidence(kegg.ids, gene.ids, illuminaHumanv2.db,database, analysis, envPos)

## End(Not run)
```

calcFuncSynexprs

Functional enrichmental analysis for a set of synexpression groups.

Description

This function performs functional enrichment for a given set of synexpression groups.

Usage

```
calcFuncSynexprs(mySynExpressionSet, myAttractorModuleSet, ontology = "BP", min.pvalue = 0.05, min.pwaysize = 10, annotation = "illuminaHumanv2.db", analysis = "microarray", expressionSetGeneFormat = "SYMBOL", ...)
```

Arguments

mySynExpressionSet	SynExpressionSet object.
myAttractorModuleSet	AttractorModuleSet object.
ontology	character string specifying which GO ontology to use, either "MF", "BP", or "CC"; defaults to "BP".
min.pvalue	numeric value specifying adjusted P-value cut-off to use, categories with P-values <= min.pvalue will be reported.
min.pwaysize	integer specifying minimum size of the pathway or category to consider for enrichment analysis.
annotation	character string specifying the annotation package that corresponds to the chip platform the data was generated from.
analysis	a character string specifying what type of experiment you performed, microarray or RNAseq.
expressionSetGeneFormat	a character string specifying the type of identifier for a gene in your expression data set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)
...	additional arguments.

Details

This function performs a functional enrichment analysis on each synexpression group using the hyperGTest from the GOSTats package. P-values are adjusted using the Benjamini-Hochberg correction method. Results are returned only if they satisfy the minimum P-value level, as specified by the min.pvalue argument.

Value

A list object.

Author(s)

Jessica Mar

References

Falcon, S. and R. Gentleman, Using GOSTats to test gene lists for GO term association. Bioinformatics, 2007. 23(2): p. 257-8.

Examples

```
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, remove.these.genes)
mapk.func <- calcFuncSynexprs(mapk.syn, attractor.states, "CC", annotation="illuminaHumanv1.db", analysis="m")
```

calcInform

Function calculates the informativeness metric (average MSS) for a set of cluster assignments.

Description

Function calculates the informativeness metric (average MSS) for a set of cluster assignments.

Usage

```
calcInform(exprs.dat, cl, class.vector)
```

Arguments

exprs.dat	a matrix of gene expression values.
cl	a vector of cluster assignments.
class.vector	a vector specifying the group membership of the samples.

Details

This function is also called internally by findSynexprs.

Value

A numeric value representing the average MSS value (informativeness metric) for a set of cluster assignments. For an informative cluster, the RSS values should be very small relative to those produced by the informativeness metric (the MSS values).

Author(s)

Jessica Mar

References

Mar, J., C. Wells, and J. Quackenbush, Defining an Informativeness Metric for Clustering Gene Expression Data. to appear, 2010.

Examples

```
## Not run:
library(cluster)
data(subset.loring.eset)
clustObj <- agnes(as.dist(1-t(cor(exprs(subset.loring.eset)))))
cinform.vals <- NULL
for( i in 1:10 ){
  cinform.vals <- c(cinform.vals, calcInform(exprs(subset.loring.eset), cutree(clustObj,i), pData(subset.loring.eset)))
}
k <- (1:10)[cinform.vals==max(cinform.vals)] # gives the optimal number of clusters

## End(Not run)
```

calcModfstat	<i>Function calculates a modified F-statistic for a set of cluster assignments.</i>
--------------	---

Description

Function calculates a modified F-statistic for a set of cluster assignments.

Usage

```
calcModfstat(exprs.dat, cl, class.vector)
```

Arguments

exprs.dat	a matrix of gene expression values.
cl	a vector of cluster assignments.
class.vector	a vector specifying group membership of the samples.

Details

This function is called internally by findSynexprs.

Value

a modified F-statistic (average MSS/average RSS) value for a set of cluster assignments.

Author(s)

Jessica Mar

Examples

```
## Not run:
library(cluster)
data(subset.loring.eset)
clustObj <- agnes(as.dist(1-t(cor(exprs(subset.loring.eset)))))
cfmod.vals <- NULL
for( i in 1:10 ){
  cfmod.vals <- c(cfmod.vals, calcModfstat(exprs(subset.loring.eset), cutree(clustObj,i), pData(subset.loring.eset)))
}
k <- (1:10)[cfmod.vals==max(cfmod.vals)]

## End(Not run)
```

calcRss

Function calculates the average RSS for a set of cluster assignments.

Description

Function calculates the average RSS for a set of cluster assignments.

Usage

