Package ‘eisa’

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Title Expression data analysis via the Iterative Signature Algorithm
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Description The Iterative Signature Algorithm (ISA) is a biclustering method; it finds correlated blocks (transcription modules) in gene expression (or other tabular) data. The ISA is capable of finding overlapping modules and it is resilient to noise. This package provides a convenient interface to the ISA, using standard BioConductor data structures; and also contains various visualization tools that can be used with other biclustering algorithms.

Depends isa2, Biobase (>= 2.17.8), AnnotationDbi, methods
Imports BiocGenerics, Category, genefilter, DBI
Suggests igraph (>= 0.6), Matrix, GOstats, GO.db, KEGG.db, biclust, MASS, xtable, ALL, hgu95av2.db, targetscan.Hs.eg.db, org.Hs.eg.db
License GPL (>= 2)

biocViews Classification, Visualization, Microarray, GeneExpression

Collate AllClasses.R AllGenerics.R ISAEexpressionSet-methods.R
            ISAModules-methods.R enrichment.R CHR.R GO.R KEGG.R miRNA.R
            generalenrichment.R

NeedsCompilation no

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The Iterative Signature Algorithm (ISA) is a biclustering method. ALLModules and ALLModulesSmall are example ISA biclusters (=modules) found in the ALL data set.

**Usage**

ALLModules

ALLModulesSmall

**Format**

Both ALLModules and ALLModulesSmall are instances of the ISAModules class.

**Source**

ISAModules was generated by calling ISA on the ALL data set, using the default parameters. ISAModulesSmall was generated the same way, but with gene threshold 2.7 and condition threshold 1.4 only.

**References**


**See Also**

The ALL BioConductor package.
condPlot

**Examples**

```r
data(ALLModules)
ALLModules
```

---

**condPlot**

*Plot sample scores of a transcription module*

---

**Description**

Creates a barplot of sample (=condition) scores, for a given transcription module. See details below.

**Usage**

```r
condPlot (modules, number, eset, col = "white", all = TRUE, sep = NULL,
           sepcol = "grey", val = TRUE, srt = 90, adj.above = c(0, 0.5),
           adj.below = c(1, 0.5), plot.only = seq_len(ncol(eset)), ...)
```

**Arguments**

- `modules`: An ISAModules object.
- `number`: An integer scalar, the module to plot.
- `eset`: An ExpressionSet or ISAExpressionSet object. This is needed for calculating the scores of the samples that are not in the module, see the `all` argument. If an ExpressionSet object is supplied, then it is normalised by calling ISANormalize on it.
- `col`: Color of the bars, it it passed to `barplot`, so it can be any format `barplot` accepts. E.g. it can be a character vector with different colors for the different bars.
- `all`: Logical scalar, whether to plot all samples (if `TRUE`, the default), or just the ones that are included in the module.
- `sep`: NULL or a numeric vector. If not NULL, then the bars are separated at the given positions with vertical lines. This is useful if you want to subdivide the samples into groups.
- `sepcol`: The color of the separating lines (see the `sep` argument), if they are plotted.
- `val`: Logical scalar, whether to add labels with the actual score values.
- `srt`: Numeric scalar, the rotation angle of the text labels, this is passed to the `text` function.
- `adj.above`: Adjustment of the text labels that are above the bars. This is passed to `text`, see its manual for details.
- `adj.below`: Adjustments of the text labels that are below the bars. This is passed to `text`, see its manual for details.
- `plot.only`: Numeric vector, if supplied it is used to plot a subset of samples only. By default all samples are plotted.
- `...`: Additional argument, to be passed to `barplot`.

---
condPlot creates a barplot for the sample scores of an ISA transcription module. Each sample is represented as a bar.

These plots are useful if you want to show that a given transcription module separates the samples into two (or more) groups. You can assign different colors to the samples, based on some external information, e.g. case and control samples can be colored differently.

In most cases the scores are between minus one and one, but this is not necessarily true. It is possible to assign scores to samples that are not part of the module, but this requires performing one (more precisely half) ISA iteration step. Currently the function always performs this extra step, even if the out-of-module samples are not plotted. Because the sample scores in a module are only approximately constant during ISA iteration, it might be possible that the plotted scores are slightly different than the ones stored in the modules variable.

Value
None.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
ISA and ISAModules.

Examples

```
data(ALLModulesSmall)
library(ALL)
data(ALL)

col <- ifelse(grepl("^B", ALL$BT), "darkolivegreen", "orange")
condPlot(ALLModulesSmall, 1, ALL, col=col)
```
enrichment

Usage

ISAEnrichment (modules, categories, ann = annotation(modules),
features = featureNames(modules), hgCutoff = 0.05,
correction = TRUE, correction.method = "holm")

Arguments

modules An ISAModules object, a set of ISA modules.
categories A named list of gene categories. The names of the entries are used as category
names. Each entry of the list must be a character vector containing Entrez gene
ids.
ann Character scalar. The annotation package to be used. By default it is taken from
the modules argument.
features Character vector. The names of the features. By default it is taken from the
modules argument.
hgCutoff Numeric scalar. The cutoff value to be used for the enrichment significance.
This can be changed later, without recalculating the test.
correction Logical scalar, whether to perform multiple hypothesis testing correction.
correction.method Character scalar, the multiple testing correction method to use. Possible values:
"holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See
the p.adjust function for details on these.

Details

This function performs enrichment analysis, based on user defined gene labels. It is useful if one
want to test ISA modules against databases, other than GO and KEGG.
The hypergeometric test, a version Fisher’s exact test, takes a gene label and a gene set (in our case
coming from an ISA module) and asks whether the number of genes in the set labelled by the label
is significantly more (or less) than what one would expect by chance.
ISAEnrichment performs the hypergeometric test for every module, for all user supplied gene
labels. The mapping from the probe ids on the array to Entrez Ids is done using the appropriate
chip annotation package.
ISAEnrichment currently cannot test for under-representation.

Value

A GeneralListHyperGResult object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

Bergmann S, Ihmels J, Barkai N: Iterative signature algorithm for the analysis of large-scale gene
Mar 11.
expPlot

Expression matrix plots for ISA modules

Description

These functions create an expression matrix plot for an ISA module. The gene and sample scores are also plotted.

Usage

expPlotCreate (eset, modules, which, norm = c("sample", "raw", "feature"))
expPlot (epo, scores = TRUE)
expPlotColbar (epo)

Arguments

eset An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied (and the norm argument is not set to ‘raw’), then it is normalised by calling ISANormalize on it. A subset of eset is selected that corresponds to the features included in modules.

norm Character constant, specifies whether and how to normalize the expression values to plot. ‘raw’ plots the raw expression values, ‘feature’ the expression values scaled and centered for each feature (=gene) separately and if ‘sample’ is specified then the expression values are centered and scaled separately for each sample.

modules An ISAModules object.

which Numeric scalar, which module to plot.

scores Logical scalar, whether to plot the scores as well.

epo An object returned by expPlotCreate.

See Also

ISAGO, ISACHR, ISAKEGG and ISAmiRNA for other enrichment calculations.

The Category package.

Examples

data(ALLModulesSmall)
library(hgu95av2.db)
entrez <- unique(unlist(mget(featureNames(ALLModulesSmall), hgu95av2ENTREZID)))
categories <- lapply(letters, function(x) sample(entrez, 100))
names(categories) <- letters
fakeEnrichment1 <- ISAEnrichment(ALLModulesSmall, categories, correction=FALSE)
fakeEnrichment2 <- ISAEnrichment(ALLModulesSmall, categories, correction=TRUE)
expPlot

Details

expPlotCreate creates an object that contains all data for performing the image plot and returns it. The reason for not plotting it directly is, that the size of the plot is usually different in different cases, and the opening of the graphics device is delayed until expPlotCreate returns.

In the returned object, the weight and height entries give the optimal size of the image, in pixels.
expPlot creates the expression plot.
expPlotColbar plots a color bar for the expression plot.

Value

expPlotCreate returns an ISAexpPlot object. It is a named list and has several entries, the important ones:

- width: Numeric scalar, the optimal width of the plot.
- height: Numeric scalar, the optimal height of the plot.

expPlot returns, invisibly, a named list with members:

- coords: A list with two entries: x and y, both numeric vectors of length two. They give the position of the actual expression matrix on the plot.
- gene.width: Numeric scalar, the width of one box on the image plot, in pixels; if the image size is exactly the suggested one.
- cond.height: Numeric scalar, the height of one box on the image plot, in pixels; if the image size is exactly the suggested one.

expPlotColbar returns NULL, invisibly.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

The vignette in the eisa package for other ISA visualizations. The ExpressionView package for an interactive version.

