Package ‘FGNet’

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Type Package

Title Functional Gene Networks derived from biological enrichment analyses

Description Build and visualize functional gene and term networks from clustering of enrichment analyses in multiple annotation spaces. The package includes a graphical user interface (GUI) and functions to perform the functional enrichment analysis through DAVID, GeneTerm Linker, gage (GSEA) and topGO.

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### FGNet-package

Functional gene networks derived from biological enrichment analyses

#### Description

Build and visualize functional gene and term networks from clustering of enrichment analyses in multiple annotation spaces. The package includes a graphical user interface (GUI) and functions to perform the functional enrichment analysis through DAVID, GeneTerm Linker, gage (GSEA) and topGO.

#### Details

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#### Author(s)


If you have any issue, you can contact us at: <jrivas at usal.es>

#### References


See Also

FGNet_GUI() provides a Graphical User Interface (GUI) to most of the functionalities of the package: Performing a Functional Enrichment Analysis (FEA) of a list of genes, and analyzing it through the functional networks.

1. The Functional Enrichment Analysis can be performed through several tools:

   - DAVID [1]: fea_david() (Requires internet connection)
   - GeneTerm Linker [2]: fea_gtLinker() & fea_gtLinker_getResults() (Requires internet connection)
   - topGO [3]: fea_topGO() (Only supports GO. For offline use requires having installed the required database packages)
   - GAGE [4]: fea_gage() (GSEA analysis. For offline use requires gene sets or installed database packages)

   There are also a few functions to import the results from a previous FEA analysis: format_david(), format_results() and readGeneTermSets().

2. FGNet_report(): automatically generates a report with the default network options. It includes the following steps, which can be executed individually to personalize or explore the networks:

   1. fea2incidMat(): Transforms the FEA output into incidence matrices. This function determines whether the network will be gene- or term-based.
   2. functionalNetwork(): Generates and plots the functional networks. These networks can be further explored by analyzeNetwork() and clustersDistance().

   Other auxiliary functions: getTerms(), keywordsTerm(), plotGoAncestors(), plotKegg().

For more info see the package tutorial: vignette("FGNet-vignette")

Examples

```r
## Not run:
# GUI:
FGNet_GUI()

# 1. FEA:

library(org.Sc.sgd.db)
geneLabels <- unlist(as.list(org.Sc.sgdGENENAME)[geneList])

# Optional: Gene expression
geneExpr <- setNames(c(rep(1,10),rep(-1,5)), geneLabels)
```
analyzeNetwork

Analyze Functional Network

Description

Analyzes the degree and betweenness of the genes in the functional network.

Usage

analyzeNetwork(incidMatrices, fNw = NULL, plotOutput = TRUE, colors = NULL)

Arguments

- incidMatrices: list or matrix. Output from fea2incidMat or the equivalent incidence matrices.
- fNw: list. Return from functionalNetwork to avoid recalculating.
- plotOutput: logical. Wether to plot the degree and betweenness boxplots.
- colors: vector. Colors for the metagroups

Value

List:

- degree, betweenness: Degree and Betweenness of the nodes in the global network (commonClusters) and within each cluster/metagroup (subsets of commonGtSets network). The degree is given as percentage, normalized based on the total number of nodes of the network. i.e. a value of 90 in a network of 10 nodes, would mean the actual degree of the node is 9: it is conneded to 9 nodes (90% of 10)).
- transitivity: Transitivity of the networks.
- betweennessMatrix: Betweenness of each node in each cluster.
hubsList: Nodes selected as potential hubs in the global network and within each cluster/metagroup (nodes with betweenness over 75% in the given network/subnetwork).

• intraHubsCount: Number of times each node was selected as potential intra-cluster hub.

See Also

Overview of the package: FGNet
Package tutorial: vignette("FGNet-vignette")

Examples

```r
# Previous Steps
jobID <- 3907019
results <- fea_gtLinker_getResults(jobID)
incidMat <- fea2incidMat(results, filterAttribute="Silhouette Width")

# Plot node degree and betweenness
analyzeNetwork(incidMat)

# Get stats without plotting
nwStats <- analyzeNetwork(incidMat, plotOutput=FALSE)
names(nwStats)
 nwStats$hubsTable
```

---

clustersDistance

**Plots distances between metagroups.**

Description

Plots the distances between metagroups taking into account the number of common genes.

Usage

```r
clustersDistance(incidenceMatices, mgCols = NULL, clustMethod="average")
```

Arguments

- `incidenceMatices`: Object returned by `fea2incidMat()`.
- `mgCols`: Colors for the metagroups.
- `clustMethod`: Clustering method. Character string (i.e. "single", "complete", "average") for function `hclust()` (argument `method`).

Value

Plot and distance matrix.

See Also

Full description of the package: FGNet
Examples

```r
results <- fea_gtLinker_getResults(jobID=1963186, jobName="gtLinker_example")
incidMatrices <- fea2incidMat(results)
clustersDistance(incidMatrices)
```

**Description**

Transforms the Functional Enrichment Analysis (FEA) results into cluster-gene incidence matrices.

**Usage**

```r
fea2incidMat(feaResults, key = "Genes", sepChar = NULL, clusterColumn = NULL,
filterAttribute = NULL, filterOperator = "<", filterThreshold = 0,
removeFilteredGtl = NULL)
```

**Arguments**

- `feaResults`: list or data.frame/matrix. Output from one of the FEA functions.
- `key`: character. "Genes" or "Terms". To build gene- or term-based networks.
- `sepChar`: character. Character separating genes or terms in the same field. By default: ",," for genes and ";" for terms.
- `clusterColumn`: character. Name of the column that contains the value to group gene-term sets. Only required if it is different than "Cluster" or "Metagroup".
- `filterAttribute`: character or data.frame. Attribute to filter the clusters/metagroups. Filtered clusters/metagroups will not be included in the matrices (and subsequent networks). Its value should be the data.frame column to use for filtering. It can be provided as character (column name) or data.frame (subset of the data.frame with drop=FALSE).
- `filterOperator`: character. Logical operator used for filtering. i.e. ">" (bigger than), "<=" (smaller or equal than), "==" (equal), "!=" (different), "%%" (included in),... The evaluation order is left to right: filterAttribute ">" filterThreshold, will filter out clusters with filter attribute bigger than the threshold.
- `filterThreshold`: numeric. Sets the value to compare to.
- `removeFilteredGtl`: logical. Only used by GeneTerm Linker term network. If FALSE, it includes generic terms filtered by GeneTerm Linker from final metagroups.