```
calcRss(exprs.dat, cl, class.vector)
```

Arguments

exprs.dat a matrix of gene expression values.
cl a vector of cluster assignments.
class.vector a vector specifying the group membership of the samples.

Details

This function is called internally by findSynexprs. For an informative cluster, the RSS values should be very small relative to those produced by the informativeness metric (the MSS values).

Value

A numeric value representing the average RSS value for this set of cluster assignments.

Author(s)

Jessica Mar

Examples

```
## Not run:
library(cluster)
data(subset.loring.eset)
clustObj <- agnes(as.dist(1-t(cor(exprs(subset.loring.eset)))))
crss.vals <- NULL
for( i in 1:10 ){
  crss.vals <- c(crss.vals, calcRss(exprs(subset.loring.eset), cutree(clustObj,i), pData(subset.loring.eset)$
}
# The RSS values are expected to be smaller than the informativeness metric values in the presence of genuine clu

## End(Not run)
```

exprs.dat

*Gene Expression Matrix of Published Data***Description**

This is a matrix object containing published gene expression data from Mueller et al. (NCBI GEO accession id GSE11508). The data set contains 11044 probes for 68 samples. From the original data set, we have selected four cell lines giving a total of 68 samples - embryonic stem cells (12 samples), neural progenitors (31 samples), neural stem cells (8 samples) and teratoma-differentiated cells (17 samples). The lines have also been restricted based on Illumina BeadChip platform, and only those collected using the WG-6 version have been used.

We also applied a quality filter to the original gene expression data where a probe was retained if it passed a 0.99 detection score in 75

Usage

```
data(exprs.dat)
```

Format

A matrix with normalized log2 expression intensities for 11044 probes on 68 samples (representing 4 different cell types).

Value

A matrix object containing published gene expression data from Mueller et al. (NCBI GEO accession id GSE11508). The data set contains 11044 probes for 68 samples.

References

Mueller F, et al., Regulatory networks define phenotypic classes of human stem cell lines. Nature, 2008. 455(7211): p. 401-405.

See Also

[samp.info](#), [loring.eset](#)

Examples

```
data(exprs.dat)
```

filterDataSet

This function filters our lowly expressed genes in RNAseq data.

Description

This function filters our lowly expressed genes in RNAseq data.

Usage

```
filterDataSet(data, filterPerc=0.75)
```

Arguments

data	A dataset with genes as rows and samples as columns.
filterPerc	a number specifying the percent of expression values that are not equal to 0 for a gene.

Details

This function removes any genes in a dataset that have an expression value of 0 for a specified percentage of samples.

Value

A data frame is returned.

Author(s)

Jessica Mar

Examples

```
data(exprs.dat)
exprs.filtered.dat <- filterDataSet(exprs.dat)
```

findAttractors

Infers the set of cell-lineage specific gene expression modules using GSEAlm and KEGG.

Description

The function infers a set of KEGG pathways that correspond to the cell-lineage specific gene expression modules, as determined using GSEA. These pathways represent those that show the greatest discrimination between the different cell types or tissues in the expression data set supplied.

Usage

```
findAttractors(myEset, cellTypeTag, min.pwaysize = 5, annotation = "illuminaHumanv2.db", database=
```

Arguments

<code>myEset</code>	ExpressionSet object.
<code>cellTypeTag</code>	character string of the variable name which stores the cell-lineages or experimental groups of interest for the samples in the data set (this string should be one of the column names of <code>pData(myEset)</code>).
<code>min.pwaysize</code>	integer specifying the minimum size of the KEGG or reactome pathways to consider in the analysis.
<code>annotation</code>	character string specifying the annotation package that corresponds to the chip platform or organism (for RNAseq data) the data was generated from.
<code>database</code>	a character string specifying what pathway database you would like to use.
<code>analysis</code>	a character string specifying what type of experiment you performed, microarray or RNAseq.
<code>databaseGeneFormat</code>	a character string specifying the type of identifier for a gene in a database (KEGG, REACTOME, MsigDB) gene set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)
<code>expressionSetGeneFormat</code>	a character string specifying the type of identifier for a gene in your expression data set. The default value is NULL. (ex. SYMBOL, ENTREZID, REFSEQ, ENSEMBL)
<code>...</code>	additional arguments.

Details

This function subsets the expression data so that only those genes with annotations in KEGG or reactome are used for the downstream gene set enrichment analysis. This subset is stored in the `eSet` slot of the `AttractoModuleSet` output object.

The `GSEAlm` algorithm finds the KEGG or reactome pathway modules which discriminate between the celltypes or experimental groups of interest. It also ranks the results of the `GSEAlm` step by significance of these pathway modules, as stored in `rankedPathways`.

The output object of the `findAttractors` function also contains the incidence matrix that was built for the KEGG or reactome pathways, stored in the slot `incidenceMatrix` and the character string denoting which column of the sample data represents the cell type or experimental groups of interest, as stored in the slot `cellTypeTag`.

Value

An `AttractorModuleSet` object.

Author(s)

Jessica Mar

References

Jiang, Z. and R. Gentleman, Extensions to gene set enrichment. *Bioinformatics*, 2007. 23(3): p. 306-313. Kanehisa, M. and S. Goto, KEGG: Kyoto Encyclopedia of Genes and Genomes. . *Nucleic Acids Res.*, 2000. 28: p. 27-30. Mar, J., C. Wells, and J. Quackenbush, Identifying the

Gene Expression Modules that Represent the Drivers of Kauffman's Attractor Landscape. to appear, 2010.

Examples

