

Package ‘CompGO’

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Title An R pipeline for .bed file annotation, comparing GO term enrichment between gene sets and data visualisation

Description This package contains functions to accomplish several tasks. It is able to download full genome databases from UCSC, import .bed files easily, annotate these .bed file regions with genes (plus distance) from aforementioned database dumps, interface with DAVID to create functional annotation and gene ontology enrichment charts based on gene lists (such as those generated from input .bed files) and finally visualise and compare these enrichments using either directed acyclic graphs or scatterplots.

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Depends RDAVIDWebService

Imports rtracklayer, Rgraphviz, ggplot2, GenomicFeatures, TxDb.Mmusculus.UCSC.mm9.knownGene, pcaMethods, reshape2, pathview

biocViews GeneSetEnrichment, MultipleComparison, GO, Visualization

NeedsCompilation no

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| | |
|-------------------|------------------------------------|
| annotateBedFromDb | <i>Annotate .bed file to genes</i> |
|-------------------|------------------------------------|

Description

Wrapper for transcriptsByOverlaps(). Returns a GRanges with the gene and transcript ids associated with the input .bed regions. Sometimes it is necessary to expand the search window a bit, because not all .bed regions directly overlap with a transcription start site, so the 'window' parameter is provided to accomplish this.

Usage

```
annotateBedFromDb(pathToBed = NULL, gRanges = NULL, db = NULL,
  window = 5000)
```

Arguments

| | |
|-----------|--|
| pathToBed | The system path to a .bed file (directory + file name) |
| gRanges | If the user has a .bed file already loaded in R, they can supply it here as a GRanges object rather than re-importing it |
| db | A TxDb object containing the transcripts of the organism (required) |
| window | The window around a .bed region to search for genes, default 5kb |

Value

A GRanges object with corresponding EntrezGene IDs in gene_id column, plus transcript IDs in tx_id

Examples

```
library(TxDb.Mmusculus.UCSC.mm9.knownGene)
txdb = TxDb.Mmusculus.UCSC.mm9.knownGene
data(bed.sample)
range = GRanges(seqnames=bed.sample$chr, IRanges(start=bed.sample$start, end=bed.sample$end))
x = annotateBedFromDb(gRanges = range, db = txdb)
x
```

| | |
|------------|--|
| bed.sample | <i>A sample of 25 rows from a .bed file of mm9 regions</i> |
|------------|--|

Description

25 regions from a .bed file for use in example code, contains regions from mm9

Usage

```
bed.sample
```

Format

A data.frame with 25 obs. of 3 variables: chromosome, start position, end position

| | |
|----------------|--|
| compareZscores | <i>Compare the Z scores of individual GO terms between two input annotation charts</i> |
|----------------|--|

Description

Accepts two fnAnot charts as args, does z score and p value calculations on them and returns a data.frame with important data. A flag, geneInfo, is provided in case the user wants to get information about the intersection and union of genes corresponding to the individual GO terms. Importantly, this function does some implicit thresholding: only terms with a minimum of 'cutoff' genes are compared, and any term present in one list but not the other is discarded.

Usage

```
compareZscores(setA, setB, geneInfo = FALSE, cutoff = 10)
```

Arguments

| | |
|----------|---|
| setA | FunctionalAnnotationChart to compare |
| setB | FunctionalAnnotationChart to compare |
| geneInfo | Whether to add gene intersection and union info to the data.frame |
| cutoff | The minimum number of genes to threshold terms by |

Value

A data.frame with columns: Term, Zscore.A, Zscore.B, ComparedZ, Pvalue (optionally geneUnion, geneIntersect as well, which are comma-separated strings).

Examples

```
data(funChart1)
data(funChart2)
cz = compareZscores(funChart1, funChart2)
str(cz)
cz = compareZscores(funChart1, funChart2, geneInfo = TRUE)
str(cz)
```

| | |
|-----------------|--|
| doZtrans.single | <i>Z transform a single functional annotation chart from DAVID</i> |
|-----------------|--|

Description

Decomposes each GO term in a functional annotation chart (returned from `getFnAnot_genome()`) to its Z-score. These tables can be merged for clustering

Usage

```
doZtrans.single(x, name)
```

Arguments

| | |
|------|--|
| x | The functional annotation chart to apply the transformation to |
| name | (optional) The name to give the Z-score column; if not supplied, name is derived from the input variable |

Value

A data.frame of GO terms and Z-scores

Examples

```
# Load example fnAnot charts from DAVID:
data(funChart1)
zscore = doZtrans.single(funChart1)
str(zscore)
```

| | |
|-------|---|
| gata4 | <i>A .bed file containing genomic ranges sampled from gata4</i> |
|-------|---|

Description

1000 genomic ranges sampled from gata4 binding sites identified in HL-1 cells (mm9)

Usage

```
gata4
```

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

getFnAnot_genome

*Get the functional annotation chart of a gene list using DAVID***Description**

Uploads a gene list to DAVID, then performs a GO enrichment analysis. Requires registration with DAVID first [here](#). Returns a DAVIDFunctionalAnnotationChart object which can be easily coerced into a data.frame. DAVID does some automatic thresholding on results. For Z-score standardisation, we found it useful to get DAVID to return all possible annotations despite non-significant P-values and perform our own thresholding.