**Value**

List:

- `clustersMatrix` or `metagroupsMatrix`: Incidence matrix with the genes or Terms in each cluster or metagroup.
- `gtSetsMatrix`: Incidence matrix with the genes or Terms in each gene-term set
- `filteredOut`: Clusters or metagroups which where filtered out and therefore not included in the incidence matrices. NULL if none.
**See Also**

Next step in the workflow: `functionalNetwork()`

Overview of the package: `FGNet`

Package tutorial: vignette("FGNet-vignette")

**Examples**

```r
jobID <- 3907019
results <- fea_gtLinker_getResults(jobID)
incidMat <- fea2incidMat(results)

# Filtering (threshold)
incidMat$filteredOut <- fea2incidMat(results,
    filterAttribute="Silhouette Width", filterThreshold=0.2)

head(incidMat$metagroupsMatrix)
head(incidMat$gtSetsMatrix)

functionalNetwork(incidMat)

# Term-based network
incidMatTerms <- fea2incidMat(results, key="Terms")
functionalNetwork(incidMatTerms, plotOutput="dynamic")

# Including generic terms filtered by GtLinker from final metagroups:
incidMatTerms <- fea2incidMat(results, key="Terms", removeFilteredGtl=FALSE)
functionalNetwork(incidMatTerms, plotOutput="dynamic", plotType="bipartite")

# Filtering by keyword
keywords <- c("rna")
selectedGroups <- sapply(getTerms(results),
    function(x)
    any(grep(paste("(", paste(keywords, collapse="|") ,")",sep=""), tolower(x)))))

resultsCbind <- results
columns <- results$metagroups
resultsCbind$metagroups <- cbind(results$metagroups, selectedKeywords=as.numeric(selectedGroups))

matSelectedGroups <- fea2incidMat(resultsCbind,
    filterAttribute="selectedKeywords", filterThreshold=1)

functionalNetwork(matSelectedGroups)
getTerms(results)[selectedGroups]
```

**Description**

Performs the functional enrichment analysis and clustering through DAVID [1] (requires internet connection).
Usage

```r
fea_david(geneList, geneIdType = "ENSEMBL_GENE_ID", geneLabels=NULL,
annotations = c("GOTERM_BP_ALL", "GOTERM_MF_ALL", "GOTERM_CC_ALL",
"KEGG_PATHWAY", "INTERPRO"),
email = NULL,
argsWS = c(overlap = 4L, initialSeed = 4L, finalSeed = 4L, linkage = 0.5,
kappa = 35L), jobName = NULL, downloadFile=TRUE)
```

Arguments

geneList character vector. List of genes to analyze.
geneIdType character vector. Type of gene identifier.
Web API: ENSEMBL_GENE_ID, ENTREZ_GENE_ID, GENE_SYMBOL, UNIPROT_ID...
For more, check DAVID's API documentation.
Web Service: run getIdTypes(DAVIDWebService$new(email=...))
geneLabels named character vector. Gene name or label to use in the report/plots instead of
the original gene ID. The vector names should be the gene ID and the content
of the vector the gene label. The resulting geneTermSets table will contain the
original gene ID column (geneIDs) and the label column (Genes).
annotations character vector. Annotation spaces for the functional analysis.
Web API: check DAVID's API documentation.
Web Service: run getAllAnnotationCategoryNames(DAVIDWebService$new(email=...)).
email character. If provided, the query will be performed through DAVID's Web Ser-
vice (recommended). Requires registration (see details).
argsWS named integer vector. Additional arguments for the clustering. Only available
using the web service.
jobName character. Folder name and prefix for the files.
downloadFile logical. If TRUE, the result files are saved in the current directory (required to
generate report).

Details


**NOTE:** Since August 2015, DAVID requires https. This causes errors in some systems. A (hope-
fully) temporary solution requires to install some certificates locally. See RDAVIDWebService help:
[https://support.bioconductor.org/p/70090/#72226](https://support.bioconductor.org/p/70090/#72226)

As an alternative, the web API allows to perform a small query without registering.
Note this option is not available in some systems, and the maximum number of genes is limited to
400. (It can be less depending on the ID types and the length of the resulting URL).
More details and full list of gene ID types and annotations are available at: [http://david.abcc.
ncifcrf.gov/content.jsp?file=DAVID_API.html](http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html).

If the functional annotation and clustering has been performed directly at DAVID's Website ([http://david.abcc.ncifcrf.gov/summary.jsp](http://david.abcc.ncifcrf.gov/summary.jsp)) fea_david() is not required. Instead, provide the file
(or the URL of the file) containing the results of the analysis to `format_david()`.
Value

**Invisible** list with the following fields:
- `queryArgs` list with the arguments for the query.
- `clusters` data.frame containing the clusters and their information:
  - **Cluster**: Cluster ID.
  - **nGenes**: Number of genes in the cluster.
  - **ClusterEnrichmentScore**: Score for the cluster.
  - **Genes**: Genes in the cluster.
  - **Terms**: Terms in the cluster.
  - **keyWordsTerm**: Term is the most representative of the terms in the cluster based on keywords.
- `geneTermSets` data.frame containing the gene-term sets that support each cluster:
  - **Cluster**: Number (id) of the cluster the gene-term set belongs to.
  - **ClusterEnrichmentScore**: Score for the cluster. Same value for all terms in each cluster.
  - **Category**: Type of annotation of the term (i.e. GO, Kegg...)
  - **Terms**: Term in the gene-term set.
  - **Genes**: Genes in the gene-term set.
  - **GenesIDs**: In case GeneLabels was provided, original gene ID.
  - **Other stats**: Count, PValue, List.Total, Pop.Hits, Pop.Total, Fold.Enrichment, Bonferroni, Benjamini, FDR.
- `fileName` .txt file with the formatted FEA results.