```
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", annotation="illuminaHumanv1.db", database=
MSigDBpath <- system.file("extdata", "c4.cgn.v5.0.entrez.gmt", package="attract")
attractor.states.cutsom <- findAttractors(subset.loring.eset, "celltype", annotation="illuminaHumanv1.db", da
```

findCorrPartners	<i>Determines Genes with Highly Correlated Expression Profiles to a Synexpression Group</i>
------------------	---

Description

This function finds genes with expression profiles highly correlated to a synexpression group.

Usage

```
findCorrPartners(mySynExpressionSet, myEset, removeGenes = NULL, cor.cutoff = 0.85, ...)
```

Arguments

mySynExpressionSet	SynExpressionSet object.
myEset	ExpressionSet object.
removeGenes	vector of probes that specify those genes who demonstrate little variability across the different celltypes and thus should be removed from downstream analysis.
cor.cutoff	numeric value specifying the correlation cut-off.
...	additional arguments.

Details

Genes with highly correlated profiles to the synexpression groups (e.g. $R > 0.85$) are also likely to be integral in maintaining cell type-specific differences, however due to their lack of inclusion in resources like KEGG, would not have been picked up by the first GSEA step using findAttractors.

Value

A SynExpressionSet object which stores the genes that are highly correlated with the synexpression group provided, and their average expression profile.

Author(s)

Jessica Mar

Examples

```
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, remove.these.genes)
mapk.cor <- findCorrPartners(mapk.syn, subset.loring.eset, remove.these.genes)
```

findOnewaySynexprs	<i>Internal function - finds the synexpression groups for a single given pathway.</i>
--------------------	---

Description

Internal function - finds the synexpression groups for a single given pathway.

Usage

```
findOnewaySynexprs(myIDs, myDataSet, cellTypeTag, min.clustersize = 5, removeGenes = NULL, ...)
```

Arguments

myIDs	a single character string denoting the KEGG or reactome ID of the pathway module to be analyzed or a character codevector of gene names of a pathway.
myDataSet	AttractorModuleSet object, output of the findAttractors step. This could also be an ExpressionSet object.
cellTypeTag	character string of the variable name which stores the cell-lineages or experimental groups of interest for the samples in the data set (this string should be one of the column names of pData(myEset)).
min.clustersize	integer specifying the minimum number of genes that must be present in clusters that are inferred.
removeGenes	vector of probes that specify those genes who demonstrate little variability across the different celltypes and thus should be removed from downstream analysis.
...	additional arguments.

Details

This function is called internally by calcFuncSynexprs. Users should use calcFuncSynexprs rather than calling findOnewaySynexprs directly.

Value

A SynExpressionSet object is returned.

Author(s)

Jessica Mar

Examples

```
## Not run:
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
map.syn <- findOnewaySynexprs("04010", attractor.states, "celltype", removeGenes=remove.these.genes)
vec.geneid <- c("GI_17999531-S", "GI_17978503-A")
custom.syn <- findSynexprs(vec.geneid, subset.loring.eset, "celltype", removeGenes=remove.these.genes)

## End(Not run)
```

findSynexprs	<i>This function finds the synexpression groups present within a core attractor pathway module.</i>
--------------	---

Description

This function takes the modules that were inferred from the GSEA step using (findAttractors) and finds a set of transcriptionally coherent set of genes associated with a particular core attractor pathway, i.e. the synexpression groups.

Usage

