# Package 'MethReg'

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Type Package

**Title** Assessing the regulatory potential of DNA methylation regions or sites on gene transcription

**Version** 1.19.0

**Description** Epigenome-wide association studies (EWAS) detects a large number of DNA methylation differences, often hundreds of differentially methylated regions and thousands of CpGs, that are significantly associated with a disease, many are located in non-coding regions.

Therefore, there is a critical need to better understand the functional impact of these CpG methylations and to further prioritize the significant changes. MethReg is an R package for integrative modeling of DNA methylation, target gene expression and transcription factor binding sites data, to systematically identify and rank functional CpG methylations. MethReg evaluates, prioritizes and annotates CpG sites with high regulatory potential using matched methylation and gene expression data, along with external TF-target interaction databases based on manually curation, ChIP-seq experiments or gene regulatory network analysis.

**License** GPL-3 **Encoding** UTF-8

LazyData true

Imports dplyr, plyr, GenomicRanges, SummarizedExperiment,
DelayedArray, ggplot2, ggpubr, tibble, tidyr, S4Vectors,
sesameData, sesame, AnnotationHub, ExperimentHub, stringr,
readr, methods, stats, Matrix, MASS, rlang, pscl, IRanges,
sfsmisc, progress, utils, openxlsx, JASPAR2024, RSQLite,
TFBSTools

Suggests rmarkdown, BiocStyle, testthat (>= 2.1.0), parallel, R.utils, doParallel, reshape2, motifmatchr, matrixStats, biomaRt, dorothea, viper, stageR, BiocFileCache, png, htmltools, knitr, jpeg, BSgenome.Hsapiens.UCSC.hg38, BSgenome.Hsapiens.UCSC.hg19, data.table, downloader

VignetteBuilder knitr

BugReports https://github.com/TransBioInfoLab/MethReg/issues/

**RoxygenNote** 7.3.1 **Depends** R (>= 4.0)

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MethReg-package MethReg: functional annotation of DMRs identified in epigenome- wide association studies	-
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# Description

To provide functional annotations for differentially methylated regions (DMRs) and differentially methylated CpG sites (DMS), MethReg performs integrative analyses using matched DNA methylation and gene expression along with Transcription Factor Binding Sites (TFBS) data. MethReg evaluates, prioritizes and annotates DNA methylation regions (or sites) with high regulatory potential that works synergistically with TFs to regulate target gene expressions, without any additional ChIP-seq data.

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#### See Also

Useful links:

• Report bugs at https://github.com/TransBioInfoLab/MethReg/issues/

clinical	TCGA-COAD clinical matrix for 38 samples retrieved from GDC using TCGAbiolinks

# Description

TCGA-COAD clinical matrix for 38 samples retrieved from GDC using TCGAbiolinks

# Usage

clinical

## **Format**

A matrix: 38 samples (rows) and variables (columns) patient, sample, gender and sample\_type

 ${\tt cor\_dnam\_target\_gene} \quad \textit{Evaluate correlation of DNA methylation region and target gene expression}$ 

#### **Description**

This function evaluate the correlation of the DNA methylation and target gene expression using spearman rank correlation test. Note that genes with RNA expression equal to 0 for all samples will not be evaluated.

#### Usage

```
cor_dnam_target_gene(
  pair.dnam.target,
  dnam,
  exp,
  filter.results = TRUE,
  min.cor.pval = 0.05,
  min.cor.estimate = 0,
  cores = 1
)
```

#### **Arguments**

pair.dnam.target

A dataframe with the following columns: regionID (DNA methylation) and tar-

get (target gene)

dnam DNA methylation matrix or SummarizedExperiment object with regions/cpgs in

rows and samples in columns are samples. Samples should be in the same order

as gene expression matrix (exp).

exp Gene expression matrix or SummarizedExperiment object (rows are genes, columns

are samples) log2-normalized (log2(exp + 1)). Samples should be in the same

order as the DNA methylation matrix.

filter.results Filter results using min.cor.pval and min.cor.estimate thresholds

min.cor.pval P-value threshold filter (default: 0.05)

min.cor.estimate

Correlation estimate threshold filter (default: not applied)

cores Number of CPU cores to be used. Default 1.

# Value

A data frame with the following information: regionID, target gene, correlation pvalue and estimate between DNA methylation and target gene expression, FDR corrected p-values.

```
\begin{array}{lll} dnam <- t(matrix(sort(c(runif(20))), ncol = 1)) \\ rownames(dnam) <- c("chr3:203727581-203728580") \\ colnames(dnam) <- paste0("Samples",1:20) \\ exp <- dnam \end{array}
```

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```
rownames(exp) <- c("ENSG00000232886")
colnames(exp) <- paste0("Samples",1:20)

pair.dnam.target <- data.frame(
    "regionID" = c("chr3:203727581-203728580"),
    "target" = "ENSG00000232886"
)

# Correlated DNAm and gene expression, display only significant associations
results.cor.pos <- cor_dnam_target_gene(
    pair.dnam.target = pair.dnam.target,
    dnam = dnam,
    exp = exp,
    filter.results = TRUE,
    min.cor.pval = 0.05,
    min.cor.estimate = 0.0
)</pre>
```

cor\_tf\_target\_gene

Evaluate correlation of TF expression and target gene expression

#### **Description**

This function evaluate the correlation of a TF and target gene expression using spearman rank correlation test. Note that genes with RNA expression equal to 0 for all samples will not be evaluated.

# Usage

```
cor_tf_target_gene(
  pair.tf.target,
  exp,
  tf.activity.es = NULL,
  cores = 1,
  verbose = FALSE
)
```

## **Arguments**

pair.tf.target	A dataframe with the following columns: TF and target (target gene)
exp	Gene expression matrix or SummarizedExperiment object (rows are genes, columns are samples) log2-normalized (log2(exp + 1)). Samples should be in the same order as the $tf$ .activity.es matrix
tf.activity.es	A matrix with normalized enrichment scores for each TF across all samples to be used in linear models instead of TF gene expression. See get_tf_ES.
cores	Number of CPU cores to be used. Default 1.
verbose	Show messages ?

# Value

A data frame with the following information: TF, target gene, correlation p-value and estimate between TF and target gene expression, FDR corrected p-values.

#### **Examples**

```
exp <- t(matrix(sort(c(runif(40))), ncol = 2))
rownames(exp) <- c("ENSG00000232886","ENSG00000232889")
colnames(exp) <- paste0("Samples",1:20)

pair.tf.target <- data.frame(
    "TF" = "ENSG00000232889",
    "target" = "ENSG00000232886"
)

# Correlated TF and gene expression
results.cor.pos <- cor_tf_target_gene(
    pair.tf.target = pair.tf.target,
    exp = exp,
)
# Correlated TF and gene expression
results.cor.pos <- cor_tf_target_gene(
    pair.tf.target = pair.tf.target,
    exp = exp,
    tf.activity.es = exp
)</pre>
```

create\_triplet\_distance\_based

Map DNAm to target genes using distance approaches, and TF to the DNAm region using JASPAR2024 TFBS.

# Description

This function wraps two other functions get\_region\_target\_gene and get\_tf\_in\_region from the package. This function will map a region to a target gene using three methods (mapping to the closest gene, mapping to any gene within a given window of distance, or mapping to a fixed number of nearby genes upstream or downstream). To find TFs binding to the region, JASPAR2024 is used.

# Usage