Warning

The web service and the API have different default arguments. To obtain the same results with both methods use:

```
API_defaults <- c(overlap=3L, initialSeed=3L, finalSeed=3L, linkage=0.5, kappa=50L)
fea_david(genesYeast, email="example@email.com", argsWS=API_defaults)
```

References


See Also

Other FEA tools:
- `fea_gtLinker()` & `fea_gtLinker_getResults()` (Requires internet connection)
- `fea_gage()`
- `fea_topGO()`
  To import results from a previous/external FEA analysis: `format_david()`, `format_results()` and `readGeneTermSets()`.

Overview of the package: FGNet

Package tutorial: vignette("FGNet-vignette")
# Load/format gene list:

library(org.Sc.sgd.db)
geneLabels <- unlist(as.list(org.Sc.sgdGENENAME)[geneList])
geneExpr <- setNames(c(rep(1,10),rep(-1,5)), geneLabels)

## Not run:
# DAVID
results_David <- fea_david(geneList, geneLabels=geneLabels, email="example@email.com")

# Available IDs and annotations:
getIdTypes(DAVIDWebService$new(email="example@email.com"))
getAllAnnotationCategoryNames(DAVIDWebService$new(email="example@email.com"))

results <- fea_david(geneList, geneIdType="ENSEMBL_GENE_ID", annotations="GOTERM_BP_ALL", email="example@email.com", jobName="yeastDavid")

# To continue the workflow... (see help for further details)
FGNet_report(results, geneExpr=geneExpr)
incidMat <- fea2incidMat(results)
functionalNetwork(incidMat, geneExpr=geneExpr)

## End(Not run)

---

**fea_gage**  
**FEA - GAGE**

## Description

Performs the functional enrichment analysis and clustering through GAGE [1] (GSEA).

### Usage

```r
fea_gage(eset, refSamples, compSamples, geneIdType, geneLabels=NULL, organism = "Hs", annotations = c("GO_BP", "GO_MF", "GO_CC", "KEGG", "REACTOME"), geneSets = NULL, sameDirection = FALSE, onlyEssentialTerms = TRUE, compareType = "as.group", jobName = NULL, ...)
```
Arguments

- **eset**: expressionSet or expression matrix.
- **refSamples**: numeric. Index of the samples to use as reference (control).
- **compSamples**: numeric. Index of the samples to analyze.
- **geneIdType**: character. Type of gene identifier should be the same as the one provided in the geneSets, or available in the organism package.
- **geneLabels**: named character vector. Gene name or label to use in the report/plots instead of the original gene ID. The vector names should be the gene ID and the content of the vector the gene label. The resulting geneTermSets table will contain the original gene ID column (geneIDs) and the label column (Genes).
- **organism**: two letter code for the organism. See: data(organisms);organisms
- **annotations**: character vector. Annotation spaces to select from the provided geneSets. Set to NULL to use the geneSets as is (i.e. geneSets not split/named by annotation)
- **geneSets**: geneSets. If NULL geneSets are calculated automatically based on the organism, gene ID and annotations. The geneSets can also be provided from a previous execution or loaded from a .gmt file. i.e.: readList("c2.cp.v4.0.symbols.gmt")
- **sameDirection**: logical. Should all the genes in the geneSet be altered in the same direction (up/down)?
- **onlyEssentialTerms**: logical. Wether to simplify the results and keep only the essential terms in the clusters.
- **compareType**: character: 'as.group', 'unpaired', 'longroup'... See GAGE for details.
- **jobName**: character. Folder name and prefix for the files.
- ... other arguments to pass to GAGE.

Value

- **Invisible** list with the following fields:
- **queryArgs**: list with the arguments for the query.
- **clusters**: data.frame containing the clusters and their information:
  - Cluster: Cluster ID.
  - nGenes: Number of genes in the cluster.
  - dir: Direction in which the term/pathway is altered (Up/Down).
  - Genes: Genes in the cluster.
  - Terms: Terms in the cluster.
- **geneTermSets**: data.frame containing the gene-term sets that support each cluster.
  - Cluster: Number (id) of the cluster the gene-term set belongs to.
  - essentialSet: Logical. Is the pathway selected as essential?
  - dir: Direction in which the term/pathway is altered (Up/Down).
  - Terms: Term in the gene-term set.
  - Genes: Genes in the gene-term set.
  - GenesIDs: In case GeneLabels was provided, original gene ID.
  - Other stats provided by GAGE: p.geomean, stat.mean, p.val, q.val, set.size
- **fileName**: .txt file with the FEA results. genesFC: Fold change.
Background and Purpose

The FEA package is provided to make gene set enrichment analysis more accessible, particularly for non-computer-literate users. It can be used in conjunction with a variety of other packages for data input, output, and visualisation. To learn about the main features and different methods for FEA, consult the package vignette (FGEF vignette), which accompanies the package.

