Package 'GRaNIE'

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Title GRaNIE: Reconstruction cell type specific gene regulatory networks including enhancers using chromatin accessibility and RNA-seq data

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Description Genetic variants associated with diseases often affect non-coding regions, thus likely having a regulatory role. To understand the effects of genetic variants in these regulatory regions, identifying genes that are modulated by specific regulatory elements (REs) is crucial. The effect of gene regulatory elements, such as enhancers, is often cell-type specific, likely because the combinations of transcription factors (TFs) that are regulating a given enhancer have celltype specific activity. This TF activity can be quantified with existing tools such as diffTF and captures differences in binding of a TF in open chromatin regions. Collectively, this forms a gene regulatory network (GRN) with cell-type and data-specific TF-RE and RE-gene links. Here, we reconstruct such a GRN using bulk RNAseq and open chromatin (e.g., using ATACseq or ChIPseq for open chromatin marks) and optionally TF activity data. Our network contains different types of links, connecting TFs to regulatory elements, the latter of which is connected to genes in the vicinity or within the same chromatin domain (TAD). We use a statistical framework to assign empirical FDRs and weights to all links using a permutation-based approach.

Imports futile.logger, checkmate, patchwork, reshape2, data.table, matrixStats, Matrix, GenomicRanges, RColorBrewer, ComplexHeatmap, DESeq2, csaw, circlize, robust, progress, utils, methods, stringr, scales, BiocManager, BiocParallel, igraph, S4Vectors, ggplot2, rlang, Biostrings, GenomeInfoDb, IRanges, SummarizedExperiment, forcats, gridExtra, limma, purrr, tidyselect, readr, grid, tidyr, dplyr, stats, grDevices, graphics, magrittr, tibble, viridis, BiocFileCache, colorspace

Depends R (>= 4.2.0), tidyverse, topGO

Suggests knitr, BSgenome. Hsapiens. UCSC. hg19,

BSgenome.Hsapiens.UCSC.hg38, BSgenome.Mmusculus.UCSC.mm10, BSgenome.Mmusculus.UCSC.mm9, TxDb.Hsapiens.UCSC.hg19.knownGene, TxDb.Hsapiens.UCSC.hg38.knownGene,

TxDb.Mmusculus.UCSC.mm10.knownGene,

TxDb.Mmusculus.UCSC.mm9.knownGene, org.Hs.eg.db, org.Mm.eg.db,

IHW, biomaRt, clusterProfiler, ReactomePA, DOSE, ChIPseeker, testthat (>= 3.0.0), BiocStyle
VignetteBuilder knitr
biocViews Software, GeneExpression, GeneRegulation, NetworkInference, GeneSetEnrichment, BiomedicalInformatics, Genetics, Transcriptomics, ATACSeq, RNASeq, GraphAndNetwork, Regression, Transcription, ChIPSeq
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LazyData false
<pre>URL https://grp-zaugg.embl-community.io/GRaNIE</pre>
<pre>BugReports https://git.embl.de/grp-zaugg/GRaNIE/issues</pre>
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addConnections_peak_gene

Add peak-gene connections to a GRN object

Description

Add peak-gene connections to a GRN object

Usage

```
addConnections_peak_gene(
   GRN,
   overlapTypeGene = "TSS",
   corMethod = "pearson",
   promoterRange = 250000,
   TADs = NULL,
   nCores = 4,
   plotDiagnosticPlots = TRUE,
   plotGeneTypes = list(c("all"), c("protein_coding"), c("protein_coding", "lincRNA")),
```

```
outputFolder = NULL,
addRobustRegression = FALSE,
forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

overlapTypeGene

Character. "TSS" or "full". Default "TSS". If set to "TSS", only the TSS of the gene is used as reference for finding genes in the neighborhood of a peak. If set to "full", the whole annotated gene (including all exons and introns) is used instead.

used instead.

corMethod Character. pearson or spearman. Default pearson. Method for calculating the

correlation coefficient. See cor for details.

promoterRange Integer >=0. Default 250000. The size of the neighborhood in bp to correlate

peaks and genes in vicinity. Only peak-gene pairs will be correlated if they are within the specified range. Increasing this value leads to higher running times and more peak-gene pairs to be associated, while decreasing results in the

opposite.

TADs Data frame with TAD domains. Default NULL. If provided, the neighborhood

of a peak is defined by the TAD domain the peak is in rather than a fixed-sized neighborhood. The expected format is a BED-like data frame with at least 3 columns in this particular order: chromosome, start, end, the 4th column is optional and will be taken as ID column. All additional columns as well as column names are ignored. For the first 3 columns, the type is checked as part

of a data integrity check.

nCores Integer >0. Default 1. Number of cores to use.

plotDiagnosticPlots

TRUE or FALSE. Default TRUE. Run and plot various diagnostic plots? If set to TRUE, PDF files will be produced and saved in the output directory (in a sub-

folder called plots).

plotGeneTypes List of character vectors. Default list(c("all"), c("protein_coding"),

c("protein_coding", "lincRNA")). Each list element may consist of one or multiple gene types that are plotted collectively in one PDF. The special keyword "all" denotes all gene types that are found (be aware: this typically contains 20+ gene types, see https://www.gencodegenes.org/pages/

biotypes.html for details).

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

addRobustRegression

TRUE or FALSE. EXPERIMENTAL. Default FALSE. Use a robust regression in addition to a non-robust one? Significantly increases overall running time.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function in different flavors.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = addConnections_peak_gene(GRN, promoterRange=10000, plotDiagnosticPlots = FALSE)
```

addConnections_TF_peak

Add TF-peak connections to a GRN object

Description

Add TF-peak connections to a GRN object

Usage

```
addConnections_TF_peak(
   GRN,
   plotDiagnosticPlots = TRUE,
   plotDetails = FALSE,
   outputFolder = NULL,
   corMethod = "pearson",
   connectionTypes = c("expression"),
   removeNegativeCorrelation = c(FALSE),
   maxFDRToStore = 0.3,
   addForPermuted = TRUE,
   useGCCorrection = FALSE,
   percBackground_size = 75,
   percBackground_resample = TRUE,
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

plotDiagnosticPlots

TRUE or FALSE. Default TRUE. Run and plot various diagnostic plots? If set to TRUE, PDF files will be produced and saved in the output directory (in a subfolder called plots).

plotDetails

TRUE or FALSE. Default FALSE. Print additional plots that may help for debugging and QC purposes? Note that these plots are currently less documented or not at all.

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

corMethod Character. pearson or spearman. Default pearson. Method for calculating the

correlation coefficient. See cor for details.

connectionTypes

Character vector. Default expression. Vector of connection types to include for the TF-peak connections. If an additional connection type is specified here, it has to be available already within the object (EXPERIMENTAL). See the function addData_TFActivity for details.

removeNegativeCorrelation

Vector of TRUE or FALSE. Default FALSE. EXPERIMENTAL. Must be a logical vector of the same length as the parameter connectionType. Should negatively correlated TF-peak connections be removed for the specific connection type? For connection type expression, the default is FALSE, while for any TF Activity related connection type, we recommend setting this to TRUE.

maxFDRToStore Numeric. Default 0.3. Maximum TF-peak FDR value to permanently store a

particular TF-peak connection in the object? This parameter has a large influence on the overall memory size of the object, and we recommend not storing

connections with a high FDR due to their sheer number.

addForPermuted TRUE or FALSE. Default FALSE. Add connections also for permuted data. Leave

at TRUE unless you know what you are doing.

useGCCorrection

TRUE or FALSE. Default FALSE. EXPERIMENTAL. Should a GC-matched background be used when calculating FDRs?

percBackground_size

Numeric (0 to 100). Default 75. EXPERIMENTAL. Description will follow. Only relevant if useGCCorrection is set to TRUE, ignored otherwise.

percBackground_resample

TRUE or FALSE. Default TRUE. EXPERIMENTAL. Should resampling be enabled for those GC bins for which not enough background peaks are available?. Only

relevant if useGCCorrection is set to TRUE, ignored otherwise.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
```

GRN = addConnections_TF_peak(GRN, plotDiagnosticPlots = FALSE, forceRerun = FALSE)

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addData

Add data to a GRN object

Description

Add data to a GRN object

Usage

```
addData(
  GRN,
  counts_peaks,
  normalization_peaks = "DESeq_sizeFactor",
  idColumn_peaks = "peakID",
  counts_rna,
  normalization_rna = "quantile",
  idColumn_RNA = "ENSEMBL",
  sampleMetadata = NULL,
  allowOverlappingPeaks = FALSE,
  forceRerun = FALSE
)
```

Arguments

GRN

Object of class GRN

counts_peaks

Data frame. No default. Counts for the peaks, with raw or normalized counts for each peak (rows) across all samples (columns). In addition to the count data, it must also contain one ID column with a particular format, see the argument idColumn_peaks below. Row names are ignored, column names must be set to the sample names and must match those from the RNA counts and the sample metadata table.

normalization_peaks

Character. Default DESeq_sizeFactor. Normalization procedure for peak data. Must be one of DESeq_sizeFactor, none, or quantile.

idColumn_peaks Character. Default peakID. Name of the column in the counts_peaks data frame that contains peak IDs. The required format must be chr:start-end", with chr denoting the abbreviated chromosome name, and start and end the begin and end of the peak coordinates, respectively. End must be bigger than start. Examples for valid peak IDs are chr1: 400-800 or chrX: 20-25.

counts_rna

Data frame. No default. Counts for the RNA-seq data, with raw or normalized counts for each gene (rows) across all samples (columns). In addition to the count data, it must also contain one ID column with a particular format, see the argument idColumn_rna below. Row names are ignored, column names must be set to the sample names and must match those from the RNA counts and the sample metadata table.