```
create_triplet_distance_based(
    region,
    genome = c("hg38", "hg19"),
    target.method = c("genes.promoter.overlap", "window", "nearby.genes", "closest.gene"),
    target.window.size = 500 * 10^3,
    target.num.flanking.genes = 5,
    target.promoter.upstream.dist.tss = 2000,
    target.promoter.downstream.dist.tss = 2000,
    target.rm.promoter.regions.from.distal.linking = TRUE,
    motif.search.window.size = 0,
    motif.search.p.cutoff = 1e-08,
    TF.peaks.gr = NULL,
    max.distance.region.target = 10^6,
    cores = 1
)
```

#### **Arguments**

region A Granges or a named vector with regions (i.e "chr21:100002-1004000")

genome Human genome reference "hg38" or "hg19"

target.method How genes are mapped to regions: regions overlapping gene promoter ("genes.promoter.overlap");

genes within a window around the region ("window"); or fixed number of nearby

genes upstream and downstream from the region

target.window.size

When method = "window", number of base pairs to extend the region (+- window.size/2). Default is 500kbp (or +/- 250kbp, i.e. 250k bp from start or end of

the region)

target.num.flanking.genes

Number of flanking genes upstream and downstream to search. For example, if target.num.flanking.genes = 5, it will return the 5 genes upstream and 5 genes downstream

target.promoter.upstream.dist.tss

Number of base pairs (bp) upstream of TSS to consider as promoter regions. Defaults to 2000 bp.

target.promoter.downstream.dist.tss

Number of base pairs (bp) downstream of TSS to consider as promoter regions. Defaults to 2000 bp.

target.rm.promoter.regions.from.distal.linking

When performing distal linking with method = "windows" or method = "nearby.genes", or "closest.gene.tss", if set to TRUE (default), probes in promoter regions will be removed from the input.

motif.search.window.size

Integer value to extend the regions. For example, a value of 50 will extend 25 bp upstream and 25 downstream the region. Default is no increase

motif.search.p.cutoff

motifmatchr pvalue cut-off. Default 1e-8.

TF.peaks.gr A granges with

A granges with TF peaks to be overlaped with input region Metadata column expected "id" with TF name. Default NULL. Note that Remap catalog can be used as shown in the examples.

max.distance.region.target

Max distance between region and target gene. Default 1Mbp.

cores Number of CPU cores to be used. Default 1.

# Value

A data frame with TF, target and RegionID information.

```
regions.names <- c("chr3:189631389-189632889","chr4:43162098-43163498")
triplet <- create_triplet_distance_based(
   region = regions.names,
   motif.search.window.size = 500,
   target.method = "closest.gene"
)</pre>
```

```
create_triplet_regulon_based
```

Map TF and target genes using regulon databases or any user provided target-tf. Maps TF to the DNAm region with TFBS using JAS-PAR2020 TFBS.

#### **Description**

This function wraps two other functions get\_region\_target\_gene and get\_tf\_in\_region from the package.

#### Usage

```
create_triplet_regulon_based(
  region,
  genome = c("hg38", "hg19"),
  regulons.min.confidence = "B",
 motif.search.window.size = 0,
 motif.search.p.cutoff = 1e-08,
  cores = 1,
  tf.target,
 TF.peaks.gr = NULL,
 max.distance.region.target = 10^6
)
```

# **Arguments**

region A Granges or a named vector with regions (i.e "chr21:100002-1004000")

genome Human genome reference "hg38" or "hg19"

regulons.min.confidence

Minimun confidence score ("A", "B", "C", "D", "E") classifying regulons based on their quality from Human DoRothEA database dorothea\_hs. The default

minimun confidence score is "B".

motif.search.window.size

Integer value to extend the regions. For example, a value of 50 will extend 25 bp upstream and 25 downstream the region. Default is no increase

motif.search.p.cutoff

motifmatchr pvalue cut-off. Default 1e-8.

Number of CPU cores to be used. Default 1. cores

A dataframe with tf and target columns. If not provided, dorothea\_hs will be tf.target

TF.peaks.gr A granges with TF peaks to be overlaped with input region Metadata column

expected "id" with TF name. Default NULL. Note that Remap catalog can be

used as shown in the examples.

max.distance.region.target

Max distance between region and target gene. Default 1Mbp.

#### Value

A data frame with TF, target and RegionID information.

dna.met.chr21

## **Examples**

```
triplet <- create_triplet_regulon_based(
    region = c("chr1:69591-69592", "chr1:898803-898804"),
    motif.search.window.size = 50,
    regulons.min.confidence = "B",
        motif.search.p.cutoff = 0.05
)</pre>
```

dna.met.chr21

TCGA-COAD DNA methylation matrix (beta-values) for 38 samples (only chr21) retrieved from GDC using TCGAbiolinks

# **Description**

TCGA-COAD DNA methylation matrix (beta-values) for 38 samples (only chr21) retrieved from GDC using TCGAbiolinks

# Usage

dna.met.chr21

#### **Format**

A beta-value matrix with 38 samples, includes CpG IDs in the rows and TCGA sample identifiers in the columns

```
export_results_to_table
```

Format MethReg results table and export to XLSX file

# **Description**

Receives a methReg results table and create a formatted XLSX file to easier readability and interpretation of the results

# Usage

```
export_results_to_table(results, file = "MethReg_results.xlsx")
```

# Arguments

results MethReg results

file xlsx filename used to save

# Value

A summarized Experiment object

#### **Examples**

```
library(dplyr)
dnam <- runif(20,min = 0,max = 1) %>%
     matrix(ncol = 1) \%\% t
rownames(dnam) <- c("chr3:203727581-203728580")
colnames(dnam) <- paste0("Samples",1:20)</pre>
exp.target <- runif(20,min = 0,max = 10) \%
    matrix(ncol = 1) %>% t
rownames(exp.target) <- c("ENSG00000252982")</pre>
colnames(exp.target) <- paste0("Samples",1:20)</pre>
exp.tf <- runif(20,min = 0,max = 10) %>%
    matrix(ncol = 1) %>% t
rownames(exp.tf) <- c("ENSG00000083937")
colnames(exp.tf) <- paste0("Samples",1:20)</pre>
exp <- rbind(exp.tf, exp.target)</pre>
triplet <- data.frame(</pre>
        "regionID" = c("chr3:203727581-203728580"),
        "target" = "ENSG00000252982",
        "TF" = "ENSG00000083937"
results <- interaction_model(</pre>
        triplet = triplet,
        dnam = dnam,
        exp = exp,
         dnam.group.threshold = 0.25,
        stage.wise.analysis = FALSE,
        sig.threshold = 1,
        filter.correlated.tf.exp.dnam = FALSE,
        filter.correlated.target.exp.dnam = FALSE,
        filter.triplet.by.sig.term = FALSE
results <- results %>% stratified_model( dnam = dnam, exp = exp)
export_results_to_table(results = results, file = "MethReg_results.xlsx")
results\RLM_DNAmGroup:TF_region_stage_wise_adj_pvalue\ <- results\RLM_DNAmGroup:TF_fdr\
results \verb|`RLM_DNAmGroup:TF_triplet_stage_wise_adj_pvalue`| <- results \verb|`RLM_DNAmGroup:TF_fdr`| <- results | RLM_DNAmGroup:TF_fdr'| <- results | RLM_D
results$`RLM_DNAmGroup:TF_fdr` <- NULL</pre>
export_results_to_table(results = results, file = "MethReg_results_stage_wise.xlsx")
```