Examples

## Not run:
# Load expressionSet:
library(gage)
data(gse16873)

# Load gene labels?
library(org.Hs.eg.db)
geneSymbols <- select(org.Hs.eg.db,columns="SYMBOL",keytype="ENTREZID",
    keys=rownames(gse16873))
head(geneSymbols)
table(table(geneSymbols$ENTREZID)) # All need to be unique identifiers

geneLabels <- geneSymbols$SYMBOL
names(geneLabels) <- geneSymbols$ENTREZID
head(geneLabels)

# FEA:
results <- fea_gage(eset=gse16873,
    refSamples=grep("HN",colnames(gse16873), ignore.case =T),
    compSamples=grep("DCIS",colnames(gse16873), ignore.case=T),
    geneIdType="ENTREZID", geneLabels=geneLabels, organism="Hs",
    annotations="REACTOME")

# To continue the workflow... (see help for further details)
FGNet_report(results)
incidMat <- fea2incidMat(results)
functionalNetwork(incidMat)

## End(Not run)
Description

Performs the functional enrichment analysis and clustering through Gene-Term Linker [1] (requires internet connection).

Since Gene-Term Linker takes a while to analyze the gene list, the process has been split into two steps:

1. fea_gtLinker(): Submits the query
2. fea_gtLinker_getResults(): Retrieves the results of the analysis. It might take a few minutes for the results to become available.

Usage

```
fea_gtLinker(geneList, organism = "Hs",
            annotations = c("GO_Biological_Process", "GO_Molecular_Function",
                             "GO_Cellular_Component", "KEGG_Pathways", "InterPro_Motifs"),
            minSupport = 4, serverWS = "http://gtlinker.cnb.csic.es:8182")
```

```
fea_gtLinker_getResults(jobID = NULL, organism = NULL, jobName = NULL,
                        alreadyDownloaded = FALSE, keepTrying = FALSE,
                        serverWeb = "http://gtlinker.cnb.csic.es",
                        serverWS = "http://gtlinker.cnb.csic.es:8182")
```

Arguments

- **geneList** character vector. List of genes to analyze.
- **minSupport** numeric. Minimum number of genes per group.
- **organism** character. "Hs" (Homo sapiens) or "Sc" (Saccharomyces cerevisiae).
- **jobID** numeric. ID of the job/analysis in GeneTerm Linker.
- **jobName** character. Folder name and prefix for the files.
- **alreadyDownloaded** logical. If the files have already been downloaded, these will be read instead of downloaded again.
- **keepTrying** logical. If true, if the job has not finished, it will keep trying to get the results every few seconds.
- **serverWeb** character. GeneTerm Linker web server. It should match the web service or web address in which the analysis was performed. Available mirrors: "http://gtlinker.cnb.csic.es"
Value

`fea_gtLinker()` returns the jobID of the analysis
`fea_gtLinker_getResults()` returns an invisible list with the following fields:

- `queryArgs` list with the arguments for the query.
- `metagroups` data.frame containing the metagroups and their information:
  - Metagroup: Metagroup ID.
  - Size: Number of gene-term sets supporting the metagroup.
  - Diameter: Maximum Cosine distance within the GeneTerm-sets of each metagroup (ranges from 0 to 1).
  - Similarity: 1 - average Cosine distance within the GeneTerm-sets of each metagroup (ranges from 0 to 1). Distance and similarity calculations are done based on the genes present in the metagroups.
  - Silhouette Width: Measures the compactness and proximity of multiple groups (ranges from 1 to -1). Metagroups with negative Silhouette Width usually include diverse annotations and genes with low functional coherence.
  - Genes: Genes in the metagroup.
  - nGenes: Number of genes in the metagroup.
  - nref_list: Number of annotated genes in the reference list.
  - pValue: Adjusted p-value.
  - Terms: Non-generic terms in the metagroup.
- `geneTermSets` data.frame containing the gene-term sets that support each metagroup.
  - Metagroup: Id of the metagroup the gene-term set belongs to.
  - Genes: Genes in the gene-term set.
  - nGenes: Number of annotated genes in the input list. In brackets: Total number of genes in the input list.
  - nref_list: Number of annotated genes in the reference list. In brackets: Total number of genes in the reference list.
  - pValue: Adjusted p-value.
  - Terms: Terms in the gene-term set.
- `fileName` .txt file with the formatted FEA results.

References


See Also

Other FEA tools:

- `fea_david()` (Requires internet connection)
- `fea_gage()`
To import results from a previous/external FEA analysis: `format_david()`, `format_results()` and `readGeneTermSets()`. 

Overview of the package: `FGNet`

Package tutorial: `vignette("FGNet-vignette")`

### Execute FEA:
```r
genesYeast <- c("ADA2", "APC1", "APC11", "APC2", "APC4", "APC5", "APC9", 
"CDC16", "CDC23", "CDC26", "CDC27", "CFT1", "CFT2", "DCP1", "DOC1", "FIP1", 
"GCN5", "GLC7", "HFI1", "KEM1", "LSM1", "LSM2", "LSM3", "LSM4", "LSM5", 
"LSM6", "LSM7", "LSM8", "MPE1", "NGG1", "PAP1", "PAT1", "PF52", "PTA1", 
"PTI1", "REF2", "RNA14", "RPN1", "RPN10", "RPN11", "RPN13", "RPN2", "RPN3", 
"RPN5", "RPN6", "RPT1", "RPT3", "RPT6", "SGF11", "SGF29", "SGF73", 
"SPT20", "SPT3", "SPT7", "SPT8", "TRA1", "YSH1", "YTH1")
# Optional expression (1=UP, -1=DW):
genesYeastExpr <- setNames(c(rep(1,29), rep(-1,30)), genesYeast)
```

```r
# Submit query
jobID <- fea_gtLinker(geneList=genesYeast,organism="Sc")
jobID
```

### Get results from FEA:
```r
jobID <- 3907019 # job ID of the query
results <- fea_gtLinker_getResults(jobID=jobID)
```

```r
# To continue the workflow... (see help for further details)
incidMat <- fea2incidMat(results)
functionalNetwork(incidMat)
```

```r
# Or full report
## Not run:
FGNet_report(results, geneExpr=genesYeastExpr)
## End(Not run)
```