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normalization_rna

Character. Default quantile. Normalization procedure for peak data. Must be one of "DESeq_sizeFactor", "none", or "quantile"

idColumn_RNA

Character. Default ENSEMBL. Name of the column in the counts_rna data frame that contains Ensembl IDs.

sampleMetadata

Data frame. Default NULL. Optional, additional metadata for the samples, such as age, sex, gender etc. If provided, the @seealso [plotPCA_all()] function can then incorporate and plot it. Sample names must match with those from both peak and RNA-Seq data. The first column is expected to contain the sample IDs, the actual column name is irrelevant.

allowOverlappingPeaks

TRUE or FALSE. Default FALSE. Should overlapping peaks be allowed (then only a warning is issued when overlapping peaks are found) or (the default) should an error be raised?

forceRerun

TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
# library(tidyverse)
# rna.df = read_tsv("https://www.embl.de/download/zaugg/GRaNIE/rna.tsv.gz")
# peaks.df = read_tsv("https://www.embl.de/download/zaugg/GRaNIE/peaks.tsv.gz")
# meta.df = read_tsv("https://www.embl.de/download/zaugg/GRaNIE/sampleMetadata.tsv.gz")
# GRN = loadExampleObject()
# We omit sampleMetadata = meta.df in the following line, becomes too long otherwise
# GRN = addData(GRN, counts_peaks = peaks.df, counts_rna = rna.df, forceRerun = FALSE)
```

addData_TFActivity

Add TF activity data to GRN object using a simplified procedure for estimating it. EXPERIMENTAL.

Description

Add TF activity data to GRN object using a simplified procedure for estimating it. EXPERIMENTAL.

Usage

```
addData_TFActivity(
   GRN,
   normalization = "cyclicLoess",
   name = "TF_activity",
   forceRerun = FALSE
)
```

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Arguments

GRN Object of class GRN

normalization Character. Default cyclicLoess. One of cyclicLoess, sizeFactors, quantile,

or none. Normalization procedure.

name Name in object under which it should be stored. This corresponds to the connectionType

afterwards that some functions iterate over.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

addTFBS

Add TFBS to a GRN object

Description

Add TFBS to a GRN object

Usage

```
addTFBS(
   GRN,
   motifFolder,
   TFs = "all",
   nTFMax = NULL,
   filesTFBSPattern = "_TFBS",
   fileEnding = ".bed",
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

motiffolder Character. No default. Path to the folder that contains the TFBS predictions.

The files must be in BED format, 6 columns, one file per TF. See the other

parameters for more details.

TFs Character vector. Default all. Vector of TF names to include. The special

keyword all can be used to include all TF found in the folder as specified by motifFolder. If all is specified anywhere, all TFs will be included. TF names must otherwise match the file names that are found in the folder, without the file

suffix.

nTFMax NULL or integer. Default NULL. Maximal number of TFs to import. Can be used

for testing purposes, e.g., setting to 5 only imports 5 TFs even though the whole

motifFolder has many more TFs defined.

filesTFBSPattern

fileEnding

Character. Default "_TFBS". Suffix for the file names in the TFBS folder that is not part of the TF name. Can be empty. For example, for the TF CTCF, if the file is called CTCF, all TFBS bod set this parameter to " all TFBS"

file is called CTCF.all.TFBS.bed, set this parameter to ".all.TFBS".

Character. Default ". bed". File ending for the files from the motif folder.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
```

```
add_TF_gene_correlation
```

Add TF-gene correlations to a GRN object. The information is currently stored in GRN@connections\$TF_genes.filtered. Note that raw p-values are not adjusted.

Description

Add TF-gene correlations to a GRN object. The information is currently stored in GRN@connections\$TF_genes.filtered. Note that raw p-values are not adjusted.

Usage

```
add_TF_gene_correlation(
   GRN,
   corMethod = "pearson",
   addRobustRegression = FALSE,
   nCores = 1,
   forceRerun = FALSE
)
```

Arguments

nCores

GRN Object of class GRN

corMethod Character. pearson or spearman. Default pearson. Method for calculating the

correlation coefficient. See cor for details.

addRobustRegression

TRUE or FALSE. EXPERIMENTAL. Default FALSE. Use a robust regression in addition to a non-robust one? Significantly increases overall running time.

Integer >0. Default 1. Number of cores to use.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = add_TF_gene_correlation(GRN, forceRerun = FALSE)
```

AR_classification_wrapper

Run the activator-repressor classification for the TFs for a GRN object

Description

Run the activator-repressor classification for the TFs for a GRN object

Usage

```
AR_classification_wrapper(
   GRN,
   significanceThreshold_Wilcoxon = 0.05,
   plot_minNoTFBS_heatmap = 100,
   deleteIntermediateData = TRUE,
   plotDiagnosticPlots = TRUE,
   outputFolder = NULL,
   corMethod = "pearson",
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

significanceThreshold_Wilcoxon

Numeric between 0 and 1. Default 0.05. Significance threshold for Wilcoxon test that is run in the end for the final classification. See the Vignette and *diffTF* paper for details.

plot_minNoTFBS_heatmap

Integer. Default 100. Minimum number of TFBS for a TF to be included in the heatmap that is part of the output of this function.

deleteIntermediateData

TRUE or FALSE. Default TRUE. Should intermediate data be deleted before returning the object after a successful run? Due to the size of the produced intermediate data, we recommend setting this to TRUE, but if memory or object size are not an issue, the information can also be kept.

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plotDiagnosticPlots

TRUE or FALSE. Default TRUE. Run and plot various diagnostic plots? If set to TRUE, PDF files will be produced and saved in the output directory (in a sub-

folder called plots).

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

corMethod Character. pearson or spearman. Default pearson. Method for calculating the

correlation coefficient. See cor for details.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
# GRN = loadExampleObject()
# GRN = AR_classification_wrapper(GRN, outputFolder = ".", forceRerun = FALSE)
```

build_eGRN_graph

Builds a graph out of a set of connections

Description

Builds a graph out of a set of connections

Usage

```
build_eGRN_graph(
   GRN,
   model_TF_gene_nodes_separately = FALSE,
   allowLoops = FALSE,
   removeMultiple = FALSE,
   directed = FALSE,
   forceRerun = FALSE
)
```

Arguments

```
GRN Object of class GRN model_TF_gene_nodes_separately
```

TRUE or FALSE. Default FALSE. Should TF and gene nodes be modeled separately? If set to TRUE, this may lead to unwanted effects in case of TF-TF connections (i.e., a TF regulating another TF)

allowLoops TRUE or FALSE. Default FALSE. Allow loops in the network (i.e., a TF that regu-

lates itself)

removeMultiple TRUE or FALSE. Default FALSE. Remove loops with the same start and end point?

This can happen if multiple TF originate from the same gene, for example.

directed TRUE or FALSE. Default FALSE. Should the network be directed?