Usage

```
getFnAnot_genome(geneList, david = NULL, email = NULL,
  idType = "ENTREZ_GENE_ID", listName = "auto_list", count = 1L,
  PVal = 1, background = NULL, bgIdType = NULL, bgListName = NULL,
  getKEGG = FALSE)
```

Arguments

| | |
|------------|---|
| geneList | Either a list of genes or a GRanges result from annotateBedFromDb to upload and functionally enrich |
| david | An RDAVIDWebService object can be passed to the function so a new one doesn't have to be requested each time |
| email | If david==NULL, an email must be supplied. DAVID requires (free) registration before users may interact with their Webservice API. This can be accomplished online (here), then the registered email supplied here. |
| idType | The type of gene IDs being uploaded (MGI, Entrez,...) |
| listName | The name to give the list when it's uploaded to the Webservice |
| count | Minimum number of genes per GO term |
| PVal | P-value threshold for GO terms |
| background | If you want to perform enrichment against a specific background instead DAVID's default (whole genome), supply it here |
| bgIdType | If the background gene ID type is different from the gene list, enter it here |
| bgListName | If you want to give the background a name, enter it here |
| getKEGG | TRUE if you want to download KEGG pathway information as well as GO |

Value

Returns a DAVIDFunctionalAnnotationChart after generating it by comparing the supplied gene list to the full genome as a background

Examples

```
## not run because registration is required
## visit http://david.abcc.ncifcrf.gov/webservice/register.htm to register
## Not run:
## You can either supply the registered email:
fnAnot = getFnAnot_genome(exp1$gene_id,
```

```

    email = "your.registered@email.com",
    idType="ENTREZ_GENE_ID", listName="My_gene_list-1")
## Or create a DAVIDWebService object with the email:
david = DAVIDWebService$new(email = "your.registered@email.com")
fnAnot = getFnAnot_genome(entrezList, david = david)

## End(Not run)

```

mef2a

A .bed file containing genomic ranges sampled from mef2a

Description

1000 genomic ranges sampled from mef2a binding sites identified in HL-1 cells (mm9)

Usage

mef2a

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

nkx25

A .bed file containing genomic ranges sampled from nkx25

Description

1000 genomic ranges sampled from NKX2-5 binding sites identified in HL-1 cells (mm9)

Usage

nkx25

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

p300

A .bed file containing genomic ranges sampled from p300

Description

1000 genomic ranges sampled from p300 binding sites identified in HL-1 cells (mm9)

Usage

p300

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

PCApplot

Plot PCA given an input list of fnAnot charts

Description

Given a list of functional annotation charts, this function outputs a PCA plot

Usage

PCApplot(input)

Arguments

input A list of functional annotation charts.

| | |
|----------------|---|
| plotDendrogram | <i>Interactive plotting function for groups of GO terms</i> |
|----------------|---|

Description

Given a list of functional annotation charts and optionally an output directory, this function can output dendrograms, PCA analysis plots and a correlation matrix to make large-scale comparisons easy.

Usage

```
plotInteractive(input, outDir = NULL, prefix = NULL, pdf = TRUE)
```

Arguments

| | |
|--------|--|
| input | A list of functional annotation charts. |
| outDir | The directory to save plots to. |
| prefix | The prefix to append to each file, if any. |
| pdf | If true, plots will be pdfs. If false, pngs. |

| | |
|----------------|---|
| plotDendrogram | <i>Plot dendrogram given an input list of fnAnot charts</i> |
|----------------|---|

Description

Given a list of functional annotation charts, this function outputs a dendrogram

Usage

```
plotDendrogram(input)
```

Arguments

| | |
|-------|---|
| input | A list of functional annotation charts. |
|-------|---|

| | |
|--------------|--|
| plotPairwise | <i>Generates a scatterplot of two sets of GO terms based on DAVID P-values</i> |
|--------------|--|

Description

Generates a $-\log_{10}$ scatterplot of two sets of GO terms by p-value or corrected p-value with linear fit and correlation. Also includes a Jaccard metric for gene overlap within each GO term. Useful as an overall metric of gene list similarity. NOTE: The plotZScores function is more statistically sound, you should use that instead of this.