Examples

data(ALLModulesSmall)
library(ALL)
data(ALL)

ep <- expPlotCreate(ALL, ALLModulesSmall, 1)
ep
if (interactive()) {
  expPlot(ep)
}
gograph  

Plot part of the Gene Ontology hierarchy

Description
These functions help creating a plot of the Gene Ontology hierarchy.

Usage

```r
gograph (table, colbar.length = 30, label.cex = 1, alpha=1, abbrev=5,  
         GOGRAPHS = NULL, go.terms = NULL)
gographPlot (graph, coords = FALSE, ...)
```

Arguments

- `table`: A data frame with one column, containing the $p$-values of the enriched GO terms. The row names of the data frame should contain the GO ids.
- `colbar.length`: Numeric scalar, the length of the color bar.
- `label.cex`: Numeric scalar, factor for the label sizes, e.g. `2` means double size compared to the default.
- `alpha`: Alpha channel for the fill color of the vertices.
- `abbrev`: Numeric scalar, the minimum length for the abbreviated GO ids.
- `GOGRAPHS, go.terms`: These are for internal use only.
- `graph`: An igraph graph, as returned by the gograph function.
- `coords`: Logical scalar, whether to return the coordinates of the vertices on the plot.
- `...`: Additional arguments. These are passed to `plot.igraph`.

Details

A GO plot can be created in two steps. `gograph` creates an igraph graph object that contains all the information about the plot; `gographPlot` creates the actual plot.

The two steps are needed, because `gograph` calculates the optimal size of the plot, and then a graphics device of this size can be created before calling `gographPlot`.

The optimal size is returned by `gograph` in the `width` and `height` graph attributes, these can be queried with

```r
G <- gograph(...)  
G$width  
G$height
```

Value

- `gograph` returns an igraph object.
- `gographPlot` by default returns NULL, invisibly. If the `coords` argument is TRUE, then it returns the coordinates of the vertices on the plot.
ISA

Iterative Signature Algorithm on Gene Expression data

Description

Run ISA on an ExpressionSet with the default parameters.

Usage

ISA (data, flist = filterfun(function(x) IQR(x) > 0.5),
uniqueEntrez = TRUE, thr.gene = seq(2, 4, by = 0.5),
thr.cond = seq(1, 3, by = 0.5), no.seeds = 100)

Arguments

- data: The input, an ExpressionSet object.
- flist: A 'list' of filter functions to apply to the array. This is passed to the genefilter function without touching it. Supply NA here if you don't want to filter the expression set before running ISA on it.
- uniqueEntrez: Logical scalar, whether to filter the input expression set to keep exactly one probeset for each Entrez gene. Probesets that are not mapped to an Entrez gene are dropped.
- thr.gene: Numeric vector. The threshold parameters for the ISA, for features (=probesets or genes). All combinations of thr.gene and thr.cond will be used to run ISA.
thr.cond Numeric vector. The threshold parameters for the ISA, for samples. All combinations of thr.gene and thr.cond will be used to run ISA.

no.seeds Number of seeds to run ISA from.

Details

Please read tutorial vignette included in this package for an introduction on ISA. The isa2-package manual page in the isa2 package is also useful.

The ISA function performs the ISA algorithm on the supplied expression data. This involves the following steps:

1. Filtering the features (i.e. probe sets) according to their variance. You will need the genefilter package for this. The default filtering function keeps the features that have an IQR of 0.5 or more. See genefilter for details on how to create filtering functions. If NA is given as the flist argument, then no filtering is performed.

2. Filtering the features by mapping them to Entrez genes. Features that do not map to Entrez genes are removed from the data set. If more features map to the same Entrez gene, then only the one with the highest variance will be kept.

3. Calling the isa function in the isa2 package to perform the Iterative Signature Algorithm. This itself performs the following steps:
   (a) Normalizing the data by calling isa.normalize.
   (b) Generating random input seeds via generate.seeds.
   (c) Running ISA with all combinations of given feature and sample thresholds, by calling isa.iterate.
   (d) Merging similar modules, separately for each threshold combination, by calling isa.unique.
   (e) Filtering the modules separately for each threshold combination, by calling isa.filter.robust in the isa2 package.
   (f) Putting all modules from the runs with different thresholds into a single object.
   (g) Merging similar modules, across all threshold combinations, if two modules are similar, then the one with the milder thresholds is kept.

4. Creates an ISAModules object from the ISA results.

Value

An ISAModules-class object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


ISA-Biclust conversion

See Also
The vignette included in the eisa package.

Examples
```r
library(ALL)
data(ALL)
modules <- ISA(ALL, thr.gene=2.7, thr.cond=1.4)
modules
```

ISA-Biclust conversion

Convert ISA modules to a Biclust object, or the opposite

Description
The biclust package implements several biclustering algorithms in a unified framework. The result of the biclustering is a Biclust object. These functions allow the conversion between Biclust and ISAModules objects.

Usage
```r
annotate(biclusters, data)
```

Arguments
- `biclusters`: A Biclust object.
- `data`: An ExpressionSet object.

Details
To convert an ISAModules object (`mods`) to a Biclust object (`bc`), you can do:
```r
bc <- as(mods, "Biclust")
```
The seed data and run data of the ISAModules object is stored in the Parameters slot of the Biclust object. The ISA scores are binarized by the conversion.
To convert a Biclust object (`bc`) to an ISAModules object (`mods`), you can call:
```r
mods <- as(bc, "ISAModules")
```
The Parameters slot of the Biclust object is used as the run data of the ISAModules object. The seed data of the new object will be an empty data frame.
The annotate function puts biological annotation into a Biclust object. It is suggested to use it before converting the Biclust object to ISAModules, so that ISA visualization functions and enrichment calculations can make use of this information.

Value
annotate returns a Biclust object.
Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

```r
if (require(biclust)) {
  library(ALL)
  data(ALL)
  ALL.filtered <- ALL[sample(1:nrow(ALL), 1000),]

  # Biclust -> ISAModules
  set.seed(1)
  Bc <- biclust(exprs(ALL.filtered), BCPlaid(),
                fit.model = ~m + a + b, verbose = FALSE)
  Bc <- annotate(Bc, ALL.filtered)
  modules <- as(Bc, "ISAModules")
  Bc
  modules
  getNoFeatures(modules)
  getNoSamples(modules)

  # ISAModules -> Biclust
  data(ALLModulesSmall)
  Bc2 <- as(ALLModulesSmall, "Biclust")
  ALLModulesSmall
  getNoFeatures(ALLModulesSmall)
  getNoSamples(ALLModulesSmall)
  Bc2
}
```

---

**ISA2heatmap**

Heatmap of a transcription module

**Description**

Create a heatmap plot for an ISA module.

**Usage**

```r
ISA2heatmap (modules, module, eset, norm = c("raw", "feature", "sample"),
             scale = c("none", "row", "column"), ...)
```
**Arguments**

- **modules**
  - An ISAModules object.
- **module**
  - Numeric scalar, the number of the module to plot.
- **eset**
  - An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied (and the `norm` argument is not set to 'raw'), then it is normalised by calling ISANormalize on it. A subset of eset is selected that corresponds to the features included in `modules`.
- **norm**
  - Character constant, specifies whether and how to normalize the expression values to plot. ‘raw’ plots the raw expression values, ‘feature’ the expression values scaled and centered for each feature (=gene) separately and if ‘sample’ is specified then the expression values are centered and scaled separately for each sample.
- **scale**
  - Character scalar, passed to the heatmap function.
- **...**
  - Additional arguments, they are passed to the heatmap function.

**Value**

The same as `heatmap`.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

heatmap

**Examples**

```r
library(ALL)
data(ALL)
data(ALLModulesSmall)

if (interactive()) {
  ISA2heatmap(ALLModulesSmall, 1, ALL, norm="feature")
}
```
Description

Hypergeometric test(s) to check whether significantly many genes of an ISA module are on the same chromosome.

Usage

ISACHR (modules, ann = annotation(modules), features = featureNames(modules),
       hgCutoff = 0.05, correction = TRUE, correction.method = "holm")

Arguments

- **modules**: An ISAModules object, a set of ISA modules.
- **ann**: Character scalar. The annotation package to be used. By default it is taken from the modules argument.
- **features**: Character vector. The names of the features. By default it is taken from the modules argument.
- **hgCutoff**: Numeric scalar. The cutoff value to be used for the enrichment significance. This can be changed later, without recalculating the test.
- **correction**: Logical scalar, whether to perform multiple hypothesis testing correction.
- **correction.method**: Character scalar, the multiple testing correction method to use. Possible values: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See the p.adjust function for details on these.