---

### fea_topGO

**FEA - topGO**

#### Description
Performs the functional enrichment analysis through topGO [1].

#### Usage
```r
fea_topGO(geneList, geneIdType = "ENSEMBL", geneLabels=NULL, organism = "Hs", 
annotations = c("GO_BP", "GO_MF", "GO_CC"), evidence=NULL, 
genesUniverse = NULL, refPackage = NULL,
```
geneID2GO = NULL, nodeSize = 5, pValThr = 0.01, testStat = NULL, jobName = NULL)

Arguments

geneList character vector. List of genes to analyze.
geneIdType character. Type of gene identifier should be available for the organism package.
geneLabels named character vector. Gene name or label to use in the report/plots instead of the original gene ID. The vector names should be the gene ID and the content of the vector the gene label. The resulting geneTermSets table will contain the original gene ID column (geneIDs) and the label column (Genes).
organism two letter code for the organism. See: data(organisms);organisms
annotations character vector. Annotation spaces for the functional analysis. Accepted values: "GO_BP", "GO_MF", "GO_CC".
evidence character vector. Required evidence code for GO annotations. If NULL no filtering is done (all annotations are used). For full list, see the organism "EVIDENCE" keys: i.e keys(org.Hs.eg.db, keytype="EVIDENCE"). For non-comprehensive code description: data(GOEvidenceCodes).
genesUniverse character vector. List of genes used for background (i.e. all genes available in the chip).
refPackage character. Name of the package to use for calculating the genes universe. A Chip package is recommended. If NULL the genes universe is set as all the genes available in the organism package.
geneID2GO GO gene sets. If NULL it is calculated automatically.
nodeSize numeric. Minimum size of GO terms. TopGo authors recommend 5-10 for more stable results, 1 for no prune.
pValThr numeric. P-value threshold.
testStat character. Folder name and prefix for the files.

Value

Invisible list with the following fields:
queryArgs list with the arguments for the query.
clusters Empty list. Only for compatibility.
geneTermSets data.frame containing the gene-term sets.
  • Ont: Ontology to which the term belongs (BP, MF or CC)
  • Terms: Term in the gene-term set.
  • Genes: Genes in the gene-term set.
  • GenesIDs: In case GeneLabels was provided, original gene ID.
  • Other stats provided by topGO: Annotated, Significant, Expected, classic.

fileName .txt file with the formatted FEA results.

References

See Also
Other FEA tools:

- `fea_david()` (Requires internet connection)
- `fea_gtLinker()` & `fea_gtLinker_getResults()` (Requires internet connection)
- `fea_gage()`

To import results from a previous/external FEA analysis: `format_david()`, `format_results()` and `readGeneTermSets()`.

Overview of the package: **FGNet**
Package tutorial: `vignette("FGNet-vignette")`

Examples

```r
## Not run:
# Load/format gene list:
geneList <- c("YBL084C", "YDL008W", "YDR118W", "YDR301W", "YDR448W", "YFR036W",  
"YGL240W", "YHR166C", "YKL022C", "YLR102C", "YLR115W", "YLR127C", "YNL172W",  
"YOL149W", "YOR249C")
library(org.Sc.sgd.db)
geneLabels <- unlist(as.list(org.Sc.sgdGENENAME)[geneList])
geneExpr <- setNames(c(rep(1,10),rep(-1,5)), geneLabels)

# FEA (using directly the gene names):
results <- fea_topGO(geneLabels, geneIdType="GENENAME", organism="Sc")

# FEA (using the gene ID, and replacing it by the label after the FEA):
results <- fea_topGO(geneList, geneIdType="ENSEMBL",  
geneLabels=geneLabels, organism="Sc")

# To continue the workflow... (see help for further details)
FGNet_report(results, geneExpr=geneExpr)
inciMat <- fea2inidMat(results, geneExpr=geneExpr)
functionalNetwork(inciMat)

## End(Not run)
```

### FGNet_GUI

**FGNet graphical user interface**

**Description**
Provides a graphical user interface (GUI) to most FGNet functionalities.

**Usage**

```r
FGNet_GUI(geneList = NULL)
```
Arguments

geneList vector. If provided, assigns the value to the genes field. It can be a character vector containing the gene list, or a named numeric vector with the gene expression.

Details

To generate the functional network, first execute or import the Functional Analysis of the gene list with one of the tools in Tab "1 - FEA", then generate the network or the report in Tab "2 - Network".

Value

Opens the GUI. No value is returned. The results of the analyses will be saved in the current working directory.

Note

Available for Windows and Linux. The current version of the GUI is not available for Mac OS X Snow Leopard.

See Also

Overview of the package: FGNet
Package tutorial: vignette("FGNet-vignette")

Examples

## Not run:

FGNet_GUI()

# To directly input a gene list (i.e. from a previous analysis):
# Optional gene expression
geneExpr <- setNames(c(rep(1,10),rep(-1,5)), genelist)

FGNet_GUI(geneExpr)

## End(Not run)
**FGNet_report**

Usage

```r
FGNet_report(feaResults, geneExpr = NULL, plotExpression = "border",
onlyGoLeaves = TRUE, plotGoTree = TRUE, plotKeggPw = TRUE,
filterAttribute = NULL, filterOperator = NULL, filterThreshold = NULL)
```

Arguments

- **feaResults**: list or data.frame/matrix. Output from one of the FEA functions.
- **geneExpr**: numeric. Named vector with the relative expression value of the gene (node). 0 is taken as reference, positive values will be plotted red, negative values green.
- **plotExpression**: character. Determines the way to plot the expression: "border" adds a red or green border to the node, "fill" colors the whole with the expression color instead of the metagroup color.
- **onlyGoLeaves**: logical. If TRUE only terminal GO terms (leaves in the ontology tree) will be included in the cluster list.
- **plotGoTree**: logical. If TRUE plots containing the terms in their position within the GO ontology (tree) will be generated.
- **plotKeggPw**: logical. If TRUE local plot for KEGG pathway terms will be generated. If FALSE KEGG pathway terms will link to KEGG website. In the current version local plots are only available for ENTREZ gene IDs.
- **filterAttribute**: character or data.frame. Attribute to filter the clusters/metagroups. Filtered clusters/metagroups will not be included in the matrices (and subsequent networks). Its value should be the data.frame column to use for filtering. It can be provided as character (column name) or data.frame (subset of the data.frame with drop=FALSE).
- **filterOperator**: character. Logical operator used for filtering. i.e. ">" (bigger than), "<=" (smaller or equal than), "==" (equal), "!=" (different), "%%" (included in),... The evaluation order is left to right: filterAttribute ">" filterThreshold, will filter out clusters with filter attribute bigger than the threshold.
- **filterThreshold**: numeric. Sets the value to compare to.