```
filter_dnam_by_quant_diff
```

Select regions with variations in DNA methylation levels above a threshold

## **Description**

For each region, computes the interquartile range (IQR) of the DNA methylation (DNAm) levels and requires the IQR to be above a threshold

#### Usage

```
filter_dnam_by_quant_diff(dnam, min.IQR.threshold = 0.2, cores = 1)
```

#### **Arguments**

 $\begin{tabular}{ll} $DNA$ methylation matrix or Summarized Experiment object \\ min.IQR.threshold \end{tabular}$ 

Threshold for minimal interquantile range (difference between the 75th and 25th

percentiles) of the DNAm

cores Number of CPU cores to be used in the analysis. Default: 1

#### Value

A subset of the original matrix only with the rows passing the filter threshold.

## **Examples**

```
data("dna.met.chr21")
dna.met.chr21.filtered <- filter_dnam_by_quant_diff(
   dna.met.chr21
)</pre>
```

```
filter\_exp\_by\_quant\_mean\_FC
```

Select genes with variations above a threshold

# **Description**

For each gene, compares the mean gene expression levels in samples in high expression (Q4) vs. samples with low gene expression (Q1), and requires the fold change to be above a certain threshold.

## Usage

```
filter_exp_by_quant_mean_FC(exp, fold.change = 1.5, cores = 1)
```

#### **Arguments**

exp Gene expression matrix or SumarizedExperiment object

fold. change Threshold for fold change of mean gene expression levels in samples with high

(Q4) and low (Q1) gene expression levels. Defaults to 1.5.

cores Number of CPU cores to be used in the analysis. Default: 1

#### Value

A subset of the original matrix only with the rows passing the filter threshold.

```
data("gene.exp.chr21.log2")
gene.exp.chr21.log2.filtered <- filter_exp_by_quant_mean_FC(
   gene.exp.chr21.log2
)</pre>
```

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```
filter_genes_zero_expression
```

Remove genes with gene expression level equal to 0 in a substantial percentage of the samples

#### **Description**

Remove genes with gene expression level equal to 0 in a substantial percentage of the samples

# Usage

```
filter_genes_zero_expression(exp, max.samples.percentage = 0.25)
```

#### **Arguments**

exp Gene expression matrix or SumarizedExperiment object max.samples.percentage

Max percentage of samples with gene expression as 0, for genes to be selected. If max.samples.percentage 100, remove genes with 0 for 100% samples. If max.samples.percentage 25, remove genes with 0 for more than 25% of the samples.

#### Value

A subset of the original matrix only with the rows passing the filter threshold.

```
gene.exp.chr21.log2 TCGA-COAD gene expression matrix (log2 (FPKM-UQ + 1)) for 38 samples (only chromosome 21) retrieved from GDC using TCGAbiolinks
```

# **Description**

TCGA-COAD gene expression matrix (log2 (FPKM-UQ + 1)) for 38 samples (only chromosome 21) retrieved from GDC using TCGAbiolinks

# Usage

```
gene.exp.chr21.log2
```

# **Format**

A log2 (FPKM-UQ + 1) gene expression matrix with 38 samples, includes Ensembl IDs in the rows and TCGA sample identifiers in the columns

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 $get\_human\_tfs$ 

Access human TF from Lambert et al 2018

# **Description**

Access human TF from Lambert et al 2018 (PMID: 29425488)

# Usage

```
get_human_tfs()
```

#### Value

A dataframe with Human TF

# **Examples**

```
human.tfs <- get_human_tfs()</pre>
```

get\_met\_probes\_info

Get HM450/EPIC manifest files from Sesame package

# Description

Returns a data frame with HM450/EPIC manifest information files from Sesame package

# Usage

```
get_met_probes_info(genome = c("hg38", "hg19"), arrayType = c("450k", "EPIC"))
```

# **Arguments**

genome Human genome of reference hg38 or hg19

arrayType "450k" or "EPIC" array

# Value

A Granges with the DNAm array manifest

```
regions.names <- c("chr22:18267969-18268249","chr23:18267969-18268249")
regions.gr <- make_granges_from_names(regions.names)
make_names_from_granges(regions.gr)</pre>
```

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get\_promoter\_avg

Summarize promoter DNA methylation beta values by mean.

# **Description**

First, identify gene promoter regions (default +-2Kkb around TSS). Then, for each promoter region calculate the mean DNA methylation of probes overlapping the region.

# Usage

```
get_promoter_avg(
  dnam,
  genome,
  arrayType,
  cores = 1,
  upstream.dist.tss = 2000,
  downstream.dist.tss = 2000,
  verbose = FALSE
)
```

# **Arguments**

dnam A DNA methylation matrix or a SummarizedExperiment object

genome Human genome of reference hg38 or hg19 arrayType DNA methylation array type (450k or EPIC)

cores A integer number to use multiple cores. Default 1 core.

upstream.dist.tss

Number of base pairs (bp) upstream of TSS to consider as promoter regions

downstream.dist.tss

Number of base pairs (bp) downstream of TSS to consider as promoter regions

verbose A logical argument indicating if messages output should be provided.

## Value

A RangedSummarizedExperiment with promoter region and mean beta-values of CpGs within it. Metadata will provide the promoter gene region and gene informations.

```
## Not run:
    data("dna.met.chr21")
    promoter.avg <- get_promoter_avg(
        dnam = dna.met.chr21,
        genome = "hg19",
        arrayType = "450k"
)
## End(Not run)</pre>
```

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```
get_region_target_gene
```

Obtain target genes of input regions based on distance

#### **Description**

To map an input region to genes there are three options: 1) map region to closest gene tss 2) map region to all genes within a window around the region (default window.size = 500kbp (i.e. +/- 250kbp from start or end of the region)). 3) map region to a fixed number of nearby genes (upstream/downstream)

## Usage