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = build_eGRN_graph(GRN, forceRerun = FALSE)
```

calculateCommunitiesEnrichment

Enrichment analysis for the genes in each community in the filtered GRN

Description

After the vertices of the filtered GRN are clustered into communities using calculateCommunitiesStats, this function will run a per-community enrichment analysis.

Usage

```
calculateCommunitiesEnrichment(
   GRN,
   ontology = c("GO_BP", "GO_MF"),
   algorithm = "weight01",
   statistic = "fisher",
   background = "neighborhood",
   selection = "byRank",
   communities = seq_len(10),
   pAdjustMethod = "BH",
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

ontology Character vector of ontologies. Default c("GO_BP", "GO_MF"). Valid values are

"GO_BP", "GO_MF", "GO_CC", "KEGG", "DO", and "Reactome", referring to GO Biological Process, GO Molecular Function, GO Cellular Component, KEGG,

Disease Ontology, and Reactome Pathways.

algorithm Character. Default "weight01". One of: "classic", "elim", "weight", "weight01",

"lea", "parentchild". Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), ignored otherwise. Name of the algorithm that handles the GO graph

structures. Valid inputs are those supported by the topGO library.

statistic Character. Default "fisher". One of: "fisher", "ks", "t", "globaltest",

"sum", "ks.ties". Statistical test to be used. Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), and valid inputs are those supported by the topGO library, ignored otherwise. For the other ontologies the test statistic is

always Fisher.

background Character. Default "neighborhood". One of: "all_annotated", "all_RNA",

"neighborhood". Set of genes to be used to construct the background for the enrichment analysis. This can either be all annotated genes in the reference genome (all_annotated), all differentially expressed genes (all_RNA), or all the genes that are within the neighborhood of a peak in the GRN (neighborhood)

selection Character. Default "byRank". One of: "byRank", "byLabel". Specify whether

the communities enrichment will by calculated based on their rank, where the largest community (with most vertices) would have a rank of 1, or by their label.

Note that the label is independent of the rank.

communities Numeric vector. Default c(1:10). Depending on what was specified in the

display parameter, this parameter would indicate either the rank or the label of the communities to be plotted. i.e. for communities = c(1,4), if display = "byRank" the GO enrichment for the first and fourth largest communities will be calculated if display = "byLabel", the results for the communities labeled

"1", and "4" will be plotted.

pAdjustMethod Character. Default "BH". One of: "holm", "hochberg", "hommel", "bonferroni",

"BH", "BY", "fdr". This parameter is only relevant for the following ontologies: KEGG, DO, Reactome. For the other ontologies, the algorithm serves as an ad-

justment.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with the enrichment results stored in the stats\$Enrichment\$byCommunity slot.

See Also

plotCommunitiesEnrichment
plotGeneralEnrichment
calculateGeneralEnrichment

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Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = calculateCommunitiesEnrichment(GRN, ontology = c("GO_BP"), forceRerun = FALSE)
```

calculateCommunitiesStats

Generate graph communities and their summarizing statistics

Description

This function generates the TF-gene graph from the filtered GRN object, and clusters its vertices into communities using established community detection algorithms.

Usage

```
calculateCommunitiesStats(GRN, clustering = "louvain", forceRerun = FALSE, ...)
```

Arguments

GRN	Object of class	GRN

clustering Character. Default louvain. One of: louvain, leiden, leading_eigen, fast_greedy,

optimal, walktrap. The community detection algorithm to be used. Please bear in mind the robustness and time consumption of the algorithms when opt-

ing for an alternative to the default.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

... Additional parameters for the used clustering method, see the igraph::cluster_*

methods for details on the specific parameters and what they do. For leiden clustering, for example, you may add a resolution_parameter to control the granularity of the community detection or n_iterations to modify the number

of iterations.

Value

The same GRN object, with a table that consists of the connections clustered into communities stored in the stats\$communities slot.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = calculateCommunitiesStats(GRN, forceRerun = FALSE)
```

 ${\tt calculateGeneralEnrichment}$

Run an enrichment analysis for the genes in the filtered GRN

Description

This function runs an enrichment analysis for the genes in the filtered network.

Usage

```
calculateGeneralEnrichment(
   GRN,
   ontology = c("GO_BP", "GO_MF"),
   algorithm = "weight01",
   statistic = "fisher",
   background = "neighborhood",
   pAdjustMethod = "BH",
   forceRerun = FALSE
)
```

Arguments

GRN	Object of class GRN
ontology	Character vector of ontologies. Default c("GO_BP", "GO_MF"). Valid values are "GO_BP", "GO_MF", "GO_CC", "KEGG", "DO", and "Reactome", referring to GO Biological Process, GO Molecular Function, GO Cellular Component, KEGG, Disease Ontology, and Reactome Pathways.
algorithm	Character. Default "weight01". One of: "classic", "elim", "weight", "weight01" "lea", "parentchild". Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), ignored otherwise. Name of the algorithm that handles the GO graph structures. Valid inputs are those supported by the topGO library.
statistic	Character. Default "fisher". One of: "fisher", "ks", "t", "globaltest", "sum", "ks.ties". Statistical test to be used. Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), and valid inputs are those supported by the topGO library, ignored otherwise. For the other ontologies the test statistic is always Fisher.
background	Character. Default "neighborhood". One of: "all_annotated", "all_RNA", "neighborhood". Set of genes to be used to construct the background for the enrichment analysis. This can either be all annotated genes in the reference genome (all_annotated), all differentially expressed genes (all_RNA), or all the genes that are within the neighborhood of a peak in the GRN (neighborhood)
pAdjustMethod	Character. Default "BH". One of: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr". This parameter is only relevant for the following ontologies: KEGG, DO, Reactome. For the other ontologies, the algorithm serves as an adjustment.

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forceRerun

TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already contains the result. Overwrites the old results.

Value

The same GRN object, with the enrichment results stored in the stats\$Enrichment\$general slot.

See Also

```
plotGeneralEnrichment
calculateCommunitiesEnrichment
plotCommunitiesEnrichment
```

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = calculateGeneralEnrichment(GRN, ontology = "GO_BP", forceRerun = FALSE)
```

calculateTFEnrichment Calculate TF-based GO enrichment

Description

This function calculates the GO enrichment per TF, i.e. for the set of genes a given TF is connected to in the filtered GRN.

Usage

```
calculateTFEnrichment(
   GRN,
   rankType = "degree",
   n = 3,
   TF.names = NULL,
   ontology = c("GO_BP", "GO_MF"),
   algorithm = "weight01",
   statistic = "fisher",
   background = "neighborhood",
   pAdjustMethod = "BH",
   forceRerun = FALSE
)
```

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Arguments

n

GRN Object of class GRN

Character. Default "degree". One of: "degree", "EV", "custom". This paramrankType

> eter will determine the criterion to be used to identify the "top" TFs. If set to "degree", the function will select top TFs based on the number of connections to genes they have, i.e. based on their degree-centrality. If set to "EV" it will select the top TFs based on their eigenvector-centrality score in the network. If set to

custom, a set of TF names will have to be passed to the "TF.names" parameter.

Numeric. Default 3. If this parameter is passed as a value between 0 and 1, it is treated as a percentage of top nodes. If the value is passed as an integer it will be treated as the number of top nodes. This parameter is not relevant if rankType

= "custom".

TF.names Character vector. Default NULL. If the rank type is set to "custom", a vector of

TF names for which the GO enrichment should be calculated should be passed

to this parameter.

ontology Character vector of ontologies. Default c("GO_BP", "GO_MF"). Valid values are

> "GO_BP", "GO_MF", "GO_CC", "KEGG", "DO", and "Reactome", referring to GO Biological Process, GO Molecular Function, GO Cellular Component, KEGG,

Disease Ontology, and Reactome Pathways.

Character. Default "weight01". One of: "classic", "elim", "weight", "weight01", algorithm

> "lea", "parentchild". Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), ignored otherwise. Name of the algorithm that handles the GO graph

structures. Valid inputs are those supported by the topGO library.

Character. Default "fisher". One of: "fisher", "ks", "t", "globaltest", statistic

> "sum", "ks.ties". Statistical test to be used. Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), and valid inputs are those supported by the topGO library, ignored otherwise. For the other ontologies the test statistic is

always Fisher.

background Character. Default "neighborhood". One of: "all_annotated", "all_RNA",

> "neighborhood". Set of genes to be used to construct the background for the enrichment analysis. This can either be all annotated genes in the reference genome (all_annotated), all differentially expressed genes (all_RNA), or all the genes that are within the neighborhood of a peak in the GRN (neighborhood)

pAdjustMethod Character. Default "BH". One of: "holm", "hochberg", "hommel", "bonferroni",

> "BH", "BY", "fdr". This parameter is only relevant for the following ontologies: KEGG, DO, Reactome. For the other ontologies, the algorithm serves as an ad-

justment.

TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already forceRerun

contains the result. Overwrites the old results.

Value

The same GRN object, with the enrichment results stored in the stats\$Enrichment\$byTF slot.

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Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = calculateTFEnrichment(GRN, n = 5, ontology = "GO_BP", forceRerun = FALSE)
```

changeOutputDirectory Change the output directory of a GRN object

Description

Change the output directory of a GRN object

Usage

```
changeOutputDirectory(GRN, outputDirectory = ".")
```

Arguments

```
GRN Object of class GRN outputDirectory
```

Character. Default .. New output directory for all output files unless overwritten via the parameter outputFolder.