Value

Generates the HTML report in the current directory.

Warning

Reactome ID change depending on the database version. Links to reactome website are created to ease the analysis, but in case the version used for the enrichment and the website’s do not match, broken links or misleading links might appear.

See Also

Functional enrichment analysis functions:

- `fea_david()` (Requires internet connection)
- `fea_gtLinker()` & `fea_gtLinker_getResults()` (Requires internet connection)
- `fea_gage()`
• **feareport**
  To import results from a previous/external FEA analysis: format_david(), format_results() and readGeneTermSets().

Overview of the package: *FGNet*

Package tutorial: vignette("FGNet-vignette")

Examples

## Not run:

### # Report with different tools:

```
# DAVID & TopGO
geneExpr <- setNames(c(rep(1,10),rep(-1,5)), geneList)
library(org.Sc.sgd.db)
geneLabels <- unlist(as.list(org.Sc.sgdGENENAME)[geneList])
names(geneExpr) <- geneLabels[1]
# DAVID
results_David <- fea_david(geneList, geneLabels, email="example@email.com")
FGNet_report(results_David, geneExpr)
# TopGO
results_topGO <- fea_topGO(geneList, geneIdType="ENSEMBL", geneLabels=geneLabels, organism="Sc")
FGNet_report(results_topGO, geneExpr)
```

### # Gage

```
library(gage); data(gse16873)
results_gage <- fea_gage(eset=gse16873, refSamples=grep("HN", colnames(gse16873), ignore.case=T),
compSamples=grep("DCIS", colnames(gse16873), ignore.case=T),
geneIdType="ENTREZID", organism="Hs", annotations="REACTOME")
FGNet_report(results_gage)
```

### # Gene-Term Linker:

```
# Execute new query:
# Optional expression (1=UP, -1=DW):
genesYeastExpr <- setNames(c(rep(1,29), rep(-1,30)), genesYeast)
```
format_david

jobID <- fea_gtLinker(geneList=genesYeast,organism="Sc")

# Load existing query:
jobID <- 3907019

results_gt linker <- fea_gtLinker_getResults(jobID=jobID)
FGNet_report(results_gtLinker, geneExpr=genesYeastExpr)

## End(Not run)

---

format_david Format DAVID output

Description

Format DAVID 'functional annotation and clustering' output to use with FGNet.

Usage

format_david(fileName, jobName = NULL, geneLabels = NULL, moveFile = FALSE, downloadFile=TRUE)

Arguments

- **fileName**: character. URL or local file with the results of a DAVID analysis, for example, performed at DAVID's Website (http://david.abcc.ncifcrf.gov/summary.jsp). In case of local file, it should be the absolute path to the .txt file (whole location from root to the file including file name: "C:\Documents\23424203.txt", "/home/user/2342342.txt").

- **jobName**: character. Folder name and prefix for the formatted files.

- **geneLabels**: named character vector. Gene name or label to use in the report/plots instead of the original gene ID. The vector names should be the gene ID and the content of the vector the gene label. The resulting geneTermSets table will contain the original gene ID column (geneIDs) and the label column (Genes).

- **moveFile**: logical. If TRUE the original file is moved to the new location. If FALSE, the file is copied.

- **downloadFile**: logical. If TRUE, the result files are saved in the current directory (required to generate report).

Value

List. Same as fea_david.

References

format_results

Description

Format the functional analysis results from external tools to use with FGNet.

Usage

format_results(fileName, newFileName = NULL, clusterCol = NULL,
               geneCol = NULL, geneSep = NULL, termDescCol = NULL, termIDCol = NULL,
               termCatCol = NULL, termCat = NULL, termSep = NULL,
               tool = "Imported text file", simplifyGage = TRUE, ...)

Arguments

fileName character. File name with the FEA results.
newFileName character. Name for the formatted files.
clusterCol character. Name of the column to use for clustering.
geneCol character. Name of the column with the genes.
geneSep character. Character separating different genes in the same field (i.e. ".", ",", ...) 
termDescCol character. Name of the column with the terms description.
termIDCol character. Name of the column with the terms ID.
termCatCol character. Name of the column with the terms type/category.

Examples

# Select file:
txtFile <- "http://david.abcc.ncifcrf.gov/data/download/901234901248.txt"

# Read:
results <- format_david(txtFile, jobName="DavidAnalysis")

# To continue the workflow... (see help for further details)
getTerms(results)
incidMat <- fea2incidMat(results)
functionalNetwork(incidMat)

?FGNet_report
**functionalNetwork**

*Creates and plots the functional gene network.*

**Description**

Plots the functional networks.

The default network links genes to genes, or terms to terms. The bipartite network links genes or terms to their clusters.

**Usage**

```r
functionalNetwork(incidMatrices, plotType = c("default", "bipartite")[[1]],
plotOutput = "static", plotTitle = "Functional Network",
plotTitleSub = NULL, legendPrefix = NULL, legendText = NULL,
geneExpr = NULL, plotExpression = c("border", "fill"),
vExprColors = c(neg="#008000", zero="white", pos="#FF2020"),
vSize = 12, vLabelCex = 2/3, vLayout = NULL, keepColors = TRUE,
bgTransparency = 0.4, eColor = "#323232", eWidth=NULL, weighted = FALSE,
keepAllNodes = FALSE, plotAllMg = FALSE)
```

**termCat** character. Name of the annotation type if it is common to all gene-term sets. Provide either termCatCol or termCat, not both.