Details

The hypergeometric test, a version Fisher’s exact test, takes a chromosome and a gene set (in our case coming from an ISA module) and asks whether the number of genes in the set that are on the given chromosome is significantly more (or less) than what one would expect by chance.

ISACHR performs the hypergeometric test for every module, for every chromosome. The chromosome mapping is taken from the annotation package of the chip.

ISACHR currently cannot test for under-representation.

Value

A CHRListHyperGResult object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

See Also

ISAGO, ISAKEGG and ISAmiRNA for other enrichment calculations.
The Category package.

Examples

data(ALLModulesSmall)
CHR <- ISACHR(ALLModulesSmall)
CHR
sigCategories(CHR)[[2]]
geneIdsByCategory(CHR)[[2]][[1]]
Two additional methods were defined to access the extra matrices: `featExprs` returns the feature-wise standardized data, `sampExprs` the sample-wise standardized one. The `hasNA` function returns TRUE if NA or NaN values appear in at least one of the expression matrices. The `prenormalized` function returns TRUE if the data was prenormalized, see `ISANormalize` for details.

**Value**

`featExprs` and `sampExprs` both return a matrix.

`hasNA` and `prenormalized` return a logical vector of length one.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

`ISANormalize`, `ExpressionSet` in the Biobase package.

**Examples**

```r
library(ALL)
data(ALL)

# Do the normalization
ALL.normed <- ISANormalize(ALL)
class(ALL.normed)
dim(exprs(ALL.normed))
dim(featExprs(ALL.normed))
dim(sampExprs(ALL.normed))

# Check that we indeed have Z-scores
all(abs(apply(featExprs(ALL.normed), 2, mean) ) < 1e-12)
all(abs(1-apply(featExprs(ALL.normed), 2, sd)) < 1e-12)

all(abs(apply(sampExprs(ALL.normed), 1, mean) ) < 1e-12)
all(abs(1-apply(sampExprs(ALL.normed), 1, sd)) < 1e-12)
```

**Description**

Robustness of ISA biclusters. The more robust biclusters are more significant, in the sense that they are less likely to be found in random data.
**Usage**

ISARobustness(data, isaresult)
ISAFilterRobust(data, isaresult, ...)

**Arguments**

- **data**: An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied, then it is normalised by calling `ISANormalize` on it.
- **isaresult**: An ISAModules object, a set of modules.
- **...**: Additional arguments, they are passed to the `isa.filter.robust` function in the `isa2` package.

**Details**

ISARobustness calculates robustness scores for ISA modules. The higher the score, the more robust the module.

ISAFilterRobust filters a set of ISA modules, by running ISA on the randomized expression data and then eliminating all modules that have a robustness score that is lower than at least one robustness score found in the randomized data.

The same feature and sample thresholds are used to calculate the randomized robustness scores. In other words the limit for the filtering depends on the feature and sample thresholds.

You can find more details in the manual of the `robustness` function in the `isa2` package.

**Value**

ISARobustness returns a numeric vector, the robustness scores of the biclusters.

ISAFilterRobust returns the filtered ISAModules instance.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

The `robustness` function in the `isa2` package.

**Examples**

data(ALLModules)
library(ALL)
data(ALL)
rob <- ISARobustness(ALL, ALLModules)
summary(rob)
Calculate Gene Ontology enrichment for transcription modules

Description
Gene Ontology enrichment is calculated for each ISA module separately. In the end the result is corrected for multiple hypothesis testing.

Usage
ISAGO (modules, ann = annotation(modules), features = featureNames(modules),
  hgCutoff = 0.05, correction = TRUE, correction.method = "holm")

Arguments
- **modules**: An ISAModules object, a set of ISA modules.
- **ann**: Character scalar. The annotation package to be used. By default it is taken from the modules argument.
- **features**: Character vector. The names of the features. By default it is taken from the modules argument.
- **hgCutoff**: Numeric scalar. The cutoff value to be used for the enrichment significance. This can be changed later, without recalculating the test.
- **correction**: Logical scalar, whether to perform multiple hypothesis testing correction.
- **correction.method**: Character scalar, the multiple testing correction method to use. Possible values: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See the p.adjust function for details on these.

Details
The Gene Ontology is a database of gene annotation. The annotating labels (these are called terms) are standardized and organized into a directed acyclic graph. In other words terms may have more specific sub-terms, that can have even more specific sub-sub-terms, and so on.

The Gene Ontology database has three big sub-graphs, the root nodes (the most general terms) of these are the direct children of the root term of the whole ontology: biological process, cellular component, molecular function. They are usually referred to as ontologies.

The hypergeometric test, a version Fisher’s exact test, takes a GO term and a gene set (in our case coming from an ISA module) and asks whether the number of genes in the set annotated by the term is significantly more (or less) than what one would expect by chance.

ISAGO performs the hypergeometric test for every module, for all GO terms of the three GO ontologies. The GO data is taken from the GO.db package and the annotation package of the chip.

ISAGO currently cannot test for under-representation and the conditional test, as implemented in the GOstats package, is not available either.

Value
A list with three **GOListHyperGResult** objects, for the three Gene Ontologies, named

- **BP** aka Biological Processes
- **CC** aka Cellular Components
- **MF** aka Molecular Function
Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
ISAKEGG, ISACHR, ISAmiRNA for other enrichment calculations.
The GO.db, GOstats and Category packages.

Examples
data(ALLModulesSmall)
GO <- ISAGO(ALLModulesSmall)
GO
summary(GO$BP)[[1]][,1:5]

Description
These functions create various sophisticated HTML pages from a set of ISA biclusters.

Usage
ISAHTMLTable (modules, target.dir, which = NULL,
template = system.file("autogen", package = "eisa"), GO = NULL,
KEGG = NULL, miRNA = NULL, CHR = NULL, htmltitle = NULL,
notes = NULL, seed = NULL, extra = list())

ISAHTMLModules (eset, modules, which = NULL, target.dir,
template = system.file("autogen", package = "eisa"), GO = NULL,
KEGG = NULL, miRNA = NULL, CHR = NULL, cond.to.include = NULL,
cond.col = "white", sep = NULL, seed = NULL, condPlot = TRUE)

ISAHTML (eset, modules, target.dir, template = system.file("autogen",
package = "eisa"), GO, KEGG, miRNA = NULL, CHR = NULL, htmltitle = NULL,
notes = NULL, seed = NULL, table.extra = list(), cond.to.include = NULL,
cond.col = "white", sep = NULL, condPlot = TRUE, which = NULL)
Arguments

modules An ISAModules object.
target.dir Character vector of length one, the directory in which the result is placed. It is created if it does not exist.
which Numeric vector, which modules to include in the table (ISAHTMLTable); or, which modules to create HTML pages for (ISAHTML and ISAHTMLModules). All modules are used if this argument is NULL, which is the default.
template The directory containing the HTML template files. By default the template included in the eisa package is used.
GO List of three GOListHyperGResult objects, as returned by the ISAGO function.
KEGG A KEGGListHyperGResult object, usually the output of the ISAKEGG function.
mirNA A mirNAListHyperGResult object, or NULL. See also the ISAmiRNA function.
CHR A CHRListHyperGResult object or NULL, see also the ISACHR function.
htmltitle Character vector of length one, the title of the HTML page.
notes Character vector of length one. Optional HTML text, on the default template it is placed on the top of the page, above the table.
seed Either NULL, or a character vector, with an optional column that is added to the module table.
extra Extra columns to put in the HTML table. It should be a named list of character vectors, each with the same length as the number of modules.
table.extra This is passed to ISAHTMLTable as the extra argument.
eset An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied, then it is normalised by calling ISANormalize on it.
cond.to.include Numeric or character vector, specifies which columns of the phenotype data of the original expression matrix are included in the tables of samples. By default the first six columns are included.
cond.col This is passed to condPlot as the col argument.
sep This is passed to condPlot as the sep argument.
condPlot Logical scalar, whether to create condition plots. If an alternative biclustering method was used to find the modules, then probably it makes no sense creating condition plots for them.

details

ISAHTMLTable creates an HTML page, a summary of the results of the modular analysis, including enrichment analysis of the modules.