```
get_region_target_gene(
  regions.gr,
  genome = c("hg38", "hg19"),
  method = c("genes.promoter.overlap", "window", "nearby.genes", "closest.gene.tss"),
  promoter.upstream.dist.tss = 2000,
  promoter.downstream.dist.tss = 2000,
  window.size = 500 * 10^3,
  num.flanking.genes = 5,
  rm.promoter.regions.from.distal.linking = TRUE
)
```

# **Arguments**

regions.gr A Genomic Ranges object (GRanges) or a SummarizedExperiment object (rowRanges

will be used)

genome Human genome of reference "hg38" or "hg19"

method How genes are mapped to regions: region overlapping gene promoter ("genes.promoter.overlap");

or genes within a window around the region ("window"); or a fixed number genes upstream and downstream of the region ("nearby.genes"); or closest gene

tss to the region ("closest.gene.tss")

promoter.upstream.dist.tss

Number of base pairs (bp) upstream of TSS to consider as promoter regions.

Defaults to 2000 bp.

promoter.downstream.dist.tss

Number of base pairs (bp) downstream of TSS to consider as promoter regions.

Defaults to 2000 bp.

window.size

When method = "window", number of base pairs to extend the region (+- win-

dow.size/2). Default is 500kbp (or +/- 250kbp, i.e. 250k bp from start or end of the region)

the re

num.flanking.genes

When method = "nearby.genes", set the number of flanking genes upstream and downstream to search.Defaults to 5. For example, if num.flanking.genes = 5, it will return the 5 genes upstream and 5 genes downstream of the given

region.

rm.promoter.regions.from.distal.linking

When performing distal linking with method = "windows", "nearby.genes" or "closest.gene.tss", if set to TRUE (default), probes in promoter regions will be removed from the input.

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#### **Details**

For the analysis of probes in promoter regions (promoter analysis), we recommend setting method = "genes.promoter.overlap".

For the analysis of probes in distal regions (distal analysis), we recommend setting either method = "window" or method = "nearby.genes".

Note that because method = "window" or method = "nearby.genes" are mainly used for analyzing distal probes, by default rm.promoter.regions.from.distal.linking = TRUE to remove probes in promoter regions.

# Value

A data frame with the following information: regionID, Target symbol, Target ensembl ID

```
library(GenomicRanges)
library(dplyr)
# Create example region
regions.gr <- data.frame(</pre>
       chrom = c("chr22", "chr22", "chr22", "chr22"),
       start = c("39377790", "50987294", "19746156", "42470063", "43817258"),
       end = c("39377930", "50987527", "19746368", "42470223", "43817384"),
       stringsAsFactors = FALSE) %>%
     {\tt makeGRangesFromDataFrame}
 # map to closest gene tss
 region.genes.promoter.overlaps <- get_region_target_gene(</pre>
                       regions.gr = regions.gr,
                       genome = "hg19",
                       method = "genes.promoter.overlap"
 )
 # map to all gene within region +- 250kbp
 region.window.genes <- get_region_target_gene(</pre>
                       regions.gr = regions.gr,
                       genome = "hg19",
                       method = "window"
                       window.size = 500 \times 10^3
 )
 # map regions to n upstream and n downstream genes
 region.nearby.genes <- get_region_target_gene(</pre>
                       regions.gr = regions.gr,
                       genome = "hg19",
                       method = "nearby.genes",
                       num.flanking.genes = 5
 )
```

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#### **Description**

Compute studentized residuals from fitting linear regression models to expression values in a data matrix

## Usage

```
get_residuals(data, metadata.samples = NULL, metadata.genes = NULL, cores = 1)
```

#### **Arguments**

data

A matrix or SummarizedExperiment object with samples as columns and features (gene, probes) as rows. Note that expression values should typically be log2(expx + 1) transformed before fitting linear regression models.

metadata.samples

A data frame with samples as rows and columns the covariates. No NA values are allowed, otherwise residual of the corresponding sample will be NA.

metadata.genes A data frame with genes (covariates) as rows and samples as columns. For each evaluated gene, each column (e.g. CNA) that corresponds to the same gene will be set as a single covariate variable. This can be used to correct copy number alterations for each gene.

cores

Number of CPU cores to be used. Defaults to 1.

#### **Details**

When only metadata. samples are provided, this function computes residuals for expression values in a data matrix by fitting model

features ~ Sample\_covariate1 + Sample\_covariate2 . . . + Sample\_covariateN where N is the index of the columns in the metadata provided, features are (typically log transformed) expression values.

When the user additionally provide metadata genes, that is, gene metadata (e.g. gene\_covariate = copy number variations/alterations) residuals are computed by fitting the following model:

features ~ Sample\_covariate1 + Sample\_covariate2 ... + Sample\_covariateN + gene\_covariate

#### Value

A residuals matrix with samples as columns and features (gene, probes) as rows

```
data("gene.exp.chr21.log2")
data("clinical")
metadata <- clinical[,c( "gender", "sample_type")]</pre>
cnv <- matrix(</pre>
   sample(x = c(-2,-1,0,1,2),
   size = ncol(gene.exp.chr21.log2) * nrow(gene.exp.chr21.log2),replace = TRUE),
   nrow = nrow(gene.exp.chr21.log2),
   ncol = ncol(gene.exp.chr21.log2)
rownames(cnv) <- rownames(gene.exp.chr21.log2)</pre>
colnames(cnv) <- colnames(gene.exp.chr21.log2)</pre>
```

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```
gene.exp.residuals <- get_residuals(
   data = gene.exp.chr21.log2[1:3,],
   metadata.samples = metadata,
   metadata.genes = cnv
)
gene.exp.residuals <- get_residuals(
   data = gene.exp.chr21.log2[1:3,],
   metadata.samples = metadata,
   metadata.genes = cnv[1:2,]
)
gene.exp.residuals <- get_residuals(
   data = gene.exp.chr21.log2[1:3,],
   metadata.samples = metadata
)</pre>
```

get\_tf\_ES

Calculate enrichment scores for each TF across all samples using dorothea and viper.

## **Description**

Calculate enrichment scores for each TF across all samples using dorothea and viper.

# Usage

```
get_tf_ES(exp, min.confidence = "B", regulons)
```

# **Arguments**

exp Gene expression matrix with gene expression counts, row as ENSG gene IDS

and column as samples

min.confidence Minimun confidence score ("A", "B", "C", "D", "E") classifying regulons based

on their quality from Human DoRothEA database. The default minimun confi-

dence score is "B"

regulons DoRothEA regulons in table format. Same as run\_viper. If not specified Bio-

conductor (human) dorothea regulons besed on GTEx will be. used dorothea\_hs.

# Value

A matrix of normalized enrichment scores for each TF across all samples

```
gene.exp.chr21.log2 <- get(data("gene.exp.chr21.log2"))
tf_es <- get_tf_ES(gene.exp.chr21.log2)</pre>
```

get\_tf\_in\_region 19

get_tf_in_region	Get human TFs for regions by either scanning it with motifmatchr us-
	ing JASPAR 2024 database or overlapping with TF chip-seq from user
	input

#### **Description**

Given a genomic region, this function maps TF in regions using two methods: 1) using motifmatchr nd JASPAR 2024 to scan the region for 554 human transcription factors binding sites. There is also an option (argument window.size) to extend the scanning region before performing the search, which by default is 0 (do not extend). 2) Using user input TF chip-seq to check for overlaps between region and TF peaks.

# Usage

```
get_tf_in_region(
  region,
  window.size = 0,
  genome = c("hg19", "hg38"),
  p.cutoff = 1e-08,
  cores = 1,
  TF.peaks.gr = NULL,
  verbose = FALSE
)
```

## **Arguments**

region	A vector of region names or	GRanges object with the	DNA methylation regions
--------	-----------------------------	-------------------------	-------------------------

to be scanned for the motifs

window.size Integer value to extend the regions. For example, a value of 50 will extend 25

bp upstream and 25 bp downstream the region. The default is not to increase the

scanned region.

genome Human genome of reference "hg38" or "hg19".

p. cutoff motifmatchr p.cutoff. Default 1e-8.

cores Number of CPU cores to be used. Default 1.