Value

The same GRN object, with the output directory being adjusted accordingly

Examples

```
GRN = loadExampleObject()
GRN = changeOutputDirectory(GRN, outputDirectory = ".")
```

deleteIntermediateData

Optional convenience function to delete intermediate data from the function AR_classification_wrapper and summary statistics that may occupy a lot of space

Description

Optional convenience function to delete intermediate data from the function AR_classification_wrapper and summary statistics that may occupy a lot of space

Usage

```
deleteIntermediateData(GRN)
```

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Arguments

GRN

Object of class GRN

Value

The same GRN object, with some slots being deleted (GRN@data\$TFs\$classification as well as GRN@stats\$connectionDetails.1)

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
# GRN = loadExampleObject()
# GRN = deleteIntermediateData(GRN)
```

filterData

Filter data from a GRN object

Description

Filter data from a GRN object

Usage

```
filterData(
   GRN,
   minNormalizedMean_peaks = 5,
   maxNormalizedMean_peaks = NULL,
   minNormalizedMeanRNA = 1,
   maxNormalizedMeanRNA = NULL,
   chrToKeep_peaks = NULL,
   minSize_peaks = NULL,
   maxSize_peaks = 10000,
   minCV_peaks = NULL,
   maxCV_peaks = NULL,
   minCV_genes = NULL,
   maxCV_genes = NULL,
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN minNormalizedMean_peaks

Numeric or NULL. Default 5. Minimum mean across all samples for a peak to be retained for the normalized counts table. Set to NULL for not applying the filter.

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maxNormalizedMean_peaks

Numeric or NULL. Default NULL. Maximum mean across all samples for a peak to be retained for the normalized counts table. Set to NULL for not applying the filter.

minNormalizedMeanRNA

Numeric or NULL. Default 5. Minimum mean across all samples for a gene to be retained for the normalized counts table. Set to NULL for not applying the filter.

maxNormalizedMeanRNA

Numeric or NULL. Default NULL. Maximum mean across all samples for a gene to be retained for the normalized counts table. Set to NULL for not applying the filter.

chrToKeep_peaks

Character vector or NULL. Default NULL. Vector of chromosomes that peaks are allowed to come from. This filter can be used to filter sex chromosomes from the peaks, for example (e.g, c(paste0("chr", 1:22), "chrX", "chrY"))

minSize_peaks Integer or NULL. Default NULL. Minimum peak size (width, end - start) for a peak

to be retained. Set to NULL for not applying the filter.

maxSize_peaks Integer or NULL. Default 10000. Maximum peak size (width, end - start) for a

peak to be retained. Set to NULL for not applying the filter.

minCV_peaks Numeric or NULL. Default NULL. Minimum CV (coefficient of variation, a unit-

less measure of variation) for a peak to be retained. Set to NULL for not applying

the filter.

maxCV_peaks Numeric or NULL. Default NULL. Maximum CV (coefficient of variation, a unit-

less measure of variation) for a peak to be retained. Set to NULL for not applying

the filter.

minCV_genes Numeric or NULL. Default NULL. Minimum CV (coefficient of variation, a unit-

less measure of variation) for a gene to be retained. Set to NULL for not applying

the filter.

maxCV_genes Numeric or NULL. Default NULL. Maximum CV (coefficient of variation, a unit-

less measure of variation) for a gene to be retained. Set to NULL for not applying

the filter.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = filterData(GRN, forceRerun = FALSE)
```

filterGRNAndConnectGenes

Filter the GRN and integrate peak-gene connections.

Description

This is one of the main integrative functions of the GRN package. It has two main functions: Filtering the TF-peak and peak-gene connections that have been identified before, and combining the 3 major elements (TFs, peaks, genes) into one data frame, with one row per connection. Here, a connection can either be a TF-peak, peak-gene or TF-peak-gene link, depending on the parameters. Internally, first, the TF-peak are filtered before the peak-gene connections are added for reasons of memory and computational efficacy: It takes a lot of time and particularly space to connect the full GRN with all peak-gene connections - as most of the links have weak support (i.e., high FDR), first filtering out unwanted links dramatically reduces the memory needed for the combined GRN

Usage

```
filterGRNAndConnectGenes(
  GRN,
  TF_peak.fdr.threshold = 0.2,
  TF_peak.connectionTypes = "all",
  peak_gene.p_raw.threshold = NULL,
  peak_gene.fdr.threshold = 0.2,
  peak_gene.fdr.method = "BH",
  peak_gene.IHW.covariate = NULL,
  peak_gene.IHW.nbins = 5,
  gene.types = c("protein_coding", "lincRNA"),
  allowMissingTFs = FALSE,
  allowMissingGenes = TRUE,
  peak\_gene.r\_range = c(0, 1),
  peak_gene.selection = "all";
  peak_gene.maxDistance = NULL,
  filterTFs = NULL,
  filterGenes = NULL,
  filterPeaks = NULL,
  TF_peak_FDR_selectViaCorBins = FALSE,
  filterLoops = TRUE,
  outputFolder = NULL,
  silent = FALSE
)
```

Arguments

GRN Object of class GRN TF_peak.fdr.threshold

Numeric[0,1]. Default 0.2. Maximum FDR for the TF-peak links. Set to 1 or NULL to disable this filter.

TF_peak.connectionTypes

Character vector. Default all. TF-peak connection types to consider. The special keyword all denotes all connection types (e.g., expression and TFActivity) that are found in the GRN object. By default, only expression is present in the object, so all and expression are usually equivalent unless calculation of TF-peak links based on TF activity has also been enabled.

peak_gene.p_raw.threshold

Numeric[0,1]. Default NULL. Threshold for the peak-gene connections, based on the raw p-value. All peak-gene connections with a larger raw p-value will be filtered out.

peak_gene.fdr.threshold

Numeric[0,1]. Default 0.2. Threshold for the peak-gene connections, based on the FDR. All peak-gene connections with a larger FDR will be filtered out.

peak_gene.fdr.method

Character. Default "BH". One of: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none", "IHW". Method for adjusting p-values for multiple testing. If set to "IHW", independent hypothesis weighting will be performed, and a suitable covariate has to be specified for the parameter peak_gene. IHW. covariate.

peak_gene.IHW.covariate

Character. Default NULL. Name of the covariate to use for IHW (column name from the table thatis returned with the function getGRNConnections. Only relevant if peak_gene.fdr.method is set to "IHW". You have to make sure the specified covariate is suitable or IHW, see the diagnostic plots that are generated in this function for this. For many datasets, the peak-gene distance (called peak_gene.distance in the object) seems suitable.

peak_gene.IHW.nbins

Integer or "auto". Default 5. Number of bins for IHW. Only relevant if peak_gene.fdr.method is set to "IHW".

gene.types

Character vector of supported gene types. Default c("protein_coding", "lincRNA"). Filter for gene types to retain, genes with other gene types are filtered.

allowMissingTFs

TRUE or FALSE. Default FALSE. Should connections be returned for which the TF is NA (i.e., connections consisting only of peak-gene links?). If set to TRUE, this generally greatly increases the number of connections but it may not be what you aim for.

allowMissingGenes

TRUE or FALSE. Default TRUE. Should connections be returned for which the gene is NA (i.e., connections consisting only of TF-peak links?). If set to TRUE, this generally increases the number of connections.

peak_gene.r_range

Numeric(2). Default c(0,1). Filter for lower and upper limit for the peak-gene links. Only links will be retained if the correlation coefficient is within the specified interval. This filter is usually used to filter out negatively correlated peak-gene links.

peak_gene.selection

"all" or "closest". Default "all". Filter for the selection of genes for each peak. If set to "all", all previously identified peak-gene are used, while

"closest" only retains the closest gene for each peak that is retained until the point the filter is applied.

peak_gene.maxDistance

Integer >0. Default NULL. Maximum peak-gene distance to retain a peak-gene

connection.

filterTFs Character vector. Default NULL. Vector of TFs (as named in the GRN object) to

retain. All TFs not listed will be filtered out.

filterGenes Character vector. Default NULL. Vector of gene IDs (as named in the GRN ob-

ject) to retain. All genes not listed will be filtered out.

filterPeaks Character vector. Default NULL. Vector of peak IDs (as named in the GRN ob-

ject) to retain. All peaks not listed will be filtered out.

TF_peak_FDR_selectViaCorBins

TRUE or FALSE. Default FALSE. Use a modified procedure for selecting TF-peak links that is based on the user-specified FDR but that retains also links that may

have a higher FDR but a more extreme correlation.

filterLoops TRUE or FALSE. Default TRUE. If a TF regulates itself (i.e., the TF and the gene

are the same entity), should such loops be filtered from the GRN?

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

silent TRUE or FALSE. Default FALSE. Print progress messages and filter statistics.

Value

The same GRN object, with the filtered and merged TF-peak and peak-gene connections in the slot connections \$all.filtered.

See Also

```
visualizeGRN
addConnections_TF_peak
addConnections_peak_gene
```

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
```

GRN = filterGRNAndConnectGenes(GRN)

generateStatsSummary Generate a summary PDF for the number of connections for a GRN object.