Usage

```
plotPairwise(setA, setB, cutoff = NULL, useRawPvals = FALSE,
             plotNA = TRUE, model = "lm", ontology = NULL)
```

Arguments

| | |
|-------------|--|
| setA | DAVIDFunctionalAnnotationChart object to compare |
| setB | DAVIDFunctionalAnnotationChart object to compare |
| cutoff | The p-value or adjusted p-value to use as a cutoff |
| useRawPvals | If false, uses adjusted p-values, otherwise uses the raw ones |
| plotNA | If true, any GO term present in only one list is considered to have a p-value of 1 in the other; otherwise, it is simply removed |
| model | The model to use when plotting linear fit, default 'lm' |
| ontology | If a specific ontology (MF, BP, CC) is wanted rather than all terms, supply it here as a string |

Examples

```
data(funChart1)
data(funChart2)
plotPairwise(funChart1, funChart2)
```

| | |
|---------------|--|
| plotTwoGODags | <i>Plots a directed acyclic graph of GO terms from two different sources</i> |
|---------------|--|

Description

Plots a directed acyclic graph of GO terms from two different sources, using colour to show intersection and difference. This is useful to see the specific functional differences between gene lists, complementing the overall metric of gene list similarity

Usage

```
plotTwoGODags(setA, setB, ont = "BP", cutoff = 0.01, maxLabel = NULL,
              fullNames = TRUE, Pvalues = TRUE)
```

Arguments

| | |
|-----------|---|
| setA | A DAVIDFunctionalAnnotationChart object |
| setB | A DAVIDFunctionalAnnotationChart object |
| ont | The ontology to use, one of BP, MF and CC |
| maxLabel | Maximum length of GO term to print |
| cutoff | The PValue cutoff to use |
| fullNames | Whether to print the full GO term label or just the GO id |
| Pvalues | Whether to print P-values alongside each label |

References

Fresno, C. and Fernandes, E. (2013) RDAVIDWebService: An R Package for retrieving data from DAVID into R objects using Web Services API. <http://david.abcc.ncifcrf.gov/>

Examples

```
data(funChart1)
data(funChart2)
plotTwoGODags(funChart1, funChart2)
```

| | |
|----------------|--|
| plotZRankedDAG | <i>Plot a directed acyclic graph (DAG) based on the corrected Pvalues generated from comparing two sets of Z scores.</i> |
|----------------|--|

Description

This function accepts two functional annotation charts as input, performs a comparison on them using compareZscores() and plots a DAG based on the results. The saturation of each node is computed based on the Pvalue, such that the more significant values are darker in colour.

Usage

```
plotZRankedDAG(setA, setB, ont = "BP", n = 100, maxLabel = NULL,
  fullNames = TRUE, Pvalues = TRUE)
```

Arguments

| | |
|-----------|--|
| setA | FunctionalAnnotationChart to compare |
| setB | FunctionalAnnotationChart to compare |
| ont | The gene ontology category for which to calculate enrichment |
| n | The number of top-ranked Pvalues to compare |
| maxLabel | The maximum number of characters in a node's label |
| fullNames | Whether to print the full GO term label or just the GO id |
| Pvalues | Whether to print P-values alongside each label |

Examples

```
## Not run:
data(funChart1)
data(funChart2)
plotZRankedDAG(funChart1, funChart2, n = 50)

## End(Not run)
```

| | |
|-------------|---|
| plotZScores | <i>Performs z transform on two sets of GO terms and plots scatterplot of result</i> |
|-------------|---|

Description

Generates a scatterplot of z transformed GO terms and plots the result along with the Jaccard metric for each GO term and linear fit + correlation.

Usage

```
plotZScores(setA, setB, cutoff = NULL, plotAbs = TRUE, plotNA = FALSE, model = "lm")
```

Arguments

| | |
|---------|---|
| setA | DAVIDFunctionalAnnotationChart object to compare |
| setB | DAVIDFunctionalAnnotationChart object to compare |
| plotAbs | Whether to plot the absolute values of z-scores or the raw values |
| plotNA | Whether to remove NAs entirely or set all NAs to 0 |
| model | The model to use when plotting linear fit, default 'lm' |
| cutoff | If you want to apply a Benjamini corrected P-value cutoff to each list before generating Z scores, supply it here |

Examples

```
data(funChart1)
data(funChart2)
plotZScores(funChart1, funChart2)
```

| | |
|----------------|--|
| slidingJaccard | <i>Plot two functional annotation charts using a sliding Jaccard coefficient</i> |
|----------------|--|

Description

This function compares two functional annotation charts using a sliding Jaccard coefficient - a ranked list of P-values is produced, and a sliding window is used to find the Jaccard coefficient of two charts at different cutoffs of the top n terms. This is useful to determine where the majority of overlapping terms is located, and can also be used to compare Jaccard profiles between multiple (up to 4) sets if C and D are supplied.