TF.peaks.gr A granges with TF peaks to be overlaped with input region Metadata column

expected "id" with TF name. Default NULL. Note that Remap catalog can be

used as shown in the examples.

verbose A logical argument indicating if messages output should be provided.

#### Value

A data frame with the following information: regionID, TF symbol, TF ensembl ID

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```
)
## Not run:
   library(ReMapEnrich)
   demo.dir <- "~/ReMapEnrich_demo"</pre>
   dir.create(demo.dir, showWarnings = FALSE, recursive = TRUE)
   # Use the function DowloadRemapCatalog
   remapCatalog2018hg38 <- downloadRemapCatalog(demo.dir, assembly = "hg38")</pre>
   # Load the ReMap catalogue and convert it to Genomic Ranges
   remapCatalog <- bedToGranges(remapCatalog2018hg38)</pre>
   regions.names <- c("chr3:189631389-189632889", "chr4:43162098-43163498")
   region.tf.remap <- get_tf_in_region(</pre>
                    region = regions.names,
                    genome = "hg38",
                    TF.peaks.gr = remapCatalog
   )
## End(Not run)
```

interaction\_model

Fits linear models with interaction to triplet data (Target, TF, DNAm), where DNAm is a binary variable (samples in Q1 or Q4)

## **Description**

Evaluates regulatory potential of DNA methylation (DNAm) on gene expression, by fitting robust linear model or zero inflated negative binomial model to triplet data. These models consist of terms to model direct effect of DNAm on target gene expression, direct effect of TF on gene expression, as well as an interaction term that evaluates the synergistic effect of DNAm and TF on gene expression.

# Usage

```
interaction_model(
    triplet,
    dnam,
    exp,
    dnam.group.threshold = 0.25,
    cores = 1,
    tf.activity.es = NULL,
    sig.threshold = 0.05,
    fdr = TRUE,
    filter.correlated.tf.exp.dnam = TRUE,
    filter.triplet.by.sig.term = TRUE,
    stage.wise.analysis = TRUE,
    verbose = FALSE
```

## **Arguments**

triplet

Data frame with columns for DNA methylation region (regionID), TF (TF), and target gene (target)

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dnam DNA methylation matrix or SummarizedExperiment object (columns: samples

in the same order as exp matrix, rows: regions/probes)

exp A matrix or SummarizedExperiment object object (columns: samples in the

same order as dnam, rows: genes represented by ensembl IDs (e.g. ENSG00000239415))

dnam.group.threshold

DNA methylation threshold percentage to define samples in the low methylated group and high methylated group. For example, setting the threshold to 0.3 (30%) will assign samples with the lowest 30% methylation in the low group and the highest 30% methylation in the high group. Default is 0.25 (25%),

accepted threshold range (0.0,0.5].

cores Number of CPU cores to be used. Default 1.

 $\hbox{tf.activity.es} \quad A \ matrix \ with \ normalized \ enrichment \ scores \ for \ each \ TF \ across \ all \ samples \ to$ 

be used in linear models instead of TF gene expression. See get\_tf\_ES.

sig. threshold Threshold to filter significant triplets. Select if interaction.pval < 0.05 or pval.dnam

< 0.05 or pval.tf < 0.05 in binary model

fdr Uses fdr when using sig.threshold. Select if interaction.fdr < 0.05 or fdr.dnam <

0.05 or fdr.tf < 0.05 in binary model

filter.correlated.tf.exp.dnam

If wilcoxon test of TF expression Q1 and Q4 is significant (pvalue < 0.05), triplet

will be removed.

filter.correlated.target.exp.dnam

If wilcoxon test of target expression Q1 and Q4 is not significant (pvalue > 0.05),

triplet will be removed.

filter.triplet.by.sig.term

Filter significant triplets ? Select if interaction.pval < 0.05 or pval.dnam < 0.05

or pval.tf < 0.05 in binary model

stage.wise.analysis

A boolean indicating if stagewise analysis should be performed to correct for

multiple comparisons. If set to FALSE FDR analysis is performed.

verbose A logical argument indicating if messages output should be provided.

# Details

This function fits the linear model

 $log2(RNA target) \sim log2(TF) + DNAm + log2(TF) * DNAm$ 

to triplet data as follow:

Model by considering DNAm as a binary variable - we defined a binary group for DNA methylation values (high = 1, low = 0). That is, samples with the highest DNAm levels (top 25 percent) has high = 1, samples with lowest DNAm levels (bottom 25 percent) has high = 0. Note that in this implementation, only samples with DNAm values in the first and last quartiles are considered.

In these models, the term log2(TF) evaluates direct effect of TF on target gene expression, DNAm evaluates direct effect of DNAm on target gene expression, and log2(TF)\*DNAm evaluates synergistic effect of DNAm and TF, that is, if TF regulatory activity is modified by DNAm.

There are two implementations of these models, depending on whether there are an excessive amount (i.e. more than 25 percent) of samples with zero counts in RNAseq data:

• When percent of zeros in RNAseq data is less than 25 percent, robust linear models are implemented using rlm function from MASS package. This gives outlier gene expression values reduced weight. We used "psi.bisqure" option in function rlm (bisquare weighting, https://stats.idre.ucla.edu/r/dae/robust-regression/).

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• When percent of zeros in RNAseq data is more than 25 percent, zero inflated negative binomial models are implemented using zeroinfl function from pscl package. This assumes there are two processes that generated zeros (1) one where the counts are always zero (2) another where the count follows a negative binomial distribution.

To account for confounding effects from covariate variables, first use the get\_residuals function to obtain RNA or DNAm residual values which have covariate effects removed, then fit interaction model. Note that no log2 transformation is needed when interaction\_model is applied to residuals data.

Note that only triplets with TF expression not significantly different in high vs. low methylation groups will be evaluated (Wilcoxon test, p > 0.05).

#### Value

A dataframe with Region, TF, target, TF\_symbo, target\_symbol, estimates and P-values, after fitting robust linear models or zero-inflated negative binomial models (see Details above).

Model considering DNAm values as a binary variable generates quant\_pval\_metGrp, quant\_pval\_rna.tf, quant\_estimates\_metGrp, quant\_estimates\_rna.tf, quant\_estimates\_metGrp.rna.tf, quant\_es

Model.interaction indicates which model (robust linear model or zero inflated model) was used to fit Model 1, and Model.quantile indicates which model(robust linear model or zero inflated model) was used to fit Model 2.