Description

Essentially, this functions calls filterGRNAndConnectGenes repeatedly and stores the total number of connections and other statistics each time to summarize them afterwards. All arguments are identical to the ones in filterGRNAndConnectGenes, see the help for this function for details.

Usage

```
generateStatsSummary(
   GRN,
   TF_peak.fdr = c(0.001, 0.01, 0.05, 0.1, 0.2),
   TF_peak.connectionTypes = "all",
   peak_gene.fdr = c(0.001, 0.01, 0.05, 0.1, 0.2),
   peak_gene.p_raw = NULL,
   peak_gene.r_range = c(0, 1),
   gene.types = c("protein_coding", "lincRNA"),
   allowMissingGenes = c(FALSE, TRUE),
   allowMissingTFs = c(FALSE),
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

TF_peak.fdr Numeric vector. Default c(0.001, 0.01, 0.05, 0.1, 0.2). TF-peak FDR val-

ues to iterate over.

TF_peak.connectionTypes

Character vector. Default all. TF-peak connection types to consider. The special keyword all denotes all connection types (e.g., expression and TFActivity) that are found in the GRN object. By default, only expression is present in the object, so all and expression are usually equivalent unless calculation of TF-peak links based on TF activity has also been enabled.

peak_gene.fdr Numeric vector. Default c(0.001, 0.01, 0.05, 0.1, 0.2). Peak-gene FDR values to iterate over.

peak_gene.p_raw

Numeric vector. Default NULL. Peak-gene raw p-value values to iterate over. Skipped if set to NULL.

peak_gene.r_range

Numeric vector of length 2 (minimum -1, maximum 1). Default c(0,1). The correlation range of peak-gene connections to keep.

gene.types Character vector of supported gene types. Default c("protein_coding", "lincRNA"). Filter for gene types to retain, genes with other gene types are filtered.

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allowMissingGenes

Logical vector. Default c(FALSE, TRUE). Allow genes to be missing for peak-

gene connections?

allowMissingTFs

Logical vector. Default c(FALSE). Allow TFs to be missing for TF-peak con-

nections?

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples GRN = loadExampleObject() GRN = generateStatsSummary(GRN, TF_peak.fdr = c(0.01, 0.1), peak_gene.fdr = c(0.01, 0.1))
```

getCounts

Get counts for the various data defined in a GRN object

Description

Get counts for the various data defined in a GRN object.

Usage

```
getCounts(GRN, type, norm, permuted = FALSE)
```

Arguments

GRN Object of class GRN

type Character. Either peaks or rna. peaks corresponds to the counts for the open

chromatin data, while rna refers to th RNA-seq counts. If set to rna, both permuted and non-permuted data can be retrieved, while for peaks, only the

non-permuted one (i.e., 0) can be retrieved.

norm Logical. TRUE or FALSE. Should original (often raw, but this may not necessarily

be the case) or normalized counts be returned?

permuted TRUE or FALSE. Default FALSE. Should the permuted data be taken (TRUE) or the

non-permuted, original one (FALSE)?

Value

Data frame of counts, with the type as indicated by the function parameters.

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Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
counts.df = getCounts(GRN, type = "peaks", norm = TRUE, permuted = FALSE)
```

getGRNConnections

Extract connections from a GRN object

Description

Extract connections from a GRN object

Usage

```
getGRNConnections(
   GRN,
   type = "all.filtered",
   permuted = FALSE,
   include_TF_gene_correlations = FALSE)
```

Arguments

GRN Object of class GRN

type Character. Default all.filtered. Must be one of TF_peaks, peak_genes,

all.filtered. The type of connections to retrieve.

permuted TRUE or FALSE. Default FALSE. Should the permuted data be taken (TRUE) or the

non-permuted, original one (FALSE)?

include_TF_gene_correlations

Logical. TRUE or FALSE. Should TFs and gene correlations be returned as well?

If set to TRUE, they must have been computed beforehand with ${\sf add_TF_gene_correlation}$.

Value

A data frame with the connections. Importantly, this function does **NOT** return a GRN object.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN_con.all.df = getGRNConnections(GRN)
```

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getParameters	Retrieve parameters for previously used function calls and general parameters for a GRN object.

Description

Retrieve parameters for previously used function calls and general parameters for a GRN object.

Usage

```
getParameters(GRN, type = "parameter", name = "all")
```

Arguments

name

GRN Object of class GRN	
type Character. Default parameter. Either function or parameter function, a valid GRaNIE function name must be given that has When set to parameter, in combination with name, returns a sp (as specified in GRN@config)).	oeen run before.

Character. Default all. Name of parameter or function name to retrieve. Set to

the special keyword all to retrieve all parameters.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
params.l = getParameters(GRN, type = "parameter", name = "all")
```

getTopNodes

Retrieve top Nodes in the filtered GRN

Description

Retrieve top Nodes in the filtered GRN

Usage

```
getTopNodes(GRN, nodeType, rankType, n = 0.1, use_TF_gene_network = TRUE)
```

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Arguments

GRN Object of class GRN

nodeType Character. One of: "gene" or "TF". Node type.

rankType Character. One of: "degree", "EV". This parameter will determine the crite-

rion to be used to identify the "top" nodes. If set to "degree", the function will select top nodes based on the number of connections they have, i.e. based on their degree-centrality. If set to "EV" it will select the top nodes based on their

eigenvector-centrality score in the network.

Numeric. Default 0.1. If this parameter is passed as a value between (0,1), it is

treated as a percentage of top nodes. If the value is passed as an integer it will

be treated as the number of top nodes.

use_TF_gene_network

TRUE or FALSE. Default TRUE. Should the TF-gene network be used (TRUE) or

the TF-peak-gene network (FALSE)?

Value

A dataframe with the node names and the corresponding scores used to rank them

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
topGenes = getTopNodes(GRN, nodeType = "gene", rankType = "degree", n = 3)
topTFs = getTopNodes(GRN, nodeType = "TF", rankType = "EV", n = 5)
```

GRaNIE

GRaNIE (Gene Regulatory Network Inference including Enhancers): Reconstruction and evaluation of data-driven, cell type specific gene regulatory networks including enhancers using chromatin accessibility and RNAseq data (general package information)

Description

Genetic variants associated with diseases often affect non-coding regions, thus likely having a regulatory role. To understand the effects of genetic variants in these regulatory regions, identifying genes that are modulated by specific regulatory elements (REs) is crucial. The effect of gene regulatory elements, such as enhancers, is often cell-type specific, likely because the combinations of transcription factors (TFs) that are regulating a given enhancer have celltype specific activity. This TF activity can be quantified with existing tools such as diffTF and captures differences in binding of a TF in open chromatin regions. Collectively, this forms a gene regulatory network (eGRN) with cell-type and data-specific TF-RE and RE-gene links. Here, we reconstruct such a eGRN using bulk RNAseq and open chromatin (e.g., using ATACseq or ChIPseq for open chromatin marks) and optionally TF activity data. Our network contains different types of links, connecting TFs to regulatory elements, the latter of which is connected to genes in the vicinity or within the same chromatin domain (TAD). We use a statistical framework to assign empirical FDRs and weights to all links using a permutation-based approach.

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Package functions

See the Vignettes for a workflow example and more generally https://grp-zaugg.embl-community.io/GRaNIE/articles/ for all project-related information.

GRN object

The GRaNIE package works with GRN objects. See GRN for details.

Contact Information

Please check out https://grp-zaugg.embl-community.io/GRaNIE for how to get in contact with us.

GRN-class Create, represent, investigate, quantify and visualize enhancermediated gene regulatory networks (eGRNs)

Description

The class GRN stores data and information related to our eGRN approach to construct enhancer-mediated gene regulatory networks out of open chromatin and RNA-Seq data. See the description below for more details, and visit our project website at https://grp-zaugg.embl-community.io/GRaNIE and have a look at the various Vignettes.

Slots

data Currently stores 4 different types of data:

- peaks:
 - counts_norm:
 - counts_orig:
 - consensusPeaks:
- RNA:
 - counts_norm.l:
 - counts_orig:
- metadata:
- TFs:
 - translationTable:
 - TF_peak_overlap:
 - classification:

config Contains general configuration data and parameters such as parameters, files, directories, flags, and recorded function parameters.

connections Stores various types of connections annotation Stores annotation data for peaks and genes

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stats Stores statistical and summary information for a GRN network. Currently, connection details are stored here.

visualization Stores visualization results, currently always empty. Feature in development.

isDev Flag whether this is an object from the development version of the package

Constructors

Currently, a GRN object is created by executing the function initializeGRN.

Accessors

In the following code snippets, GRN is a GRN object.