ISAHTMLModules creates a separate HTML page for each module, including the following elements:

• An expression plot of the genes and samples in the module, including the ISA scores. This is done by calling expPlot.
• Gene Ontology tree plots for the enriched GO terms, separately for the three ontologies. These are produced by calling gograph.
• Tables for the enriched Gene Ontology terms, separately for the three ontologies.
• A table for the enriched KEGG pathways.
• A table for the enriched miRNA families.
The list of genes in the module.
The list of samples in the module.
A condition plot (if the condPlot argument is TRUE), see `condPlot`.

By default, clicking on the rows of the table generated by `ISAHTMLTable` is linked to the HTML page of the module, generated by `ISAHTMLModules`.

`ISAHTML` calls both `ISAHTMLTable` and `ISAHTMLModules`.

**Value**

These functions do not return a value. (They return `NULL`, invisibly.)

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

The vignette included in the eisa package.

**Examples**

```r
## Not run:
# Load data
library(ALL)
data(ALL)
data(ALLModulesSmall)

# Calculate enrichment
GO <- ISAGO(ALLModulesSmall)
KEGG <- ISAKEGG(ALLModulesSmall)
CHR <- ISACHR(ALLModulesSmall)

# Generate HTML summary
htmldir <- tempdir()
ISAHTML(ALL, modules=ALLModulesSmall, target.dir=htmldir,
          GO=GO, KEGG=KEGG, CHR=CHR)

# Open a browser to view the summary
if (interactive()) {
  browseURL(URLencode(paste("file://", htmldir, "/maintable.html", sep="")))
}

## End(Not run)
```
ISAIterate

Perform the Iterative Signature Algorithm

Description

ISAIterate performs the ISA on an ExpressionSet object, from the given input seeds.

Usage

ISAIterate(data, feature.seeds, sample.seeds, thr.feat, 
    thr.samp = thr.feat, ...) 

Arguments

data An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object 
is supplied, then it is normalised by calling ISANormalize on it.
feature.seeds A matrix of feature seeds. The number of rows should match the number of 
  features in the ExpressionSet, each column is a seed. Either this, or the 
sample.seeds argument must be given.
sample.seeds A matrix of sample seeds. The number of rows should match the number of 
  samples in the ExpressionSet, each column in a seed. Either this, or the 
  feature.seeds argument must be given.
thr.feat Numeric scalar or vector giving the threshold parameter for the features. Higher 
  values indicate a more stringent threshold and the result biclusters will contain 
  less features on average. The threshold is measured by the number of standard 
  deviations from the mean, over the values of the feature vector. If it is a vector, 
  then it must contain an entry for each seed.
thr.samp Numeric scalar or vector giving the threshold parameter for the columns. The 
  analogue of thr.feat.
... Additional arguments, these are passed to the isa.iterate function in the isa2 
  package. See also details below.

Details

Performs the ISA from the given seeds. It is allowed to specify both type of seeds, then a half-
iteration is performed on the sample.seeds and they are appended to the feature.seeds.

The isa.iterate function of the isa2 package is called to do all the work, this has the follow-
ning extra parameters: direction, convergence, cor.limit, eps, corx, oscillation, maxiter. 
Please see the isa.iterate manual for details about them.

Value

An ISAModules object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
References


See Also

The **ISA** function for an easier interface with parameters.

Examples

```r
library(ALL)
data(ALL)

# Only use a small sample, to make this example finish faster
ALL.normed <- ISANormalize(ALL)[sample(1:nrow(ALL), 1000),]

# Generate seeds and do ISA
seeds <- generate.seeds(nrow(ALL.normed), count=100)
modules <- ISAIterate(ALL.normed, seeds, thr.feat=3, thr.samp=2)
modules
```

**ISAKEGG**

*Calculate KEGG Pathway enrichment for transcription modules*

Description

KEGG pathway enrichment is calculated for each ISA module separately. In the end the result is corrected for multiple hypothesis testing.

Usage

```r
ISAKEGG(modules, ann = annotation(modules), features = featureNames(modules),
        hgCutoff = 0.05, correction = TRUE, correction.method = "holm")
```

Arguments

- `modules` : An ISAModules object, a set of ISA modules.
- `ann` : Character scalar. The annotation package to be used. By default it is taken from the modules argument.
- `features` : Character vector. The names of the features. By default it is taken from the modules argument.
- `hgCutoff` : Numeric scalar. The cutoff value to be used for the enrichment significance. This can be changed later, without recalculating the test.
- `correction` : Logical scalar, whether to perform multiple hypothesis testing correction.
- `correction.method` : Character scalar, the multiple testing correction method to use. Possible values: “holm”, “hochberg”, “hommel”, “bonferroni”, “BH”, “BY”, “fdr”, “none”. See the `p.adjust` function for details on these.
Details

KEGG (Kyoto Encyclopedia of Genes and Genomes) is a collection of online databases dealing with genomes, enzymatic pathways, and biological chemicals. The PATHWAY database records networks of molecular interactions in the cells, and variants of them specific to particular organisms. The hypergeometric test, a version Fisher’s exact test, takes a KEGG pathway and a gene set (in our case coming from an ISA module) and asks whether the number of genes in the set participating in the pathway, is significantly more (or less) than what one would expect by chance.

ISAKEGG performs the hypergeometric test for every module, for all KEGG pathways. The KEGG data is taken from the KEGG.db package and the annotation package of the chip.

ISAKEGG currently cannot test for under-representation.

Value

A KEGGListHyperGResult object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

http://www.genome.jp/kegg/


See Also

ISAGO, ISACHR, ISAmiRNA for other enrichment calculations.

The KEGG.db and Category packages.

Examples

data(ALLModulesSmall)
KEGG <- ISAKEGG(ALLModulesSmall)
KEGG
sigCategories(KEGG)[[1]]
summary(KEGG)[[1]][,1:5]

Description

This function performs enrichment calculations with respect to predicted miRNA targets to check whether an ISA module contains many genes that are targets of the same miRNA.
Usage

ISAmiRNA (modules, ann = annotation(modules), features = featureNames(modules),
          hgCutoff = 0.05, correction = TRUE, correction.method = "holm")

Arguments

modules       An ISAModules object, a set of ISA modules.
ann           Character scalar. The annotation package to be used. By default it is taken from
              the modules argument.
features      Character vector. The names of the features. By default it is taken from the
              modules argument.
hgCutoff      Numeric scalar. The cutoff value to be used for the enrichment significance.
              This can be changed later, without recalculating the test.
correction    Logical scalar, whether to perform multiple hypothesis testing correction.
correction.method
              Character scalar, the multiple testing correction method to use. Possible values:
              "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". See
              the p.adjust function for details on these.

Details

miRNAs are short RNA fragments that specifically regulate (usually inhibit) the expression
of genes. Some genes have been experimentally validated as targets of a given miRNA, but we
currently don’t know the target genes of most miRNAs.

TargetScan is a database of predicted miRNA targets. The predictions are done based many factors,
including the conservation of the target region during evolution.

The hypergeometric test, a version Fisher’s exact test, takes a miRNA and a gene set (in our case
coming from an ISA module) and asks whether the number of genes in the set regulated by the
miRNA is significantly more (or less) than what one would expect by chance.

ISAmiRNA performs the hypergeometric test for every module, for all miRNAs in the TargetScan
database.

In order to use this function, TargetScan annotation packages are needed.

Value

A miRNAListHyperGResult object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

Conserved Seed Pairing, Often Flanked by Adenosines, Indicates that Thousands of Human Genes
are MicroRNA Targets Benjamin P Lewis, Christopher B Burge, David P Bartel. Cell, 120:15-20
(2005).

Bergmann S, Ihmels J, Barkai N: Iterative signature algorithm for the analysis of large-scale gene
Mar 11.
See Also

ISAGO, ISAKEGG and ISACHR for other enrichment calculations.

The Category package.

Examples

data(ALLModulesSmall)

if (require(targetscan.Hs.eg.db)) {
  miRNA <- ISAmiRNA(ALLModulesSmall)
  summary(miRNA, p=0.1)[[7]]
}

---

**ISAModules-class**

* A set of ISA modules

**Description**

An ISAModules object stores the results of one ISA run. It contains a set of biclusters (=modules or transcription modules) and some meta information about the ISA run and the input data.