```
findSynexprs(myIDs, myDataSet, cellTypeTag, removeGenes = NULL, min.clustersize = 5, ...)
```

Arguments

myIDs	either a single character string or vector of character strings denoting the KEGG or reactome IDs of the pathway modules to be analyzed. It may also be a character codevector of gene names of a pathway if defining a custom pathway.
myDataSet	AttractorModuleSet object, output of the findAttractors step. This could also be an ExpressionSet object if using a custom pathway.
cellTypeTag	character string of the variable name which stores the cell-lineages or experimental groups of interest for the samples in the data set (this string should be one of the column names of pData(myEset)).
removeGenes	vector of gene names that specify those genes who demonstrate little variability across the different celltypes and thus should be removed from downstream analysis.
min.clustersize	integer specifying the minimum number of genes that must be present in clusters that are inferred.
...	additional arguments.

Details

This function performs a hierarchical cluster analysis of the genes in a core attractor pathway module, and uses an informativeness metric to determine the number of optimal clusters (synexpression groups) that describe the data.

Value

If a single KEGG or reactome ID is specified in pwayIds, then a SynExpressionSet object is returned. If a multiple KEGG or reactome IDs are specified, then an environment object is returned where the keys are labeled "pwayIDsynexprs" (e.g. for MAPK KEGGID = 04010, the key is pway04010synexprs). The value associated with each key is a [SynExpressionSet](#) object.

Author(s)

Jessica Mar

References

Mar, J., C. Wells, and J. Quackenbush, Identifying the Gene Expression Modules that Represent the Drivers of Kauffman's Attractor Landscape. to appear, 2010.

Examples

```
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, "celltype", remove.these.genes)
top5.syn <- findSynexprs(attractor.states@rankedPathways[1:5,1], attractor.states, "celltype", removeGenes=remove.these.genes)
vec.geneid <- c("GI_17999531-S", "GI_17978503-A")
custom.syn <- findSynexprs(vec.geneid, subset.loring.eset, "celltype", removeGenes=remove.these.genes)
```

flagPwayExists

Internal function - flags a gene if it exists in a pathway.

Description

Internal function - flags a gene if it exists in a pathway.

Usage

```
flagPwayExists(x)
```

Arguments

x A vector of pathway ids.

Details

This function is called internally by the findAttractors function. Function returns TRUE if the probe exists in at least one pways, FALSE if there are no pway annotations.

Value

A logical value.

Author(s)

Jessica Mar

Examples

```
## Not run:
library(illuminaHumanv2.db)
flag.check <- flagPwayExists(ls(illuminaHumanv2PATH2PROBE))

## End(Not run)
```

getCustomGenes	<i>Function removes genes from the custom pathway that demonstrate little variation across the cell types.</i>
----------------	--

Description

Function removes genes from the custom pathway that demonstrate little variation across the cell types.

Usage

```
getCustomGenes(vec.geneid, removeGenes = NULL)
```

Arguments

vec.geneid	a vector of character strings denoting a custom gene set.
removeGenes	vector of gene names that specify those genes who demonstrate little variability across the different cell types and thus should be removed from downstream analysis.

Details

This function is also called internally by findSynexprs.

Value

A vector of gene names that have variable expression across the different cell types.

Author(s)

Jessica Mar

References

Mar, J., C. Wells, and J. Quackenbush, Defining an Informativeness Metric for Clustering Gene Expression Data. to appear, 2010.

Examples

```
## Not run:

data(subset.loring.eset)
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
vec.geneid <- c("GI_17999531-S", "GI_17978503-A")
customGenes <- getCustomGenes(vec.geneid, removeGenes=NULL)

## End(Not run)
```

getPwayGenes	<i>Function removes genes from the chosen pathway that demonstrate little variation across the cell types.</i>
--------------	--

Description

Function removes genes from the chosen pathway that demonstrate little variation across the cell types.

Usage

```
getPwayGenes(pathwayIds, myAttractorModuleSet, removeGenes = NULL)
```

Arguments

pathwayIds	a single character string denoting the KEGG or reactome ID of the pathway module to be analyzed.
myAttractorModuleSet	AttractorModuleSet object, output of the findAttractors step.
removeGenes	vector of probes or gene IDs (RNAseq) that specify those genes who demonstrate little variability across the different celltypes and thus should be removed from downstream analysis.

Details

This function is also called internally by findSynexprs.

Value

A vector of gene names that have variable expression across the different cell types in a pathway.

Author(s)

Jessica Mar

References

Mar, J., C. Wells, and J. Quackenbush, Defining an Informativeness Metric for Clustering Gene Expression Data. to appear, 2010.

Examples