```
library(dplyr)
dnam <- runif(20,min = 0,max = 1) %>%
  matrix(ncol = 1) %>% t
rownames(dnam) <- c("chr3:203727581-203728580")
colnames(dnam) <- paste0("Samples",1:20)</pre>
exp.target <- runif(20,min = 0,max = 10) %>%
  matrix(ncol = 1) %>% t
rownames(exp.target) <- c("ENSG00000252982")</pre>
colnames(exp.target) <- paste0("Samples",1:20)</pre>
exp.tf <- runif(20,min = 0,max = 10) %>%
  matrix(ncol = 1) %>% t
\texttt{rownames}(\texttt{exp.tf}) \mathrel{<\!\!\!-} \texttt{c("ENSG00000083937")}
colnames(exp.tf) <- paste0("Samples",1:20)</pre>
exp <- rbind(exp.tf, exp.target)</pre>
triplet <- data.frame(</pre>
   "regionID" = c("chr3:203727581-203728580"),
   "target" = "ENSG00000252982",
   "TF" = "ENSG00000083937"
results <- interaction_model(</pre>
   triplet = triplet,
   dnam = dnam,
   exp = exp,
    dnam.group.threshold = 0.25,
   stage.wise.analysis = FALSE,
   sig.threshold = 1,
   filter.correlated.tf.exp.dnam = FALSE,
```

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```
filter.correlated.target.exp.dnam = FALSE,
filter.triplet.by.sig.term = FALSE
)
```

make\_dnam\_se

Transform DNA methylation array into a summarized Experiment object

# Description

Transform DNA methylation array into a summarized Experiment object

# Usage

```
make_dnam_se(
  dnam,
  genome = c("hg38", "hg19"),
  arrayType = c("450k", "EPIC"),
  betaToM = FALSE,
  verbose = FALSE
)
```

# **Arguments**

dnam	DNA methylation matrix with beta-values or m-values as data, row as cpgs
	"cg07946458" or regions ("chr1:232:245") and column as samples

genome Human genome of reference: hg38 or hg19 arrayType DNA methylation array type (450k or EPIC)

betaToM indicates if converting methylation beta values to mvalues

verbose A logical argument indicating if messages output should be provided.

# Value

A summarized Experiment object with DNA methylation probes mapped to genomic regions

```
library(dplyr)
dnam <- runif(20, min = 0,max = 1) %>% sort %>%
  matrix(ncol = 1) %>% t
rownames(dnam) <- c("chr3:203727581-203728580")
colnames(dnam) <- paste0("Samples",1:20)
  se <- make_dnam_se(dnam)</pre>
```

make_exp_se	Transform gene expression matrix into a Summarized Experiment object
-------------	--

# **Description**

Transform gene expression matrix into a Summarized Experiment object

#### Usage

```
make_exp_se(exp, genome = c("hg38", "hg19"), verbose = FALSE)
```

#### **Arguments**

exp Gene expression matrix with gene expression counts, row as ENSG gene IDS

and column as samples

genome Human genome of reference: hg38 or hg19

verbose A logical argument indicating if messages output should be provided.

#### Value

A summarized Experiment object

#### **Examples**

```
gene.exp.chr21.log2 <- get(data("gene.exp.chr21.log2"))
gene.exp.chr21.log2.se <- make_exp_se(gene.exp.chr21.log2)</pre>
```

```
make_granges_from_names
```

Create a Granges object from a genmic region string

# Description

Given a region name such as chr22:18267969-18268249, we will create a Granges object

#### Usage

```
make_granges_from_names(names)
```

# Arguments

names

A region name as "chr22:18267969-18268249" or a vector of region names.

#### Value

A GRanges

```
regions.names <- c("chr22:18267969-18268249","chr23:18267969-18268249")
regions.gr <- make_granges_from_names(regions.names)</pre>
```

```
make_names_from_granges
```

Create region name from Granges

# **Description**

Given a GRanges returns region name such as chr22:18267969-18268249

# Usage

```
make_names_from_granges(region)
```

# **Arguments**

region

A GenomicRanges object

#### Value

A string

# **Examples**

```
regions.names <- c("chr22:18267969-18268249","chr23:18267969-18268249")
regions.gr <- make_granges_from_names(regions.names)
make_names_from_granges(regions.gr)</pre>
```

methReg\_analysis

Wrapper for MethReg functions

# Description

Wrapper for the following MethReg functions: 1) DNAm vs Target gene spearman correlation 2) TF vs Target gene spearman correlation 3) interaction\_model 4) stratified model

# Usage

```
methReg_analysis(
    triplet,
    dnam,
    exp,
    tf.activity.es = NULL,
    dnam.group.percent.threshold = 0.25,
    perform.correlation.analaysis = TRUE,
    remove.nonsig.correlated.dnam.target.gene = FALSE,
    remove.nonsig.correlated.dnam.target.gene.threshold.pvalue = 0.01,
    remove.nonsig.correlated.dnam.target.gene.threshold.estimate = 0.2,
    remove.sig.correlated.tf.exp.dnam = TRUE,
    filter.triplet.by.sig.term = TRUE,
    filter.triplet.by.sig.term.using.fdr = TRUE,
    filter.triplet.by.sig.term.pvalue.threshold = 0.05,
```