**Usage**

```r
## S4 method for signature 'ISAModules'
dim(x)
## S4 method for signature 'ISAModules'
featureNames(modules)
## S4 method for signature 'ISAModules'
sampleNames(modules)
## S4 method for signature 'ISAModules'
annotation(modules)
## S4 method for signature 'ISAModules'
getOrganism(modules)
## S4 method for signature 'ISAModules'
pData(modules)
## S4 method for signature 'ISAModules'
seedData(modules)
## S4 method for signature 'ISAModules'
runData(modules)
## S4 method for signature 'ISAModules'
featureThreshold(modules, mods)
## S4 method for signature 'ISAModules'
sampleThreshold(modules, mods)
## S4 method for signature 'ISAModules'
length(x)
## S4 method for signature 'ISAModules'
getNoFeatures(modules, mods)
## S4 method for signature 'ISAModules'
getNoSamples(modules, mods)
```
## S4 method for signature 'ISAModules'
getFeatures(modules, mods)
## S4 method for signature 'ISAModules'
getSamples(modules, mods)
## S4 method for signature 'ISAModules'
getFeatureNames(modules, mods)
## S4 method for signature 'ISAModules'
getSampleNames(modules, mods)
## S4 method for signature 'ISAModules'
getFeatureScores(modules, mods)
## S4 method for signature 'ISAModules'
getSampleScores(modules, mods)
## S4 method for signature 'ISAModules'
getFeatureMatrix(modules, binary = FALSE,
sparse = FALSE, mods)
## S4 method for signature 'ISAModules'
getSampleMatrix(modules, binary = FALSE,
sparse = FALSE, mods)
## S4 method for signature 'ISAModules'
getFullFeatureMatrix(modules, eset, mods)
## S4 method for signature 'ISAModules'
getFullSampleMatrix(modules, eset, mods)

## S4 method for signature 'ISAModules,ANY,ANY'
x[i, j, ..., drop = FALSE]
## S4 method for signature 'ISAModules,ANY,ANY'
x[[i, j, ..., drop = FALSE]]

### Arguments

- **x, modules**: An ISAModules object.
- **mods**: An optional numeric index vector for the modules. If given, the information is only returned only for the specified modules.
- **binary**: Logical scalar. Whether to binarize the feature or sample scores.
- **sparse**: Logical scalar. Whether to return a sparse matrix. The Matrix package is required for sparse matrices.
- **eset**: An ExpressionSet or ISAExpressionSet object. This is needed for calculating the scores of the features/samples that are not in the module. If an ExpressionSet object is supplied, then it is normalised by calling ISANormalize on it.
- **i**: For ‘[’ an index vector for selecting features (=probes, genes). For ‘[[’ an index vector for selecting modules.
- **j**: For ‘[’ an index vector for selecting samples. It is ignored for ‘[[’.
- **...**: Additional indexing arguments, they are not used, just ignored.
- **drop**: This argument is currently not used, just silently ignored.

### Details

An ISAModules object contains a set of biclusters, obtained using one run of the Iterative Signature Algorithm.

Various operations are defined such an object, here we document all of them, in several groups.
Value

dim returns a numeric vector of length two. featureNames and sampleNames return a character vector. annotation and getOrganism return a character vector of length one. pData returns a data frame.

seedData returns a data frame, see more below. runData returns a named list, see more below. featureThreshold and sampleThreshold return a numeric vector.

length returns a numeric scalar. getNoFeatures and getNoSamples return a numeric vector. getFeatures and getSamples return a list of named numeric vectors. getFeatureNames and getSampleNames return a list of character vectors. getFeatureScores and getSampleScores return a list of named numeric vectors. getFeatureMatrix, getSampleMatrix, getFullFeatureMatrix and getFullSampleMatrix return a numeric matrix.

Information about the input data.

dim returns the dimension of the input expression matrix, number of features times number of samples.

featureNames returns a character vector, the names of the features in the original input matrix. I.e. in the input was an ExpressionSet for an Affymetrix array, then the Affymetrix probe IDs are returned.

sampleNames returns a character vector, the names of the samples in the original expression set.

annotation returns a character scalar, the name of the array for the input expression set. More precisely, the annotation slot of the input ExpressionSet is returned, this is the name of the annotation package to use for the ExpressionSet.

getOrganism returns the scientific name of the organism for which the input expression data was measures. This is obtained by loading the annotation package of the input ExpressionSet object, so that must be installed.

pData returns the phenotypic data attached to the input ExpressionSet object, in a data frame, samples as rows and various phenotypic variables as columns.

Information about the ISA run

seedData returns information about the modules. Each row of the returned data frame corresponds to one module, the columns are various variables:

iterations The number of ISA iterations needed to find the module.

oscillation The length of the oscillation cycle for oscillating modules, zero for others.

thr.row The feature (=gene) threshold used for finding the module.

thr.col The sample (=condition) threshold used for finding the module.

freq The number of times the module was found. This is always one, unless ISAUnique was performed.

rob The robustness score of the module. See ISARobustness for details.

rob.limit The robustness limit that was used for filtering the modules. As this depends of the feature and sample thresholds, it may be different for different modules.

runData returns information about the ISA runs, it is a named list with elements:

annotation The annotation package corresponding to the input expression set.

organism The scientific name of the organism.
**direction**  The direction parameter of the ISA. Please see **ISAIterate** for details.

**convergence**  The method to determine ISA convergence, a character scalar. Please see **ISAIterate** for details.

**cor.limit**  Correlation limit for the "cor" convergence criterium, see **ISAIterate** for details.

**eps**  Difference limit for the "eps" convergence criterium, see **ISAIterate** for details.

**corx**  Size of the time window for the "corx" convergence criterium, see **ISAIterate** for details.

**maxiter**  The maximum number of ISA iterations that was allowed.

**oscillation**  Logical, whether oscillating modules were considered during the ISA iteration.

**N**  Numeric scalar, the number of input seeds that were used to find the modules.

**unique**  Logical scalar, whether **ISAUnique** was run on the modules.

**prenormalize**  Logical scalar, whether the input data was prenormalized during ISA normalization, see **ISANormalize**.

**hasNA**  Logical scalar, whether the normalized input data contained some NA or NaN values.

**rob.perms**  Numeric scalar, the number of times the input data was scrambled when the modules were filtered according to robustness.

Note that some of these might be missing, i.e. **rob.perms** is only present if **ISAFilterRobust** was performed.

**featureThreshold** returns the feature thresholds that were used to find the modules.

**sampleThreshold** returns the sample thresholds that were used to find the modules.

**Information about the modules**

**length** returns the number of modules.

**getNoFeatures** returns the number of features (=genes) in the input data. The number of features after filtering is returned if the input data was filtered.

**getNoSamples** returns the number of samples (=conditions) in the input data.

**Retrieve the modules**

**getFeatures** returns the indices of the features included in the modules. It returns a list, with one entry for each module. Each entry contains the indices of the features (=genes) in the corresponding module.

**getSamples** does the same as **getFeatures**, but for samples.

**getFeatureNames** is similar to **getFeatures**, but returns feature names instead of feature indices.

**getSampleNames** is similar to **getFeatures**, but returns sample names instead of sample indices.

**getFeatureScores** returns the feature scores for the selected modules (all modules by default). It returns a list, with one entry for each module. Each list entry contains the feature scores for one module, in a named numeric vector.

**getSampleScores** is similar to **getFeatureScores**, but for samples and sample scores.

**getFeatureMatrix** returns feature scores for the specified modules (all modules by default) in a matrix form. The number of rows is the number of features and the number of columns is the number of modules requested. It can optionally binarize the values.

**getSampleMatrix** is similar to **getFeatureMatrix**, but for sample scores.

**getFullFeatureMatrix** is similar to **getFeatureMatrix**, but is also calculates scores for the features that were not included in the module. For this it performs one ISA iteration and omits the
thresholding step. You need to supply the normalized (or the original) expression data to make this possible.

getFullSampleMatrix is the same as getFullFeatureMatrix, but for sample scores.

Indexing

A couple of indexing operations were defined to make it easier selecting subsets of modules, features or samples from an ISAModules object.

The ‘[]’ double bracket indexing operator can be used with a single index vector to select a subset of modules.

The ‘[’ single bracket indexing operator can be used to restrict an ISAModules object to a subset of features and/or samples. The first index corresponds to features, the second to samples. Indices can be numeric, logical or character vectors, for the latter feature and sample names are used.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

The vignette included in the eisa package.

Examples

data(ALLModulesSmall)
ALLModulesSmall

length(ALLModulesSmall)
dim(ALLModulesSmall)
annotation(ALLModulesSmall)
getOrganism(ALLModulesSmall)

seedData(ALLModulesSmall)

getNoFeatures(ALLModulesSmall)
getNoSamples(ALLModulesSmall)

getFeatureScores(ALLModulesSmall, 1)[[1]]
**ISANormalize**

**Normalize expression data for the Iterative Signature Algorithm**

**Description**

ISA works best if the input data is centered and scaled. ISANormalize performs this transformation.