Get general annotation of a GRaNIE object

nPeaks(GRN)) and nGenes(GRN)): Retrieve the number of peaks and genes, respectively, that have been added to the object (both before and after filtering)

importTFData

Import externally derived TF Activity data. EXPERIMENTAL.

Description

Import externally derived TF Activity data. EXPERIMENTAL.

Usage

```
importTFData(
   GRN,
   data,
   name,
   idColumn = "ENSEMBL",
   nameColumn = "TF.name",
   normalization = "none",
   forceRerun = FALSE
)
```

Arguments

GRN	Object of class GRN
GIAIT	Object of class diff

data Data frame. No default. Data with TF data.

name Name in object under which it should be stored. This corresponds to the connectionType

afterwards that some functions iterate over.

idColumn Character. Default ENSEMBL. Name of the ID column. Must not be unique as

some TFs may correspond to the same ID.

nameColumn Character. Default TF. name. Must be unique for each TF / row.

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normalization Character. Default cyclicLoess. One of cyclicLoess, sizeFactors, quantile,

or none. Normalization procedure.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

initializeGRN Initialize a GRN object

Description

Initialize a GRN object

Usage

```
initializeGRN(objectMetadata = list(), outputFolder, genomeAssembly)
```

Arguments

objectMetadata List. Default list(). Optional (named) list with an arbitrary number of el-

ements, all of which capture metadata for the object. This is mainly used to distinguish GRN objects from one another by storing object-specific metadata

along with the data.

outputFolder Output folder, either absolute or relative to the current working directory. No de-

fault. Default output folder where all pipeline output will be put unless specified otherwise. We recommend specifying an absolute path. Note that for Windowsbased systems, the path must be correctly specified with "/" as path separator.

genomeAssembly Character. No default. The genome assembly of all data that to be used within

this object. Currently, supported genomes are: hg19, hg38, and mm10.

Value

Empty GRN object

Examples

```
meta.l = list(name = "exampleName", date = "01.03.22")
GRN = initializeGRN(objectMetadata = meta.l, outputFolder = "output", genomeAssembly = "hg38")
```

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loadExampleObject

Load example GRN dataset

Description

Load example GRN dataset

Usage

```
loadExampleObject(
  forceDownload = FALSE,
  fileURL = "https://www.embl.de/download/zaugg/GRaNIE/GRN.rds"
)
```

Arguments

forceDownload

TRUE or FALSE. Default FALSE. Should the download be enforced even if the

local cached file is already present?

fileURL

Character. Default https://www.embl.de/download/zaugg/GRaNIE/GRN.rds.

URL to the GRN example object in rds format.

Value

An example GRN object

Examples

```
GRN = loadExampleObject()
```

nGenes

Get the number of genes for a GRN object.

Description

Return the number of genes (all or only non-filtered ones) that are defined in the GRN object.

Usage

```
nGenes(GRN, filter = TRUE)
```

Arguments

GRN Object of class GRN

filter TRUE or FALSE. Default TRUE. Should genes marked as filtered be included

in the count?

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Value

Integer. Number of genes that are defined in the GRN object, either by excluding (filter = TRUE) or including (filter = FALSE) genes that are currently marked as *filtered*.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
nGenes(GRN, filter = TRUE)
nGenes(GRN, filter = FALSE)
```

nPeaks

Get the number of peaks for a GRN object.

Description

Return the number of peaks (all or only non-filtered ones) that are defined in the GRN object.

Usage

```
nPeaks(GRN, filter = TRUE)
```

Arguments

GRN Object of class GRN

filter TRUE or FALSE. Default TRUE. Should peaks marked as filtered be included

in the count?

Value

Integer. Number of peaks that are defined in the GRN object, either by excluding (filter = TRUE) or including (filter = FALSE) peaks that are currently marked as *filtered*.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
nPeaks(GRN, filter = TRUE)
nPeaks(GRN, filter = FALSE)
```

overlapPeaksAndTFBS

Overlap peaks and TFBS for a GRN object

Description

Overlap peaks and TFBS for a GRN object

Usage

```
overlapPeaksAndTFBS(GRN, nCores = 2, forceRerun = FALSE)
```

Arguments

GRN Object of class GRN

nCores Integer >0. Default 1. Number of cores to use.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = overlapPeaksAndTFBS(GRN, nCores = 2, forceRerun = FALSE)
```

performAllNetworkAnalyses

Perform all network-related statistical and descriptive analyses, including community and enrichment analyses.

Description

A convenience function that calls all network-related functions in one-go, using selected default parameters and a set of adjustable ones also. For full adjustment, run the individual functions separately.

Usage

```
performAllNetworkAnalyses(
  GRN,
  ontology = c("GO_BP", "GO_MF"),
  algorithm = "weight01",
  statistic = "fisher",
  background = "neighborhood",
  clustering = "louvain",
  communities = seq_len(10),
  display = "byRank",
  topnGenes = 20,
  topnTFs = 20,
  maxWidth_nchar_plot = 50,
  display_pAdj = FALSE,
  outputFolder = NULL,
  forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

ontology Character vector of ontologies. Default c("GO_BP", "GO_MF"). Valid values are

"GO_BP", "GO_MF", "GO_CC", "KEGG", "DO", and "Reactome", referring to GO Biological Process, GO Molecular Function, GO Cellular Component, KEGG,

Disease Ontology, and Reactome Pathways.

algorithm Character. Default "weight01". One of: "classic", "elim", "weight", "weight01",

"lea", "parentchild". Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), ignored otherwise. Name of the algorithm that handles the GO graph

structures. Valid inputs are those supported by the topGO library.

statistic Character. Default "fisher". One of: "fisher", "ks", "t", "globaltest",

"sum", "ks.ties". Statistical test to be used. Only relevant if ontology is GO related (GO_BP, GO_MF, GO_CC), and valid inputs are those supported by the topGO library, ignored otherwise. For the other ontologies the test statistic is

always Fisher.

background Character. Default "neighborhood". One of: "all_annotated", "all_RNA",

"neighborhood". Set of genes to be used to construct the background for the enrichment analysis. This can either be all annotated genes in the reference genome (all_annotated), all differentially expressed genes (all_RNA), or all the

genes that are within the neighborhood of a peak in the GRN (neighborhood)

clustering Character. Default louvain. One of: louvain, leiden, leading_eigen, fast_greedy,

optimal, walktrap. The community detection algorithm to be used. Please bear in mind the robustness and time consumption of the algorithms when opt-

ing for an alternative to the default.

communities Numeric vector. Default seq_len(10). Depending on what was specified in the display parameter, this parameter would indicate either the rank or the label

of the communities to be plotted. i.e. for communities = c(1,4), if display =

"byRank" the results for the first and fourth largest communities will be plotted. if display = "byLabel", the results for the communities labeled "1", and "4"

will be plotted. If set to NULL, all communities will be plotted

display Character. Default "byRank". One of: "byRank", "byLabel". Specify whether

the communities will be displayed based on their rank, where the largest community (with most vertices) would have a rank of 1, or by their label. Note that

the label is independent of the rank.

topnGenes Integer. Default 20. Number of genes to plot, sorted by their rank or label.

topnTFs Integer. Default 20. Number of TFs to plot, sorted by their rank or label.

maxWidth_nchar_plot

Integer (>=10). Default 50. Maximum number of characters for a term before it

is truncated.

display_pAdj TRUE or FALSE. Default FALSE. Is the p-value being displayed in the plots the

adjusted p-value? This parameter is relevant for KEGG, Disease Ontology, and

Reactome enrichments, and does not affect GO enrichments.

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
```

GRN = loadExampleObject()

GRN = performAllNetworkAnalyses(GRN, outputFolder = ".", forceRerun = FALSE)

plotCommunitiesEnrichment

Plot community-based enrichment results

Description

Similarly to plotGeneralEnrichment, the results of the community-based enrichment analysis are plotted. By default, the results for the 10 largest communities are displayed. Additionally, if a general enrichment analysis was previously generated, this function plots an additional heatmap to compare the general enrichment with the community based enrichment. A reduced version of this heatmap is also produced where terms are filtered out to improve visibility and display and highlight the most significant terms.

Usage

```
plotCommunitiesEnrichment(
  GRN,
  outputFolder = NULL,
  basenameOutput = NULL,
  display = "byRank",
  communities = NULL,
  topn_pvalue = 30,
  p = 0.05,
  nSignificant = 2,
  nID = 10,
 maxWidth_nchar_plot = 50,
  display_pAdj = FALSE,
  plotAsPDF = TRUE,
  pdf_width = 12,
  pdf_height = 12,
  pages = NULL,
  forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

display Character. Default "byRank". One of: "byRank", "byLabel". Specify whether

the communities will be displayed based on their rank, where the largest community (with most vertices) would have a rank of 1, or by their label. Note that

the label is independent of the rank.

communities NULL or numeric vector. Default NULL. If set to NULL, the default, all communi-

ties enrichments that have been calculated before are plotted. If a numeric vector is specified: Depending on what was specified in the display parameter, this parameter indicates either the rank or the label of the communities to be plotted. i.e. for communities = c(1,4), if display = "byRank" the results for the first and fourth largest communities are plotted. if display = "byLabel", the results

for the communities labeled "1", and "4" are plotted.

topn_pvalue Numeric. Default 30. Maximum number of ontology terms that meet the p-value

significance threshold to display in the enrichment dot plot

p Numeric. Default 0.05. p-value threshold to determine significance.

nSignificant Numeric. Default 3. Threshold to filter out an ontology term with less than

nSignificant overlapping genes.