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```
multiple.correction.by.stage.wise.analysis = TRUE,
  tf.dnam.classifier.pval.threshold = 0.001,
  verbose = FALSE,
  cores = 1
)
```

#### **Arguments**

triplet Data frame with columns for DNA methylation region (regionID), TF (TF), and

target gene (target)

dnam DNA methylation matrix or SummarizedExperiment object (columns: samples

in the same order as exp matrix, rows: regions/probes)

exp A matrix or SummarizedExperiment object object (columns: samples in the

same order as dnam, rows: genes represented by ensembl IDs (e.g. ENSG00000239415))

tf.activity.es A matrix with normalized enrichment scores for each TF across all samples to be used in linear models instead of TF gene expression. See <a href="mailto:get\_tf\_ES">get\_tf\_ES</a>.

dnam.group.percent.threshold

DNA methylation threshold percentage to define samples in the low methylated group and high methylated group. For example, setting the threshold to 0.3 (30%) will assign samples with the lowest 30% methylation in the low group and the highest 30% methylation in the high group. Default is 0.25 (25%), accepted threshold range (0.0,0.5].

perform.correlation.analaysis

Perform correlation analysis?

remove.nonsig.correlated.dnam.target.gene

If spearman correlation of target expression and DNAm for all samples is not significant (pvalue > 0.05), triplet will be removed If wilcoxon test of target expression Q1 and Q4 is not significant (pvalue > 0.05), triplet will be removed.

remove.nonsig.correlated.dnam.target.gene.threshold.pvalue

Cut-off for remove.nonsig.correlated.dnam.target.gene in the spearman test remove.nonsig.correlated.dnam.target.gene.threshold.estimate

Cut-off for remove.nonsig.correlated.dnam.target.gene in the spearman test remove.sig.correlated.tf.exp.dnam

If wilcoxon test of TF expression Q1 and Q4 is significant (pvalue < 0.05), triplet will be removed.

filter.triplet.by.sig.term

Filter significant triplets ? Select triplets if any term is significant 1) interaction (TF x DNAm) p-value < 0.05 or 2) DNAm p-value < 0.05 or 3) TF p-value < 0.05 in binary model

filter.triplet.by.sig.term.using.fdr

Uses FRD instead of p-value when using filter.triplet.by.sig.term.

filter.triplet.by.sig.term.pvalue.threshold

P-values/FDR Threshold to filter significant triplets.

multiple.correction.by.stage.wise.analysis

A boolean indicating if stagewise analysis should be performed to correct for multiple comparisons. If set to FALSE then FDR analysis is performed.

tf.dnam.classifier.pval.threshold

P-value threshold to consider a linear model significant of not. Default 0.001. This will be used to classify the TF role and DNAm effect.

verbose A logical argument indicating if messages output should be provided.

cores Number of CPU cores to be used. Default 1.

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#### **Details**

This function fits the linear model

 $log2(RNA target) \sim log2(TF) + DNAm + log2(TF) * DNAm$ 

to triplet data as follow:

Model by considering DNAm as a binary variable - we defined a binary group for DNA methylation values (high = 1, low = 0). That is, samples with the highest DNAm levels (top 25 percent) has high = 1, samples with lowest DNAm levels (bottom 25 percent) has high = 0. Note that in this implementation, only samples with DNAm values in the first and last quartiles are considered.

In these models, the term log2(TF) evaluates direct effect of TF on target gene expression, DNAm evaluates direct effect of DNAm on target gene expression, and log2(TF)\*DNAm evaluates synergistic effect of DNAm and TF, that is, if TF regulatory activity is modified by DNAm.

There are two implementations of these models, depending on whether there are an excessive amount (i.e. more than 25 percent) of samples with zero counts in RNAseq data:

- When percent of zeros in RNAseq data is less than 25 percent, robust linear models are implemented using rlm function from MASS package. This gives outlier gene expression values reduced weight. We used "psi.bisqure" option in function rlm (bisquare weighting, https://stats.idre.ucla.edu/r/dae/robust-regression/).
- When percent of zeros in RNAseq data is more than 25 percent, zero inflated negative binomial models are implemented using zeroinfl function from pscl package. This assumes there are two processes that generated zeros (1) one where the counts are always zero (2) another where the count follows a negative binomial distribution.

To account for confounding effects from covariate variables, first use the get\_residuals function to obtain RNA or DNAm residual values which have covariate effects removed, then fit interaction model. Note that no log2 transformation is needed when interaction\_model is applied to residuals data.

Note that only triplets with TF expression not significantly different in high vs. low methylation groups will be evaluated (Wilcoxon test, p > 0.05).

# Value

A dataframe with Region, TF, target, TF\_symbo, target\_symbol, estimates and P-values, after fitting robust linear models or zero-inflated negative binomial models (see Details above).

Model considering DNAm values as a binary variable generates quant\_pval\_metGrp, quant\_pval\_rna.tf, quant\_estimates\_metGrp, quant\_estimates\_rna.tf, quant\_estimates\_metGrp.rna.tf, quant\_es

Model.interaction indicates which model (robust linear model or zero inflated model) was used to fit Model 1, and Model.quantile indicates which model(robust linear model or zero inflated model) was used to fit Model 2.

plot\_interaction\_model

Plot interaction model results

#### **Description**

Create several plots to show interaction data TF expression with target gene interaction using a linear model

$$log2(RNAtarget) = log2(TF) + DNAm + log2(TF) * DNAm$$

To consider covariates, RNA can also be the residuals.

```
log2(RNAtargetresiduals) = log2(TFresidual) + DNAm + log2(TFresidual) * DNAm
```

#### Usage

```
plot_interaction_model(
    triplet.results,
    dnam,
    exp,
    metadata,
    tf.activity.es = NULL,
    tf.dnam.classifier.pval.thld = 0.001,
    dnam.group.threshold = 0.25,
    label.dnam = "beta-value",
    label.exp = "expression",
    genome = "hg38",
    add.tf.vs.exp.scatter.plot = FALSE
)
```

#### **Arguments**

triplet.results

Output from function interaction\_model with Region ID, TF (column name: TF), and target gene (column name: target), p-values and estimates of interaction

dnam DNA methylation matrix or SummarizedExperiment object (columns: samples

same order as met, rows: regions/probes)

exp gene expression matrix or a SummarizedExperiment object (columns: samples

same order as met, rows: genes)

metadata A data frame with samples as rownames and one columns that will be used to

color the samples

tf.activity.es A matrix with normalized enrichment scores for each TF across all samples to be used in linear models instead of TF gene expression.

tf.dnam.classifier.pval.thld

P-value threshold to consider a linear model significant of not. Default 0.001. This will be used to classify the TF role and DNAm effect.

 ${\tt dnam.group.threshold}$ 

DNA methylation threshold percentage to define samples in the low methylated group and high methylated group. For example, setting the threshold to 0.3 (30%) will assign samples with the lowest 30% methylation in the low group and the highest 30% methylation in the high group. Default is 0.25 (25%), accepted threshold range (0.0,0.5].

label.dnam Used for label text. Option "beta-value" and "residuals" label.exp Used for label text. Option "expression" and "residuals"

```
\begin{tabular}{ll} {\tt genome} & {\tt Genome} & {\tt Genome} & {\tt ference} & {\tt to} & {\tt be} & {\tt added} & {\tt to} & {\tt the} & {\tt plot} & {\tt as} & {\tt text} \\ {\tt add.tf.vs.exp.scatter.plot} & & & & & & & & & \\ \end{tabular}
```

Add another row to the figure if the target gene expression vs TF expression stratified by DNA methylation groups (DNAmLow - low quartile, DNAmHigh - high quartile)

#### Value

A ggplot object, includes a table with results from fitting interaction model, and the the following scatter plots: 1) TF vs DNAm, 2) Target vs DNAm, 3) Target vs TF, 4) Target vs TF for samples in Q1 and Q4 for DNA methylation, 5) Target vs DNAm for samples in Q1 and Q4 for the TF

```
library(dplyr)
dnam <- runif(20, min = 0, max = 1) \%
  matrix(ncol = 1) %>% t
rownames(dnam) <- c("chr3:203727581-203728580")
colnames(dnam) <- paste0("Samples",1:20)</pre>
exp.target <- runif(20,min = 0,max = 10) %>%
 matrix(ncol = 1) %>% t
rownames(exp.target) <- c("ENSG00000252982")</pre>
colnames(exp.target) <- paste0("Samples",1:20)</pre>
exp.tf <- runif(20,min = 0,max = 10) %>%
  matrix(ncol = 1) %>% t
rownames(exp.tf) <- c("ENSG00000083937")
colnames(exp.tf) <- paste0("Samples",1:20)</pre>
exp <- rbind(exp.tf, exp.target)</pre>
triplet <- data.frame(</pre>
   "regionID" = c("chr3:203727581-203728580"),
   "target" = "ENSG00000252982",
   "TF" = "ENSG00000083937"
)
results <- interaction_model(</pre>
   triplet = triplet,
   dnam = dnam,
   exp = exp,
    dnam.group.threshold = 0.25,
   stage.wise.analysis = FALSE,
   sig.threshold = 1,
   filter.correlated.tf.exp.dnam = FALSE,
   filter.correlated.target.exp.dnam = FALSE,
   filter.triplet.by.sig.term = FALSE
plots <- plot_interaction_model(</pre>
    triplet.results = results,
    dnam = dnam,
    exp = exp
)
```

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```
plot_stratified_model Plot stratified model results
```

## **Description**

Create several plots to show interaction data TF expression with target gene interaction using a linear model

```
log2(RNAtarget) log2(TF)
```

to samples with highest DNAm values (top 25 percent) and lowest DNAm values (bottom 25 percent), separately.