**Usage**

```
ISANormalize (data, prenormalize = FALSE)
```

**Arguments**

- `data`: An ExpressionSet object.
- `prenormalize`: If this argument is set to TRUE, then feature-wise scaling is calculated on the sample-wise scaled matrix and not on the input matrix directly.

**Details**

It was observed that the ISA works better if the input matrix is scaled and its rows have mean zero and standard deviation one.

An ISA step consists of two sub-steps, and this implies two different normalizations, in the first the rows (=features), in the second the columns (=samples) of the input matrix will be scaled and centered.

**Value**

An ISAExpressionSet object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

The ISA function for an easier ISA workflow.

**Examples**

```r
library(ALL)
data(ALL)

# Do the normalization
ALL.normed <- ISANormalize(ALL)
class(ALL.normed)
dim(exprs(ALL.normed))
```
dim(featExprs(ALL.normed))
dim(sampExprs(ALL.normed))

# Check that we indeed have Z-scores
all(abs(apply(featExprs(ALL.normed), 2, mean) ) < 1e-12)
all(abs(1-apply(featExprs(ALL.normed), 2, sd)) < 1e-12)

all(abs(apply(sampExprs(ALL.normed), 1, mean) ) < 1e-12)
all(abs(1-apply(sampExprs(ALL.normed), 1, sd)) < 1e-12)

---

**ISASweep**

*Create an ISA module tree*

---

**Description**

These functions create and plot the hierarchical description of an expression data set, by applying the ISA with various thresholds, and connecting the related modules. See details below.

**Usage**

```
ISASweep (expset, modules, ...)
ISASweepGraph (sweep.result)
ISASweepGraphPlot (graph, vertex.label=V(graph)$id,
  vertex.label.topleft=NA, vertex.label.topright=NA,
  vertex.label.bottomleft=NA, vertex.label.bottomright=NA,
  vertex.label.cex=0.8, edge.label=NA, asp=FALSE, rescale=FALSE,
  xlim=range(graph$layout[,1]), ylim=range(graph$layout[,2]),
  thresholds=TRUE, xlab=NA, ylab=NA, ...)
```

**Arguments**

- `expset` The expression set object, if it is not an ISAExpressionSet, then ISANormalize is called on it.
- `modules` An ISAModules object.
- `...` Additional arguments. ISASweep passes these to isa.sweep; ISASweepGraphPlot passes additional arguments to plot.igraph.
- `sweep.result` An ISAModules object that contains the sweep tree information as well.
- `graph` An igraph graph object, the sweep tree.
- `vertex.label` Vertex labels, by default the ids of the modules.
- `vertex.label.topleft` Vertex labels to put at the top left corner.
- `vertex.label.topright` Vertex labels to put at the top right corner.
- `vertex.label.bottomleft` Vertex labels to put at the bottom left corner.
- `vertex.label.bottomright` Vertex labels to put at the bottom right corner.
- `vertex.label.cex` Magnification factor for the vertex labels.
edge.label  Edge labels.
as  Logical scalar, whether the plot should have 1:1 aspect ratio.
rescale  Logical scalar, whether to rescale the layout coordinates to the [-1,1] interval.
xlim  Numeric vector of length two, the X limits of the plot.
ylim  Numeric vector of length two, the Y limits of the plot.
thresholds  Logical scalar, whether to add the (non-constant) thresholds to the plot.
xlab  The label of the horizontal axis, by default omitted.
ylab  The label of the vertical axis, by default omitted.

Details

The ISA uses two threshold parameters that tune the sizes of the transcription modules. The sweep graph of an expression set is defined as the following. It is a directed graph, where the vertices are ISA modules, found at some threshold parameter values. There is an edge from module A to module B, if using 1) (the genes of) module A as the seed vector and 2) the threshold parameters used to find module B, the ISA converges to module B.

The ISASweep function creates an ISA sweep tree, in which one threshold parameter is kept fixed and the other varies. It starts from the modules found at the most stringent (=highest) threshold parameters, and uses them individually as seeds at the next less stringent threshold level. If this ISA iteration converges to an already known module, then an edge of the sweep tree is found. If the iteration converges to a new module, then this is added to the module list, together with the sweep tree edge. Then we proceed with the next level of modules, towards the less stringent threshold parameters.

The ISASweepGraph function creates a graph object that corresponds to the sweep tree of the expression set.

The ISASweepGraphPlot function plots a graph created with ISASweepGraph.

Value

ISASweep returns an ISAModules object, with some seed data added.
ISASweepGraph returns an igraph graph object.
ISASweepGraphPlot returns NULL, invisibly.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

library(genefilter)
library(ALL)
data(ALL)

varLimit <- 0.5
kLimit <- 4
ALimit <- 5
flist <- filterfun(function(x) var(x)>varLimit, kOverA(kLimit,ALimit))
ALL.filt <- ALL[genefilter(ALL, flist),]
ALL.filt2 <- ALL.filt[, grepl("^B", ALL.filt$BT)]

# Run ISA
set.seed(2)
modules <- ISA(ALL.filt2, flist=NA, thr.gene=seq(2,4,by=0.5), thr.cond=1)

# Do the sweep
modules2 <- ISASweep(ALL.filt2, modules)
modules2

# Plot it
## Not run:
    G <- ISASweepGraph(modules2)
    ISASweepGraphPlot(G)
## End(Not run)

---

**ISAUnique**  
Remove duplicated ISA modules

**Description**

From a potentially non-unique set of ISA modules remove all modules that are similar to another module that was found earlier.

**Usage**

ISAUnique(data, isaresult, ...)

**Arguments**

data    An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied, then it is normalised by calling ISANormalize on it.

isaresult    An ISAModules object to be filtered.

...    Additional arguments, these are passed to the isa.unique function in the Isa2 package. See also details below.

**Details**

The ISA algorithm might very well find the same modules from many different input seeds, so the output of the ISAIterate function is usually not unique: many modules are very similar to each other.

ISAUnique eliminates the duplicates and potentially also the non-convergent modules.

The work is performed by calling the isa.iterate function in the isa2 package. The following additional arguments can be specified to be passed to this function:

**method** Character scalar giving the method to be used to determine if two biclusters are similar. Right now only ‘cor’ is implemented, this keeps both biclusters if their Pearson correlation is less than cor.limit, both for their row and column scores. See also the neg.cor argument.
ignore.div Logical scalar, if TRUE, then the divergent biclusters will be removed.
cor.limit Numeric scalar, giving the correlation limit for the ‘cor’ method.
neg.cor Logical scalar, if TRUE, then the ‘cor’ method considers the absolute value of the correlation.
drop.zero Logical scalar, whether to drop biclusters that have all zero scores.

Value
Another ISAModules object, with unique modules.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
The ISA function for an easier ISA workflow.

Examples
library(ALL)
data(ALL)

# Only use a small sample, to make this example finish faster
ALL.normed <- ISANormalize(ALL)[sample(1:nrow(ALL), 1000),]

# Generate seeds and do ISA
seeds <- generate.seeds(nrow(ALL.normed), count=100)
modules <- ISAIterate(ALL.normed, seeds, thr.feat=3, thr.samp=2)
modules

# Merge the modules
modules2 <- ISAUnique(ALL.normed, modules)
modules2
Usage

```r
## S4 method for signature 'ListHyperGParams'
makeValidParams(object)
## S4 method for signature 'ListHyperGParams'
drive(p)
## S4 replacement method for signature 'ListHyperGParams,logical'
drive(p) <- dri

## S4 method for signature 'GOListHyperGParams'
ontology(object)
## S4 replacement method for signature 'GOListHyperGParams,character'
ontology(object) <- go
## S4 method for signature 'GOListHyperGParams'
conditional(r)
## S4 replacement method for signature 'GOListHyperGParams,logical'
conditional(r) <- cond

## S4 method for signature 'ListHyperGParams'
hyperGTest(p)
```

Arguments

- `object`: A `ListHyperGParams` object.
- `p`: An object of class `ListHyperGParams`.
- `r`: A logical scalar, whether to store the genes that are in the intersection of the specified gene set and the annotation category.
- `go`: A character scalar, the ontology for GO, possible values: 'BP', 'CC', 'MF'.
- `cond`: A logical scalar, whether to perform conditional enrichment calculation. Currently this option is ignored.

Details

The `ListHyperGParams` abstract class extends `HyperGParams` and allows to specify a list of gene sets for the enrichment calculation instead of a single set.

`ListHyperGParams` calculates the enrichment much faster than the original `HyperGParams` classes in the `Category` package, especially if the calculation is performed against the same gene universe for many gene sets.