**termSep** character. Character separating different terms in the same field (i.e. ",", ";", ...) for functionalNetwork.

**tool** character. Tool used for the FEA (row name from data(FEA_tools)).

**simplifyGage** logical. For internal use, only for GAGE. Determines whether to keep non essential terms in the final clusters.

**Value**

Saves the formatted file and returns an invisible list with the appropriate format to use with `FGNet_report()` and `fea2incidMat()` (fields "clusters", "geneTermSets" and "fileName").

**See Also**

Overview of the package: **FGNet**

Package tutorial: vignette("FGNet-vignette")

**Examples**

```r
## Not run:

results <- format_results("/home/user/feaResults.txt", clusterCol="Cluster",
geneCol="Genes", termDescCol="Terms", sep="\t")

## End(Not run)
```

---

`functionalNetwork` is used to create and plot the functional gene network. It takes an `incidMatrices` (input) and generates a network based on the provided parameters. The network can be either of type “default” or “bipartite”. Additional parameters are used to customize the appearance of the network, such as the title, legend, gene expression colors, and layout. The function returns a list that can be used for further analysis with `FGNet_report()` and `fea2incidMat()`. Example usage is shown where the results from a formatted file are passed to `functionalNetwork` to create a network with specified parameters.
Arguments

incidMatrices list or matrix. Raw output (list) from fea2incidMat: a list with slots: "gtSetsMatrix", "filteredOut" and either "metagroupsMatrix" or "clustersMatrix". If only a matrix is provided, it will be assumed to be the clusters matrix, and all nodes will be connected to every other node in the metagroup.

plotType "default" or "bipartite".
Default network: Nodes are either genes or terms. Edges join nodes in common gene-term sets. Background and node color represent cluster/metagroup. White nodes are in several clusters/metagroups.
Bipartite network: Nodes are genes or terms (circles) and their clusters (squares). By default it keeps only the genes or terms in more than one cluster or metagroup, which represents a simplified version of the functional network. Node shape is only available in the "static" output.

plotOutput "static", "dynamic" or "none". "static" will generate a standard R plot. "dynamic" will produce an interactive tkplot (metagroups background cannot be drawn). "none" will not plot the network.

plotTitle character. Title to show on the plot.

plotTitleSub character. Text to show at the bottom of the plot (sub-title).

legendPrefix character. Label to show next to the cluster/metagroup id in the legend. In the bipartite network the legends replaces the cluster node label.

legendText character. Description of each cluster (shown as the legend). If FALSE, legend is not shown.

geneExpr numeric. Named vector with the relative expression value of the gene (node). 0 is taken as reference, positive values will be plotted red, negative values green.

plotExpression character. Determines the way to plot the expression: "border" adds a red or green border to the node, "fill" colors the whole with the expression color instead of the metagroup color.

vExprColors character. Vector with the colors for expression: first color for negative values, second for zero, and third for positive.

vSize numeric. Vertex size. If named, it allows to set a value for each gene. Name as "default" to set a default value, otherwise the default value is the mean.

vLabelCex numeric. A numerical value giving the amount by which plotting text and symbols should be magnified relative to the default label size.

vLayout 2 x n matrix or character. Where n is the number of nodes in the graph, each column gives the (x, y)-coordinates for the corresponding node. The bipartite network accepts "kk" (Kamada Kawai), "circle", or "sugiyama" (hierarquical).

keepColors logical. If TRUE, it will keep the same colors for all the plots, independently of the filtered groups. Only available if incidMatrices is the raw result from fea2incidMat()．

bgTransparency numeric. Value between 0 and 1 for the transparency of the metagroups background (only default network).

eColor character. Color for the edges.

eWidth numeric. Edge width. Not to plot edges, set eWidth=0 or eColor=NA.

weighted logical. If TRUE, edges width will be based on the number of shared gene-term sets.

keepAllNodes logical. Only used in bipartite network. If FALSE, nodes in only one cluster are not plotted. If TRUE, all nodes in the clusters are shown.
plotAllMg logical. Only used in bipartite network. If FALSE, non-connected clusters are not plotted. If TRUE, all non-filtered clusters are shown.

Value
Plots the functional networks.
An invisible list with the igraph networks and incidence matrices, to collect it assign it to a variable.

See Also
Previous step in the workflow: fea2incidMat()
To see the terms included in each cluster or metagroup: getTerms()
Overview of the package: FGNet
Package tutorial: vignette("FGNet-vignette")

Examples

# Previous steps
# Set gene list:
# Optional gene expression
genesYeastExpr <- setNames(c(rep(1,29), rep(-1,30)), genesYeast) # 1=UP, -1=DW

# FEA:
# jobID <- query_gtLinker(genesYeast, organism = "Sc")
jobID <- 3907019
results <- fea_gtLinker_getResults(jobID)

# Gene-based networks:
incidMat <- fea2incidMat(results, filterAttribute="Silhouette Width")
functionalNetwork(incidMat, geneExpr=genesYeastExpr)
functionalNetwork(incidMat, plotType="bipartite", plotOutput="dynamic", vSize=c(default=10, GLC7=20, PTA1=20))

getTerms(results)

# To modify the layout and plot as static network (with metagroup background)...
library(igraph)
# saveLayout <- tkplot.getcoords(1) # tkp.id (ID of the tkplot window)
# functionalNetwork(incidMat, vLayout=saveLayout, plotType="bipartite")

# Only return the network, without plotting
fNw <- functionalNetwork(incidMat, plotOutput="none")
class(fNw)
names(fNw)
betweenness(fNw$iGraph$commonClusters)
getTerms

getTerms

Get terms in the metagroups/clusters.

Description

Gets the terms in each metagroup/cluster (simplifies the raw output from GeneTermLinker or DAVID).