Usage

```
slidingJaccard(setA, setB, increment = 50, setC = NULL, setD = NULL)
```

Arguments

| | |
|-----------|--|
| setA | A DAVIDFunctionalAnnotationChart to compare |
| setB | A DAVIDFunctionalAnnotationChart to compare |
| increment | The number of terms (n) to increment for each sliding window |
| setC | A DAVIDFunctionalAnnotationChart to compare, optional |
| setD | A DAVIDFunctionalAnnotationChart to compare, optional |

Examples

```
data(funChart1)
data(funChart2)
slidingJaccard(funChart1, funChart2, 50, FALSE)
```

srf

A .bed file containing genomic ranges sampled from srf

Description

1000 genomic ranges sampled from srf binding sites identified in HL-1 cells (mm9)

Usage

```
srf
```

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

| | |
|------|--|
| tbx5 | <i>A .bed file containing genomic ranges sampled from tbx5</i> |
|------|--|

Description

1000 genomic ranges sampled from tbx5 binding sites identified in HL-1 cells (mm9)

Usage

tbx5

Format

A data frame with 1000 observations of 3 variables: chromosome, start position, end position.

Source

He, A. (2011) *Co-occupancy by multiple cardiac transcription factors identifies transcriptional enhancers active in heart*. PNAS 108(14), 5632-5637

| | |
|----------|---|
| viewKegg | <i>Compare KEGG pathways between two functional annotation charts</i> |
|----------|---|

Description

viewKegg uses pathview to compare the gene lists visually by KEGG pathway. You can either supply a pathway id or the function will pick the most differentially enriched pathway between the two inputs. As functional annotation charts don't have differential gene expression information, a boolean scale is used - genes in the pathway are coloured green if from setA, yellow if from both, and red if from setB. We recommend you supply a working directory, as pathview will download an XML and PNG file as well as output an additional PNG of the pathway.

Usage

```
viewKegg(setA, setB, keggTerm = NULL, species = NULL, workingDir = NULL,
  sortByCount = FALSE, ...)
```

Arguments

| | |
|-------------|--|
| setA | FunctionalAnnotationChart to compare |
| setB | FunctionalAnnotationChart to compare |
| keggTerm | If a specific KEGG pathway is of interest, input the name here; otherwise, the most differentially expressed pathway will be used. |
| species | The program can usually figure out the species from the KEGG terms, but if it can't, supply the species ID here. From pathview vignette, run 'data(bods); bods' to find species codes. |
| workingDir | The directory to output into. Recommended, since pathview will put a few different files there each time. |
| sortByCount | Set TRUE if you want the function to automatically choose the pathway with the most number of genes |
| ... | Arguments to be passed to pathview |

Value

Output from pathview: a list of 2, plot.data.gene and plot.data.cpd

Examples

```
## Not run:
# Since this function requires writing to a directory, it won't be run here
data(funChart1)
data(funChart2)
viewKegg(funChart1, funChart2)

## End(Not run)
```

| | |
|---------------------|---|
| zTransformDirectory | <i>Z-score transformation of DAVID functional annotation charts in a supplied directory</i> |
|---------------------|---|

Description

Given a directory of functional annotation charts, this function iterates over them and generates Odds Ratio, St. Error and Z scores. This is useful for batch processing, as all the charts can be written to disk somewhere then iterated over by this function automatically. Two options are provided for dealing with absent terms: either the NAs are set as 0 (a pseudo-representation of a Z-score with no enrichment), or incomplete rows are removed. The final table can be used for clustering analyses.

Usage

```
zTransformDirectory(inputDir, cutoff = NULL, pattern = NULL,
  removeNA = FALSE)
```

Arguments

| | |
|----------|--|
| inputDir | The directory to search for functional annotation charts |
| pattern | The regex pattern to match files in inputDir |
| cutoff | Reduce the computation to the top n GO terms ranked by variance |
| removeNA | True to only generate the Z-transform table based on GO terms common to all input enrichment analyses, False to set all NAs as 0 |

Value

Returns a data.frame of z scores, ORs and SEs

Examples

```
## Not run:
#not run as dir required
z.merge = zTransformDirectory("./fnAnot_charts", pattern = "-fnAnot.txt")
# To plot a dendrogram based on Z-scores:
d <- cor(abs(z.merge[2:(ncol(z.merge)-1)]))
dist.cor <- hclust(dist(1-d), method="complete")
```

```
plot(dist.cor, xlab="Complete linkage", sub = NA)  
## End(Not run)
```

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