`ListHyperGParams` is an abstract class, it is not possible to instantiate objects from it. Instead, its various extensions must be used: `GOListHyperGParams`, `KEGListHyperGParams`, `CHRListHyperGParams` and `miRNAListHyperGParams`.

The various `ListHyperGParams` objects can be created with the standard `new` command, by giving all necessary arguments. Please see the examples below.

Value

- `makeValidParams` returns another `ListHyperGParams` instance that has the same class as its arguments.
- `ontology` returns a character vector of length one.
- `conditional` returns a logical vector of length one.
- `drive` returns a logical vector of length one.
Member functions

Most of these functions are analogous to the ones defined in the Category package, the only difference is that they handle ListHyperGParams objects.

makeValidParams validates ListHyperGParams object, in particular, it removes duplicate genes, both from the gene universe and the specified gene sets; and it also makes sure that all genes in the gene sets are included in the universe.

ontology can be used to query or set the ontology for enrichment calculated against the GO database.

conditional queries or sets whether conditional GO enrichment will be performed. This feature is not implemented yet, see the Category and GOstats packages for a working implementation and more information.

drive queries or sets whether the intersections of the gene sets and the universe are stored in the result object. This information can be calculated later as well, but it is faster to store it at the same time when the hypergeometric test is performed.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

Functions for enrichment calculation of ISA modules: ISAGO, ISAKEGG, ISACHR, ISAmiRNA.
Perhaps see also the vignette in the GOstats package.

Examples

# GO enrichment, "by hand"
# Load data first
data(ALLModulesSmall)

# Create gene sets
library(hgu95av2.db)
genes <- getFeatureNames(ALLModulesSmall)
extrez <- lapply(genes, function(x) na.omit(unlist(mget(x, hgu95av2ENTREZID))))

# Create universe
universe <- na.omit(unlist(mget(featureNames(ALLModulesSmall), hgu95av2ENTREZID)))

# Create parameter object
param <- new("GOListHyperGParams", geneIds = entrez, universeGeneIds = universe,
 pvalueCutoff = 0.01, drive = FALSE, ontology = "BP",
 conditional = FALSE, testDirection = "over",
 annotation = annotation(ALLModulesSmall))

# Do the calculation
GOBP <- hyperGTest(param)

# Inspect the result
GOBP
summary(GOBP)[[1]]
# How to create other parameter objects

```r
paramKEGG <- new("KEGGListHyperGParams", geneIds=entrez,
                  universeGeneIds=universe, drive=FALSE,
                  annotation=annotation(ALLModulesSmall))

paramCHR <- new("CHRListHyperGParams", geneIds=entrez,
                 universeGeneIds=universe, drive=FALSE,
                 annotation=annotation(ALLModulesSmall))
```

# Enrichment with user-supplied categories, we use a list of
# hand-picked genes that are involved in myelin formation

```r
mygenes <- c("YARS", "NFKB2", "NGFR", "COH1", "NFAT5", "NDRG1", "GAP43",
             "EGR2", "MSN", "ROCK1", "SREBF2", "SOX10", "FIG4", "EGR1", "PIK3R1",
             "CDC42", "EDN3", "EDNRB", "NCAM1", "DHH", "OMG", "PMP22", "LAMA4",
             "MPDZ", "MTMR2", "REL", "S100A1", "ITGA4", "GFAP", "FGF2", "RPSA",
             "CADM1", "CDH19", "DNM2", "PAX3", "SREBF1", "DAG1", "DRP2", "SOD2",
             "MBO", "RELA", "RELB", "JUN", "NAB1", "MOBP", "SIX", "COL5A2", "RHOA",
             "NFASC", "NFE2L2", "MPO", "MAG", "EDNRA", "ERBB4", "LITAF", "MMP2",
             "PLP1", "CDKN1A", "PAK1", "ROX", "GB1", "LAMA5", "JAM3", "ITGB1",
             "PAR03", "FABP7", "LAMA2", "ERBB3", "CADM4", "FOXO4", "TSPAN31",
             "GPR126", "PTK2", "RAC1", "CDKN2A", "CLDN5", "D2", "LAMC1", "SOX2",
             "CNTN2", "ERBB2", "NFKB1", "NAB2", "EN2", "MMP9", "CCND1", "L1CAM",
             "MOG")
```

```r
library(org.Hs.eg.db)
myentrez <- na.omit(mapIds(org.Hs.eg.db, mygenes, 'ENTREZID',
                            keytype='SYMBOL'))
categories <- list(myelin=myentrez)
```

```r
data(ALLModules)
genes2 <- getFeatureNames(ALLModules)
entrez2 <- lapply(genes2, function(x) na.omit(unlist(mget(x, hgu95av2ENTREZID))))
```

# Create universe

```r
universe2 <- na.omit(unlist(mget(featureNames(ALLModules), hgu95av2ENTREZID)))
```

```r
paramMY <- new("GeneralListHyperGParams", geneIds=entrez2,
                universeGeneIds=universe2, drive=FALSE,
                annotation=annotation(ALLModulesSmall),
                categories=categories)
```

```r
MY <- hyperGTest(paramMY)
MY
summary(MY)[[1]]
```

---

**ListHyperGResult-class**

*Classes for quick GO/KEGG/CHR/miRNA target or other enrichment calculation for multiple gene sets*

**Description**

These classes extend the HyperGResult class from the Category package to perform enrichment calculation quickly for multiple gene sets.
ListHyperGResult-class

Usage

## S4 method for signature 'ListHyperGResult'
summary(object, pvalue = pvalueCutoff(object),
categorySize = NULL)
## S4 method for signature 'ListHyperGResult'
htmlReport(r, file = "", append = FALSE,
label = "", digits = 3, summary.args = NULL)
## S4 method for signature 'ListHyperGResult'
pvalues(r)
## S4 method for signature 'ListHyperGResult'
sigCategories(r, p)

## S4 method for signature 'ListHyperGResult'
geneCounts(r)
## S4 method for signature 'ListHyperGResult'
expectedCounts(r)
## S4 method for signature 'ListHyperGResult'
oddsRatios(r)
## S4 method for signature 'ListHyperGResult'
universeCounts(r)
## S4 method for signature 'ListHyperGResult'
geneMappedCount(r)
## S4 method for signature 'ListHyperGResult'
universeMappedCount(r)
## S4 method for signature 'ListHyperGResult'
geneIdsByCategory(r, catids = NULL)

## S4 method for signature 'ListHyperGResult'
geneIdUniverse(r, cond = FALSE)

Arguments

object,r A ListHyperGResult object.
pvalue,p Numeric vector of length one, the \( p \)-value limit, up to which the terms are listed.
categorySize A numeric vector of length one, or NULL. If not NULL, then it gives the minimum number of annotated genes in the universe, in order to list the term.
file A file name, or a connection object. The result is written here. If it is "", then the result is written to the standard output. If it is NULL, then the result is not written anywhere. (But it is always returned, invisibly, see below.)
append Logical scalar, whether to append the HTML code to the given file, or remove its previous contents if it already exists.
label An HTML label (<A LABEL="""> tag) to add.
digits The number of digits to use for the numeric columns.
summary.args A list of arguments to pass to the summary method.
catids The categories for which the genes are listed. All categories will be listed if this argument is NULL.
cond Currently not used.
Details

A ListHyperGResult object can store the results of hypergeometric tests, several gene sets against the same universe. ListHyperGResult is an extension of HyperGResult, as defined in the Category package.

More precisely, ListHyperGResult is an abstract class, it is not possible to instantiate objects from it. Its extensions be used instead: GOListHyperGResult, KEGGListHyperGResult, CHRListHyperGResult and miRNAListHyperGResult.

Value

pvalues, geneCounts, expectedCounts, oddsRatios and universeCounts return a list of named numeric vectors.

geneMappedCount returns a numeric vector, universeMappedCount returns a numeric vector of length one.

sigCategories returns a list of character vectors.

geneIdsByCategory returns a list of lists of character vectors.

geneIdUniverse returns a list of character vectors.

summary returns a list of data frames with columns: 'Pvalue', 'OddsRatio', 'ExpCount', 'Count', 'Size' and optionally 'drive'.

htmlReport returns a list of character vectors, invisibly.

conditional returns a logical vector of length one. ontology returns a character vector of length one.