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nID Numeric. Default 10. For the reduced heatmap, number of top terms to select per community.

maxWidth_nchar_plot

Integer (>=10). Default 50. Maximum number of characters for a term before it

is truncated.

display_pAdj TRUE or FALSE. Default FALSE. Is the p-value being displayed in the plots the

adjusted p-value? This parameter is relevant for KEGG, Disease Ontology, and

Reactome enrichments, and does not affect GO enrichments.

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, without modifications. A single PDF file is produced with the results.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotCommunitiesEnrichment(GRN, plotAsPDF = FALSE)
```

 ${\it plot Communities Stats} \quad {\it Plot general structure \& connectivity statistics for each community in} \\ a {\it filtered GRN}$

Description

Similarly to the statistics produced by plotGeneralGraphStats, summaries regarding the vertex degrees and the most important vertices per community are generated. Note that the communities need to first be calculated using the calculateCommunitiesStats function

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Usage

```
plotCommunitiesStats(
   GRN,
   outputFolder = NULL,
  basenameOutput = NULL,
  display = "byRank",
  communities = seq_len(10),
  topnGenes = 20,
  topnTFs = 20,
  plotAsPDF = TRUE,
  pdf_width = 12,
  pdf_height = 12,
  pages = NULL,
  forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

display Character. Default "byRank". One of: "byRank", "byLabel". Specify whether

the communities will be displayed based on their rank, where the largest community (with most vertices) would have a rank of 1, or by their label. Note that

the label is independent of the rank.

communities Numeric vector. Default seq_len(10). Depending on what was specified in the

display parameter, this parameter would indicate either the rank or the label of the communities to be plotted. i.e. for communities = c(1,4), if display = "byRank" the results for the first and fourth largest communities will be plotted. if display = "byLabel", the results for the communities labeled "1", and "4"

will be plotted. If set to NULL, all communities will be plotted

topnGenes Integer. Default 20. Number of genes to plot, sorted by their rank or label.

topnTFs Integer. Default 20. Number of TFs to plot, sorted by their rank or label.

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, without modifications. A single PDF file is produced with the statistics.

See Also

```
plotGeneralGraphStats
calculateCommunitiesStats
calculateCommunitiesEnrichment
```

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotCommunitiesStats(GRN, plotAsPDF = FALSE)
```

plotDiagnosticPlots_peakGene

Plot diagnostic plots for peak-gene connections for a GRN object

Description

Plot diagnostic plots for peak-gene connections for a GRN object

Usage

```
plotDiagnosticPlots_peakGene(
   GRN,
   outputFolder = NULL,
   basenameOutput = NULL,
   gene.types = list(c("protein_coding", "lincRNA")),
   useFiltered = FALSE,
   plotDetails = FALSE,
   plotPerTF = FALSE,
   plotAsPDF = TRUE,
   pdf_width = 12,
   pdf_height = 12,
   pages = NULL,
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

gene.types List of character vectors. Default list(c("protein coding", "lincRNA")). Vectors

of gene types to consider for the diagnostic plots. Multiple distinct combinations of gene types can be specified. For example, if set to list(c("protein_coding", "lincRNA"), c("protein_coding"), c("all")), 3 distinct PDFs will be produced, one for each element of the list. The first file would only consider protein-coding and lincRNA genes, while the second plot only considers protein-coding ones. The special keyword "all" denotes all gene types found (usually, there are

many gene types present, also more exotic and rare ones).

useFiltered Logical. TRUE or FALSE. Default FALSE. If set to FALSE, the diagnostic plots

will be produced based on all peak-gene connections. This is the default and will usually be best to judge whether the background behaves as expected. If set to TRUE, the diagnostic plots will be produced based on the filtered set of connections. For this, the function link{filterGRNAndConnectGenes} must

have been run before.

plotDetails TRUE or FALSE. Default FALSE. Print additional plots that may help for debug-

ging and QC purposes? Note that these plots are currently less documented or

not at all.

plotPerTF Logical. TRUE or FALSE. Default FALSE. If set to FALSE, the diagnostic plots

will be done across all TF (the default), while setting it to TRUE will generate the QC plots TF-specifically, including "all" TF, sorted by the number of connec-

tions.

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
# GRN = loadExampleObject()
# types = list(c("protein_coding"))
# GRN = plotDiagnosticPlots_peakGene(GRN, gene.types=types, plotAsPDF = FALSE)
```

plotDiagnosticPlots_TFPeaks

Plot diagnostic plots for TF-peak connections for a GRN object

Description

Due to the number of plots that this functions produces, we currently provide only the option to plot as PDF. This may change in the future.

Usage

```
plotDiagnosticPlots_TFPeaks(
   GRN,
   outputFolder = NULL,
   basenameOutput = NULL,
   plotDetails = FALSE,
   dataType = c("real", "permuted"),
   nTFMax = NULL,
   plotAsPDF = TRUE,
   pdf_width = 12,
   pdf_height_base = 8,
   pages = NULL,
   forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

plotDetails TRUE or FALSE. Default FALSE. Print additional plots that may help for debug-

ging and QC purposes? Note that these plots are currently less documented or

not at all.

dataType Character vector. One of, or both of, "real" or "permuted". For which data

type, real or permuted data, to produce the diagnostic plots?

nTFMax NULL or Integer. Default NULL. Maximum number of TFs to process. Can be

used for testing purposes by setting this to a small number i(.e., 10)

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

pdf_height_base

Number. Default 8. Base height of the PDF, in cm, per connection type. The total height is automatically determined based on the number of connection types that are found in the object (e.g., expression or TF activity). For example, when

two connection types are found, the base height is multiplied by 2.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, with added data from this function.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotDiagnosticPlots_TFPeaks(GRN, outputFolder = ".", dataType = "real", nTFMax = 2)
```

plotGeneralEnrichment Plot the general enrichement results

Description

This function plots the results of the general enrichment analysis for every specified ontology.

Usage

```
plotGeneralEnrichment(
   GRN,
   outputFolder = NULL,
   basenameOutput = NULL,
   ontology = NULL,
   topn_pvalue = 30,
   p = 0.05,
```

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```
display_pAdj = FALSE,
maxWidth_nchar_plot = 50,
plotAsPDF = TRUE,
pdf_width = 12,
pdf_height = 12,
pages = NULL,
forceRerun = FALSE
)
```

Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

ontology Character. NULL or vector of ontology names. Default NULL. Vector of ontologies

to plot. The results must have been previously calculated otherwise an error is

thrown.

topn_pvalue Numeric. Default 30. Maximum number of ontology terms that meet the p-value

significance threshold to display in the enrichment dot plot

p Numeric. Default 0.05. p-value threshold to determine significance.

display_pAdj TRUE or FALSE. Default FALSE. Is the p-value being displayed in the plots the

adjusted p-value? This parameter is relevant for KEGG, Disease Ontology, and

Reactome enrichments, and does not affect GO enrichments.

maxWidth_nchar_plot

Integer (>=10). Default 50. Maximum number of characters for a term before it

is truncated.

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm. pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, without modifications. A single PDF file is produced with the results.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotGeneralEnrichment(GRN, plotAsPDF = FALSE)
```

plotGeneralGraphStats Plot general structure and connectivity statistics for a filtered GRN

Description

This function generates graphical summaries about the structure and connectivity of the TF-peak-gene and TF-gene graphs. These include, distribution of vertex types (TF, peak, gene) and edge types (tf-peak, peak-gene), the distribution of vertex degrees, and the most "important" vertices according to degree centrality and eigenvector centrality scores.

Usage

```
plotGeneralGraphStats(
   GRN,
   outputFolder = NULL,
   basenameOutput = NULL,
   plotAsPDF = TRUE,
   pdf_width = 12,
   pdf_height = 12,
   pages = NULL,
   forceRerun = FALSE
)
```

Arguments

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

 $base name \texttt{Output} \quad \texttt{NULL} \ or \ character. \ Default \ \texttt{NULL}. \ Base name \ of \ the \ output \ files \ that \ are \ produced.$

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

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pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object with no changes. The results are output to a file.

See Also

```
plotGeneralEnrichment
plotCommunitiesStats
plotCommunitiesEnrichment
```

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotGeneralGraphStats(GRN, plotAsPDF = FALSE)
```

plotPCA_all

Produce a PCA plot of the data from a GRN object

Description

Produce a PCA plot of the data from a GRN object

Usage

```
plotPCA_all(
   GRN,
   outputFolder = NULL,
   basenameOutput = NULL,
   data = c("rna", "peaks"),
   topn = c(500, 1000, 5000),
   type = c("raw", "normalized"),
   plotAsPDF = TRUE,
   pdf_width = 12,
   pdf_height = 12,
   pages = NULL,
   forceRerun = FALSE
)
```

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Arguments

GRN Object of class GRN

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

data Character. Either "peaks" or "rna" or "all". Default c("rna", "peaks").