```
## Not run:
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
map.syn <- getPwayGenes("04010", attractor.states, removeGenes=remove.these.genes)
## End(Not run)
```

loring.eset

An ExpressionSet Object containing published data from M?ller et al.

Description

This is an ExpressionSet object containing the published data from M?ller et al. (NCBI GEO accession id GSE11508). The expression data set contains 11044 probes for 68 samples.

Usage

```
data(loring.eset)
```

Format

An ExpressionSet object.

Value

An ExpressionSet object containing the published data from M?ller et al. (NCBI GEO accession id GSE11508). The expression data set contains 11044 probes for 68 samples.

References

M?ller, F, et al., Regulatory networks define phenotypic classes of human stem cell lines. Nature, 2008. 455(7211): p. 401-405.

See Also

[exprs.dat](#), [samp.info](#)

Examples

```
data(loring.eset)
exprs.dat <- exprs(loring.eset) # gene expression matrix
```

plotsynexprs

Visualizing the Average Expression Profile of a Synexpression Group.

Description

This function plots the average expression profile for a specific synexpression group.

Usage

```
plotsynexprs(mySynExpressionSet, tickMarks, tickLabels, vertLines, index=1, ...)
```

Arguments

mySynExpressionSet	<code>SynExpressionSet</code> object.
tickMarks	numeric vector of specifying the location of the tick marks along the x-axis. There should be one tick for each cell type or group.
tickLabels	character vector specifying the labels to be appear underneath the tick marks on the x-axis. These should correspond to the cell type or group names.
vertLines	numeric vector specifying the location of the vertical lines that indicate the cell type or group-specific regions along the x-axis.
index	numeric value specifying which synexpression group should be plotted.
...	additional arguments.

Details

Generic plotting parameters can be passed to this function to create a more sophisticated plot, e.g `col="blue", main="Synexpression Group 1"`.

Value

A plot showing the average expression profile for the synexpression group specified.

Author(s)

Jessica Mar

Examples

```
data(subset.loring.eset)
attractor.states <- findAttractors(subset.loring.eset, "celltype", nperm=10, annotation="illuminaHumanv1.db")
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
mapk.syn <- findSynexprs("04010", attractor.states, remove.these.genes)
par(mfrow=c(2,2))
pretty.col <- rainbow(3)
for( i in 1:3 ){
  plotsynexprs(mapk.syn, tickMarks=c(6, 28, 47, 60), tickLabels=c("ESC", "PRO", "NSC", "TER"), vertLines=c(12.5, 28.5, 47.5, 60.5),
    main=paste("Synexpression Group ", i, sep=""), col=pretty.col[i])
}
```

removeFlatGenes	<i>Flags a set of genes which demonstrates little variation across the cell-types or experimental groups of interest.</i>
-----------------	---

Description

This function uses a linear model set up in `limma` to assess the degree of association between celltype and a gene's expression profile. In this way, we can flag those genes whose profiles show very little change across different celltype groups, or in other words are "flat".

Usage

```
removeFlatGenes(eSet, cellTypeTag, contrasts = NULL, limma.cutoff = 0.05, ...)
```

Arguments

eSet	ExpressionSet object.
cellTypeTag	character string of the variable name which stores the cell-lineages or experimental groups of interest for the samples in the data set (this string should be one of the column names of pData(myEset)).
contrasts	optional vector of contrasts that specify the comparisons of interest. By default, all comparisons between the different groups are generated.
limma.cutoff	numeric specifying the P-value cutoff. Genes with P-values greater than this value are considered "flat" and will be included in the set of flat genes.
...	additional arguments.

Details

Flat genes are removed from the analysis after the core attractor pathway modules are first inferred (i.e. the findAttractors step).

Value

A vector with gene names (as defined in the eset) of those genes with expression profiles that hardly vary across different celltype or experimental groups.

Author(s)

Jessica Mar

References

limma package.

Smyth, G. K. (2004). Linear models and empirical Bayes methods for assessing differential expression in microarray experiments. *Statistical Applications in Genetics and Molecular Biology* 3, No. 1, Article 3.

Examples

```
data(subset.loring.eset)
remove.these.genes <- removeFlatGenes(subset.loring.eset, "celltype", contrasts=NULL, limma.cutoff=0.05)
```

samp.info

samp.info Contains the Sample Information for the Mueller data set.

Description

This is sample information data frame for the samples in the Mueller data set (NCBI GEO accession id GSE11508). The data frame contains the cell type groups for the 68 samples.

Usage