Member functions

Most of the member functions are analogous to the ones defined for HyperGResult in the Category package. Usually the only difference is that they return a list of vectors, with one entry for each gene set, instead of just a single vector.

pvalues returns the $p$-values of the hypergeometric tests. A list is returned, with one numeric vector entry for each input gene set. The $p$-values for each gene set are ordered according to decreasing significance.

geneCounts returns the number of genes from the gene set that are annotated with the given term. This is returned for all input gene sets, in a list.

expectedCounts returns the number of genes that are expected to be annotated with the given term, just by chance. This is calculated for all input gene sets, and returned as a list.

oddsRatios returns the odds ratios for each term tested, for all gene sets, in a list of numeric vectors.

universeCounts returns the number of genes from the universe that are annotated with the given term, for all gene sets, in a list.

geneMappedCount gives the size of the gene sets, as used in the algorithm. This can be different than the size of the input gene sets, because of the elimination of duplicates and genes that are not in the universe, before the actual computation.

universeMappedCount gives the size of the gene universe, as used in the computation. This can be different than the size given by the user, because duplicates are eliminated before the computation.

sigCategories returns the significant terms, at the given $p$-value threshold, for all gene sets, as a list.
geneIdsByCategory returns a list of lists, one entry for each input gene set. Every entry is a list itself and for each tested term it gives the gene ids from the gene set that are annotated with the given term.

geneIdUniverse returns a list of character vectors, one for each term that was tested, giving the ids of the genes from the universe that are annotated with that term.


htmlReport creates a HTML summary from a ListHyperGParams object. This consists of one table for each input gene set. The summary can be written to a file, but it is also returned in a list of character vectors. There is one list entry for each input gene set, and each element of the character vector corresponds to one line of HTML code. You need the xtable package to use this function.

The following functions are defined for GOlistHyperGResult objects only.

conditional returns a logical vector of length one, whether the test was conditional or not. Conditional testing is currently not implemented, please see the GOstats package for a working implementation.

ontology returns a character vector of length one, the name of the ontology for the GO test.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
Functions for enrichment calculation of ISA modules: ISAGO, ISAKEGG, ISACHR, ISAmiRNA, ISAEnrichment. Perhaps see also the vignette in the GOstats package.

Examples

data(ALLModulesSmall)
GO <- ISAGO(ALLModulesSmall)
GO$CC
sigCategories(GO$CC)[[1]]
summary(GO$CC)[[1]][,1:5]

mnplot  
Plot group means against each other, for an ISA module

Description
Plot mean expression values for two sets of samples, against each other.

Usage

mnplot (x, expset, group, ...)
ISAmnplot (modules, number, eset, norm = c("raw", "feature", "sample"), group, ...)
Arguments

x    A character vector, the feature names for which the plot is created.
expset    An ExpressionSet object (Biobase package), or an expression matrix, with row names as feature names.
eset    An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied (and the norm argument is not set to ’raw’), then it is normalised by calling ISANormalize on it. A subset of eset is selected that corresponds to the features included in modules.
norm    Character constant, specifies whether and how to normalize the expression values to plot. ’raw’ plots the raw expression values, ’feature’ the expression values scaled and centered for each feature (=gene) separately and if ’sample’ is specified then the expression values are centered and scaled separately for each sample.
group    A factor that defines two groups to plot one against the other.
modules    An ISAModules object.
number    A numeric scalar, the number of the module for which the plot is created.
...    Additional arguments, they are passed to the plot function.

Details

mnplot plots two group-means against each other, the mean expression of all the specified probes. The two groups are specified as a factor with two levels.

ISAmnplot calls mnplot and plots the mean expression of genes in an ISA module, again, for two groups.

Value

Both functions return invisibly a matrix with two lines, the mean expression values for the two groups, for all the specified genes.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

The GOmnplot and KEGGmnplot functions in the annotate package.

Examples

data(ALLModulesSmall)
library(ALL)
data(ALL)
group <- ifelse(grepl("B", ALL$BT), "B-cell", "T-cell")
ISAmnplot(ALLModulesSmall, 2, ALL, norm="feature", group=group)
**Overlap of ISA biclusters**

**Description**

Plots a network, where each node is a module and modules that overlap are closer to each other.

**Usage**

```r
overlap (modules, algorithm = c("mds", "fr", "drl"), edge.limit = 0.5)
overlapPlot (graph, xsize = 400, ysize = 400, vertex.size = 20,
             vertex.size2 = 10, ...)
```

**Arguments**

- `modules` An `ISAModules` object.
- `algorithm` The algorithm to use for placing the vertices, a character scalar. See details below.
- `edge.limit` Numeric constant between zero and one, only edges between modules that have a Pearson correlation higher than `edge.limit` will be drawn.
- `graph` An igraph object, as returned by `overlap`.
- `xsize` The width of the plot in pixels, only used to calculate the return value, it does not influence the plot itself.
- `ysize` The height of the plot in pixels, only used to calculate the return value, it does not influence the plot itself.
- `vertex.size` The width of the vertices on the plot.
- `vertex.size2` The height of the vertices on the plot.
- `...` Additional arguments, these are passed to the `plot.igraph` function from the igraph package.

**Details**

An `ISAModules` object may potentially contain many modules that overlap. These functions visualize the overlapping relationships of a set of modules.

`overlap` creates an igraph graph with additional information on how to plot this graph in a way that nodes representing overlapping modules are close to each other.

`overlapPlot` takes such a graph and plots it.

`overlap` can use various algorithms, depending on the `algorithm` argument. If it is ‘mds’, then multi-dimensional scaling is used, by calling the `isaMDS` function in the MASS package. If it is ‘fr’, then the Fruchterman-Reingold algorithm is used, through the `layout.fruchterman.reingold` function of the igraph package. If it is ‘drl’, then the DrL graph layout algorithm is used, see the `layout.drl` function in the igraph0 package.

**Value**

- `overlap` returns an igraph graph.
- `overlapPlot` returns the coordinates of the vertices in a two-column matrix, invisibly.
profilePlot

Profile plots for ISA biclusters

Description

Line plots to compare biclusters to the background, i.e. the rest of the expression matrix.

Usage

profilePlot(modules, module, eset, plot = c("samples", "features", "both"), norm = "default", background = TRUE, col = gray(0.7), col.mod = 1, type = "l", type.mod = type, mean = TRUE, meancol = "green", meancol.mod = "red", xlabs = c("Features", "Samples"), ylab = "Expression", ...)

Arguments

- **modules**: An ISAModules object.
- **module**: Numeric scalar, the module to plot.
- **eset**: An ExpressionSet or ISAExpressionSet object. If an ExpressionSet object is supplied (and the norm argument is not set to 'raw'), then it is normalised by calling ISANormalize on it. A subset of eset is selected that corresponds to the features included in modules.
- **plot**: Character constant, specifies what to plot. ‘sample’ plots sample scores, ‘features’ plots feature scores. If ‘both’ is given, then the plot is divided into two subplots and both scores are plotted.
- **norm**: Character constant, specifies how to normalize the expression matrix for plotting. It can be of length one or two, the latter for the case when plots are made both for features and samples. Possible values: ‘raw’ uses the raw expression values; ‘feature’ uses featExprs to extract the expression values from the expression set object; ‘sample’ uses sampExprs; ‘default’ means ‘feature’ for sample plots and ‘sample’ for feature plots.

Examples

data(ALLModulesSmall)
G <- overlap(ALLModulesSmall, algorithm="drl", edge.limit=0.3)
if (interactive()) {
  overlapPlot(G)
}

References

profilePlot

background Logical scalar, whether to plot the features/samples that are not in the module.
col Color of lines corresponding to the background features/samples.
col.mod Color of the lines corresponding to the features/samples included in the module.
type Type of the plot, for the background features/samples. It is passed to `plot`.
type.mod Type of the plot, for the features/samples included in the module. It is passed to `plot`.
mean Logical scalar, whether to plot the mean expression for each feature/sample, separately for the samples/features that are in the module and the ones that are not.
meancol Color of the line for the mean expression values, background.
meancol.mod Color of the line for the mean expression values, module.
xlabs Character vector of length one or two. The labels of the horizontal axes of the plot, the second value is used if both the feature and the sample plots are drawn.
ylab Character vector of length one. The label of the vertical axes.
... Additional graphical arguments. They are passed to the `lines` function that creates the lines of the plot.

Details

plot="both" uses the `mfrow` graphical parameter to create the two subplots. This does not work properly if you already have subplots.

Value

None. (Well, NULL, invisibly.)

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

The similar `parallelCoordinates` function in the biclust package.

Examples

data(ALLModulesSmall)
library(ALL)
data(ALL)
if (interactive()) {
  profilePlot(ALLModulesSmall, 2, ALL, plot="samples")
}
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