Type of data to plot a PCA for. "peaks" corresponds to the the open chromatin data, while "rna" refers to the RNA-seq counts. If set to "all", PCA will be done for both data modalities. In any case, PCA will be based on the original provided data before any additional normalization has been run (i.e., usually the

raw data).

topn Integer vector. Default c(500,1000,5000). Number of top variable features to

do PCA for. Can be a vector of different numbers (see default).

type Character. One of or a combination of "raw", "normalized", "all". Default

c("raw", "normalized"). Should the PCA plots be done based on the raw or

normalized data, respectively?

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.

pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, without modifications. In addition, for each specified type, a PDF file is produced with a PCA. We refer to the Vignettes for details and further explanations.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
```

GRN = loadExampleObject()

GRN = plotPCA_all(GRN, topn = 500, type = "rna", plotAsPDF = FALSE)

plotTFEnrichment 49

plotTFEnrichment

Plot TF-based GO enrichment results

Description

This function plots the enrichment results. The result consist of a dot plot per specified TF, as well as two comparative heatmaps. The first heatmap displays the p value for each GO term across the TFs. Terms that The second heatmap is a subset of the first, where select terms are kept or filtered out for better visibility and display.

Usage

```
plotTFEnrichment(
  GRN,
  rankType = "degree",
  n = NULL,
 TF.names = NULL,
  topn_pvalue = 30,
  p = 0.05,
  nSignificant = 2,
 nID = 10,
  display_pAdj = FALSE,
 maxWidth_nchar_plot = 50,
 outputFolder = NULL,
  basenameOutput = NULL,
  plotAsPDF = TRUE,
  pdf_width = 12,
 pdf_height = 12,
 pages = NULL,
  forceRerun = FALSE
)
```

Arguments

GRN	Object of class GRN
rankType	Character. One of: "degree", "EV", "custom". This parameter will determine the criterion to be used to identify the "top" nodes. If set to "degree", the function will select top nodes based on the number of connections they have, i.e. based on their degree-centrality. If set to "EV" it will select the top nodes based on their eigenvector-centrality score in the network.
n	NULL or numeric. Default NULL. If set to NULL, all previously calculated TF enrichments will be plotted. If set to a value between (0,1), it is treated as a percentage of top nodes. If the value is passed as an integer it will be treated as the number of top nodes. This parameter is not relevant if rankType = "custom".
TF.names	NULL or character vector. Default NULL. For rankType="custom" the names of the TFs to plot. Ignored otherwise.

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topn_pvalue Numeric. Default 30. Maximum number of ontology terms that meet the p-value

significance threshold to display in the enrichment dot plot

p Numeric. Default 0.05. p-value threshold to determine significance.

nSignificant Numeric. Default 3. Threshold to filter out an ontology term with less than

nSignificant overlapping genes.

nID Numeric. Default 10. For the reduced heatmap, number of top terms to select

per community.

display_pAdj TRUE or FALSE. Default FALSE. Is the p-value being displayed in the plots the

adjusted p-value? This parameter is relevant for KEGG, Disease Ontology, and

Reactome enrichments, and does not affect GO enrichments.

maxWidth_nchar_plot

Integer (>=10). Default 50. Maximum number of characters for a term before it

is truncated.

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.
pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

The same GRN object, without modifications. A single PDF file is produced with the results.

See Also

calculateTFEnrichment

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plotTFEnrichment(GRN, n = 5, plotAsPDF = FALSE)
```

```
plot_stats_connectionSummary
```

Plot various network connectivity summaries for a GRN object

Description

Plot various network connectivity summaries for a GRN object

Usage

```
plot_stats_connectionSummary(
   GRN,
   type = "heatmap",
   outputFolder = NULL,
   basenameOutput = NULL,
   plotAsPDF = TRUE,
   pdf_width = 12,
   pdf_height = 12,
   pages = NULL,
   forceRerun = FALSE
)
```

Arguments

type Character. Either "heatmap" or "boxplot". Default "heatmap". Which plot

type to produce?

outputFolder Character or NULL. Default NULL. If set to NULL, the default output folder as spec-

ified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We

recommend specifying an absolute path.

basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm. pdf_height Number. Default 12. Height of the PDF, in cm.

pages Integer vector or NULL. Default NULL. Page number(s) to plot. Can be used to

plot only specific pages to a PDF or the currently active graphics device.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

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Value

The same GRN object, without modifications. In addition, for the specified type, a PDF file (default filename is GRN.connectionSummary_{type}.pdf) is produced with a connection summary.

Examples

```
# See the Workflow vignette on the GRaNIE website for examples
GRN = loadExampleObject()
GRN = plot_stats_connectionSummary(GRN, forceRerun = FALSE, plotAsPDF = FALSE)
```

visualizeGRN

Visualize a filtered GRN.

Description

Visualize a filtered GRN.

Usage

```
visualizeGRN(
  GRN,
  outputFolder = NULL,
  basenameOutput = NULL,
  plotAsPDF = TRUE,
  pdf_width = 12,
  pdf_height = 12,
  title = NULL,
 maxRowsToPlot = 500,
  graph = "TF-gene",
  colorby = "type",
  layout = "fr",
  vertice_color_TFs = list(h = 10, c = 85, l = c(25, 95)),
  vertice\_color\_peaks = list(h = 135, c = 45, l = c(35, 95)),
  vertice_color_genes = list(h = 260, c = 80, 1 = c(30, 90)),
  vertexLabel_cex = 0.4,
  vertexLabel_dist = 0,
  forceRerun = FALSE
)
```

Arguments

GRN

Object of class GRN

outputFolder

Character or NULL. Default NULL. If set to NULL, the default output folder as specified when initiating the object in link{initializeGRN} will be used. Otherwise, all output from this function will be put into the specified folder. We recommend specifying an absolute path.

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basenameOutput NULL or character. Default NULL. Basename of the output files that are produced.

If set to NULL, a default basename is chosen. If a custom basename is specified, all output files will have the chosen basename as file prefix, be careful with not

overwriting already existing files (if forceRerun is set to TRUE)

plotAsPDF TRUE or FALSE. Default TRUE. Should the plots be printed to a PDF file? If set

to TRUE, a PDF file is generated, the name of which depends on the value of basenameOutput. If set to FALSE, all plots are printed to the currently active device. Note that most functions print more than one plot, which means you

may only see the last plot depending on your active graphics device.

pdf_width Number. Default 12. Width of the PDF, in cm.
pdf_height Number. Default 12. Height of the PDF, in cm.

title NULL or Character. Default NULL. Title to be assigned to the plot.

maxRowsToPlot Numeric. Default 500. Refers to the maximum number of connections to be

plotted.

graph Character. Default TF-gene. One of: TF-gene, TF-peak-gene. Whether to plot

a graph with links from TFs to peaks to gene, or the graph with the inferred TF

to gene connections.

colorby Character. Default type. One of type, code community. Color the vertices by

either type (TF/peak/gene) or community. See calculateCommunitiesStats

layout Character. Dfault fr. One of star, fr, sugiyama, kk, lgl, graphopt, mds,

sphere

vertice_color_TFs

Named list. Default list(h = 10, c = 85, l = c(25, 95)). The list must specify the color in hcl format (hue, chroma, luminence). See the colorspace pack-

age for more details and examples

vertice_color_peaks

Named list. Default list(h = 135, c = 45, l = c(35, 95)).

vertice_color_genes

Named list. Default list(h = 260, c = 80, 1 = c(30, 90)).

vertexLabel_cex

Numeric. Default 0.4. Font size (multiplication factor, device-dependent)

vertexLabel_dist

Numeric. Default 0 vertex. Distance between the label and the vertex.

forceRerun TRUE or FALSE. Default FALSE. Force execution, even if the GRN object already

contains the result. Overwrites the old results.

Value

the GRN object

Examples

```
GRN = loadExampleObject()
GRN = visualizeGRN(GRN, maxRowsToPlot = 700, graph = "TF-gene", colorby = "type")
```

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