# Usage

```
plot_stratified_model(
    triplet.results,
    dnam,
    exp,
    metadata,
    label.dnam = "beta-value",
    label.exp = "expression",
    tf.activity.es = NULL,
    dnam.group.threshold = 0.25
)
```

# **Arguments**

triplet.results

Output from function stratified\_model with Region ID, TF (column name: TF), and target gene (column name: target), p-values and estimates of interaction

dnam DNA methylation matrix or SummarizedExperiment object (columns: samples

same order as met, rows: regions/probes)

exp A gene expression matrix or SummarizedExperiment object (columns: samples

same order as met, rows: genes)

metadata A data frame with samples as row names and one columns that will be used to

color the samples

label.dnam Used for label text. Option "beta-value" and "residuals"

label.exp Used for label text. Option "expression" and "residuals"

tf.activity.es A matrix with normalized enrichment scores for each TF across all samples to

be used in linear models instead of TF gene expression.

dnam.group.threshold

DNA methylation threshold percentage to define samples in the low methylated group and high methylated group. For example, setting the threshold to 0.3 (30%) will assign samples with the lowest 30% methylation in the low group and the highest 30% methylation in the high group. Default is 0.25 (25%), accepted threshold range (0.0,0.5].

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#### Value

A ggplot object, includes a table with results from fitting stratified model, and the following scatter plots: 1) TF vs DNAm, 2) Target vs DNAm, 3) Target vs TF, 4) Target vs TF for samples in Q1 and Q4 for DNA methylation, 5) Target vs DNAm for samples in Q1 and Q4 for the TF

readRemap2022

Access REMAP2022 non-redundant peaks

# **Description**

Access REMAP2022 non-redundant peaks

# Usage

```
readRemap2022(cell_line)
```

# **Arguments**

cell\_line

filter peaks using cell line description field

stratified\_model

Fits linear models to triplet data (Target, TF, DNAm) for samples with high DNAm or low DNAm separately, and annotates TF (activator/repressor) and DNam effect over TF activity (attenuate, enhance).

# **Description**

Should be used after fitting interaction\_model, and only for triplet data with significant TF\*DNAm interaction. This analysis examines in more details on how TF activities differ in samples with high DNAm or low DNAm values.

# Usage

```
stratified_model(
   triplet,
   dnam,
   exp,
   cores = 1,
   tf.activity.es = NULL,
   tf.dnam.classifier.pval.thld = 0.001,
   dnam.group.threshold = 0.25
)
```

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## **Arguments**

triplet Data frame with columns for DNA methylation region (regionID), TF (TF), and

target gene (target)

dnam DNA methylation matrix or SummarizedExperiment (columns: samples in the

same order as exp matrix, rows: regions/probes)

exp A matrix or SummarizedExperiment (columns: samples in the same order as

dnam matrix, rows: genes represented by ensembl IDs (e.g. ENSG00000239415))

cores Number of CPU cores to be used. Default 1.

tf.activity.es A matrix with normalized enrichment scores for each TF across all samples to

be used in linear models instead of TF gene expression.

tf.dnam.classifier.pval.thld

P-value threshold to consider a linear model significant of not. Default 0.001.

This will be used to classify the TF role and DNAm effect.

dnam.group.threshold

DNA methylation threshold percentage to define samples in the low methylated group and high methylated group. For example, setting the threshold to 0.3 (30%) will assign samples with the lowest 30% methylation in the low group and the highest 30% methylation in the high group. Default is 0.25 (25%),

accepted threshold range (0.0,0.5].

#### **Details**

This function fits linear model log2(RNA target) = log2(TF)

to samples with highest DNAm values (top 25 percent) or lowest DNAm values (bottom 25 percent), separately.

There are two implementations of these models, depending on whether there are an excessive amount (i.e. more than 25 percent) of samples with zero counts in RNAseq data:

- When percent of zeros in RNAseq data is less than 25 percent, robust linear models are implemented using rlm function from MASS package. This gives outlier gene expression values reduced weight. We used "psi.bisqure" option in function rlm (bisquare weighting, https://stats.idre.ucla.edu/r/dae/robust-regression/).
- When percent of zeros in RNAseq data is more than 25 percent, zero inflated negative binomial models are implemented using zeroinfl function from pscl package. This assumes there are two processes that generated zeros (1) one where the counts are always zero (2) another where the count follows a negative binomial distribution.

To account for confounding effects from covariate variables, first use the get\_residuals function to obtain RNA residual values which have covariate effects removed, then fit interaction model. Note that no log2 transformation is needed when interaction\_model is applied to residuals data.

This function also provides annotations for TFs. A TF is annotated as activator if increasing amount of TF (higher TF gene expression) corresponds to increased target gene expression. A TF is annotated as repressor if increasing amount of TF (higher TF gene expression) corresponds to decrease in target gene expression. A TF is annotated as dual if in the Q1 methylation group increasing amount of TF (higher TF gene expression) corresponds to increase in target gene expression, while in Q4 methylation group increasing amount of TF (higher TF gene expression) corresponds to decrease in target gene expression (or the same but changing Q1 and Q4 in the previous sentence).

In addition, a region/CpG is annotated as enhancing if more TF regulation on gene transcription is observed in samples with high DNAm. That is, DNA methylation enhances TF regulation on

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target gene expression. On the other hand, a region/CpG is annotated as attenuating if more TF regulation on gene transcription is observed in samples with low DNAm. That is, DNA methylation reduces TF regulation on target gene expression.

#### Value

A data frame with Region, TF, target, TF\_symbol target\_symbol, results for fitting linear models to samples with low methylation (DNAmlow\_pval\_rna.tf, DNAmlow\_estimate\_rna.tf), or samples with high methylation (DNAmhigh\_pval\_rna.tf, DNAmhigh\_pval\_rna.tf.1), annotations for TF (class.TF) and (class.TF.DNAm).

```
library(dplyr)
dnam <- runif (20,min = 0,max = 1) %>%
  matrix(ncol = 1) %>% t
rownames(dnam) <- c("chr3:203727581-203728580")
colnames(dnam) <- paste0("Samples",1:20)</pre>
exp.target <- runif (20,min = 0,max = 10) %>%
 matrix(ncol = 1) %>% t
\verb|rownames(exp.target)| <- c("ENSG00000232886")|
colnames(exp.target) <- paste0("Samples",1:20)</pre>
exp.tf <- runif (20,min = 0,max = 10) %>%
  matrix(ncol = 1) %>% t
rownames(exp.tf) <- c("ENSG00000232888")</pre>
colnames(exp.tf) <- paste0("Samples",1:20)</pre>
exp <- rbind(exp.tf, exp.target)</pre>
triplet <- data.frame(</pre>
   "regionID" = c("chr3:203727581-203728580"),
   "target" = "ENSG00000232886",
   "TF" = "ENSG00000232888"
)
results <- stratified_model(</pre>
  triplet = triplet,
  dnam = dnam,
  exp = exp
)
```

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