```
data(samp.info)
```

Format

A data.frame object with one column of sample IDs (these are the column IDs of the exprs.dat expression matrix object) and second column indicating which cell type each sample represents.

ChipID A vector of sample IDs.

celltype A vector denoting the cell type a sample represents.

Value

A sample data frame for the samples in the Mueller data set (NCBI GEO accession id GSE11508). The data frame contains the cell type groups for the 68 samples.

References

Mueller F, et al., Regulatory networks define phenotypic classes of human stem cell lines. Nature, 2008. 455(7211): p. 401-405.

See Also

[exprs.dat](#), [loring.eset](#)

Examples

```
data(samp.info)
```

subset.loring.eset	<i>An ExpressionSet Object containing published data from Mueller et al.</i>
--------------------	--

Description

This is an ExpressionSet object containing a subset of the published data from Mueller et al. (NCBI GEO accession id GSE11508). The expression data set contains 5522 probes for 68 samples. This ExpressionSet object was created specifically to demonstrate the functions in this package. If you're looking for the full Mueller data set, see [loring.eset](#).

Usage

```
data(subset.loring.eset)
```

Format

An ExpressionSet object.

Value

An ExpressionSet object containing a subset of the published data from Mueller et al. (NCBI GEO accession id GSE11508). The expression data set contains 5522 probes for 68 samples.

References

Mueller, F, et al., Regulatory networks define phenotypic classes of human stem cell lines. Nature, 2008. 455(7211): p. 401-405.

See Also

[exprs.dat](#), [samp.info](#), [loring.eset](#)

Examples

```
data(subset.loring.eset)
subset.exprs.dat <- exprs(subset.loring.eset) # gene expression matrix
```

SynExpressionSet-class

Class SynExpressionSet

Description

This is a class representation for storing synexpression group information.

Objects from the Class

Objects are output by the function [findSynexprs](#). Objects can also be created by using `new("SynExpressionSet", ...)`.

Slots

groups: A list object denoting the probes or gene IDs (rnaseq) belonging to each synexpression group.

profiles: A matrix of average expression profiles for each synexpression group, each profile is stored as a row.

Methods

No methods have yet been defined with class "SynExpressionSet" in the signature.

Note

This class is described in more detail in the vignette.

Author(s)

Jessica Mar <jess@jimmy.harvard.edu>

Examples

```
new.synexpressionset <- new("SynExpressionSet", groups=list(), profiles=matrix(0))
```


Index

- * **aplot**
 - plotsynexprs, 20
- * **classes**
 - AttractorModuleSet-class, 3
 - SynExpressionSet-class, 24
- * **datasets**
 - exprs.dat, 11
 - loring.eset, 20
 - samp.info, 22
 - subset.loring.eset, 23
- * **internal**
 - buildCorMatrix, 4
 - buildKeggIncidenceMatrix, 6
 - findOnepwaySynexprs, 15
 - flagPwayExists, 17
 - getCustomGenes, 18
 - getPwayGenes, 19
- * **methods**
 - buildCustomIncidenceMatrix, 5
 - calcFuncSynexprs, 7
 - calcInform, 8
 - calcModfstat, 9
 - calcRss, 10
 - filterDataSet, 12
 - findAttractors, 12
 - findCorrPartners, 14
 - findSynexprs, 16
 - removeFlatGenes, 21
- * **package**
 - attract-package, 2

attract (attract-package), 2

attract-package, 2

AttractorModuleSet
(AttractorModuleSet-class), 3

AttractorModuleSet-class, 3

buildCorMatrix, 4

buildCustomIncidenceMatrix, 5

buildKeggIncidenceMatrix, 6

calcFuncSynexprs, 7

calcInform, 8

calcModfstat, 9

calcRss, 10

exprs.dat, 11, 20, 23, 24

filterDataSet, 12

findAttractors, 3, 12

findCorrPartners, 14

findOnepwaySynexprs, 15

findSynexprs, 16, 24

flagPwayExists, 17

getCustomGenes, 18

getPwayGenes, 19

loring.eset, 11, 20, 23, 24

plotsynexprs, 20

removeFlatGenes, 21

samp.info, 11, 20, 22, 24

subset.loring.eset, 23

SynExpressionSet, 17, 21

SynExpressionSet
(SynExpressionSet-class), 24

SynExpressionSet-class, 24