Usage

getTerms(feaResults, returnValue = "description")

Arguments

feaResults Output returned by any of the fea functions.

returnValue "description" Returns term description, "GO" or "KEGG" returns GO or KEGG term IDs.

Value

List of matrices

Each matrix contains the terms in each metagroup. This matrix contains only the term description. To get the term ID, annotation type, number of genes, or any other information, see the raw results returned by getResults.

See Also

Full description of the package: FGNet

Examples

results <- fea_gtLinker_getResults(jobID=1963186)
getTerms(results)
**keywordsTerm**

Select keyword term

**Description**

Selects a term as the most representative of the terms in the cluster based on keywords.

**Usage**

```r
keywordsTerm(termsDescriptions, nChar = 30)
```

**Arguments**

- `termsDescriptions` 
  List with the terms in each cluster. Output from `getTerms`.
- `nChar` 
  numeric. Maximum number of chars to show in the term. If the selected term is longer, it will be trimmed.

**Value**

Character vector with the term selected for each cluster.

**See Also**

Overview of the package: [FGNet](#)

Package tutorial: vignette("FGNet-vignette")

**Examples**

```
# Previous Steps: FEA
results <- fea_gtLinker_getResults(3907019)

# Select keywords
keywordsTerm(getTerms(results), nChar=100)
```

**organisms**

FGNet data

**Description**

FGNet package includes the following data files: organisms, groupTypes and FEA_tools
**plotGoAncestors**

### Usage
```
data(organisms)
organisms

data(groupTypes)
groupTypes

data(FEA_tools)
FEA_tools

data(GOEvidenceCodes)
GOEvidenceCodes
```

### Value
- **organisms**: Matrix with the supported organisms' name, package and KEGG prefix.
- **groupTypes**: Matrix with the group types supported by FGNet (cluster, metagroup and gene-term set).
- **FEA_tools**: Matrix with info about the FEA tools supported by FGNet.

### Description
Plots the ancestors in the tree ontology for the given GO terms.

### Usage
```
plotGoAncestors(goIds, tColor = NULL, ontology = NULL,
plotOutput = "static", nCharTerm = 50, nSize = NULL, labelCex = NULL,
asp = NULL, fileName = NULL, height = 1000)
```

### Arguments
- **goIds**: character vector. GO IDs of the terms to plot.
- **tColor**: character. Color for the term (i.e. based on expression).
- **ontology**: character. If character determines which ontology to plot ("BP", "MF" or "CC").
- **plotOutput**: "static", "dynamic" or "none". "static" will generate a standard R plot. "dynamic" will produce an interactive tkplot. "none" will not plot the network.
- **nCharTerm**: numeric. Max term size (number of characters). Longer terms will be trimmed.
- **nSize**: numeric. Determines the node size.
- **labelCex**: numeric. Determines the node label size.
- **fileName**: character. If provided, the plot is saved as png with this fileName.
- **asp**: character. If fileName is provided, asp argument for plot.
- **height**: numeric. If fileName is provided, height argument for png().
Value

An invisible list with the nodes identified as leaves (leaves) and the graph (iGraph).

See Also

Overview of the package: FGNet
Package tutorial: vignette("FGNet-vignette")

Examples

plotGoAncestors(c("GO:0000152", "GO:0043234", "GO:0044446", "GO:0043227"))

plotGoAncestors(c("GO:0051603", "GO:0019941", "GO:0051128", "GO:0044265"),
               plotOutput="dynamic")

# From analysis:
txtFile <- paste(file.path(system.file('examples', package='FGNet')),
                 "DAVID_Yeast_raw.txt", sep="/"
results <- format_david(txtFile, jobName="DavidAnalysis")
plotGoAncestors(getTerms(results, returnValue="GO")$"Cluster 7", ontology="MF")

plotKegg

Plot KEGG pathway

Description

Plots KEGG pathway with the given genes.

Usage

plotKegg(keggIDs, geneExpr, geneIDtype = "ENSEMBL",
         colType = c("continuous", "discrete"))

Arguments

keggIDs character vector. KEGG IDS with prefix.
geneExpr named numeric vector. Names should contain the gene ID, the value the gene
          expression. NA is accepted as expression value.
geneIDtype character vector. Type of gene identifier.
colType "continuous" or "discrete" determines the color of the genes.

Details

Color code:

- Green: Negative expression.
- Red: Positive expression.
- Light yellow: Expression zero or very close to zero.
- Blue: Expression not available.
- White: Gene not in list.
Value

The pathway plot is saved in the current working directory.

See Also

Overview of the package: FGNet

Package tutorial: vignette("FGNet-vignette")

Examples

```
# Gene info

genesYeastExpr <- setNames(c(rep(1,29), rep(-1,30)), genesYeast)
```

```
# Gene expression (1=UP, -1=DW)
genesYeastExpr <- setNames(c(rep(1,29), rep(-1,30)), genesYeast)
```

```
# Plot pathway

# Specific pathway
plotKegg("sce04111", genesYeastExpr, geneIDtype="GENENAME")
```

```
# Pathways from analysis:

txtFile <- paste(file.path(system.file("examples", package="FGNet")),
              "DAVID_Yeast_raw.txt", sep="/"")
results <- format_david(txtFile, jobName="DavidAnalysis")
keggTerms <- unlist(getTerms(results, returnValue="KEGG"))

# plotKegg(keggTerms, ... geneIDtype="ENSEMBL")
```
readGeneTermSets

Usage

readGeneTermSets(fileName, tool = NULL, simplifyGage = TRUE)

Arguments

fileName character. File name of the .txt file.
tool character. Tool used for the FEA (row name from data(FEA_tools); FEA_tools)
simplifyGage logical. Only for GAGE: Wether to simplify the results and keep only the essential terms in the clusters.

Value

List formatted in the same way as the fea functions.

See Also

Overview of the package: FGNet
Package tutorial: vignette("FGNet-vignette")
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