Package ‘epiregulon’

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Title   Gene regulatory network inference from single cell epigenomic data

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Description   Gene regulatory networks model the underlying gene regulation hierarchies that drive gene expression and observed phenotypes. Epiregulon infers TF activity in single cells by constructing a gene regulatory network (regulons). This is achieved through integration of scATAC-seq and scRNA-seq data and incorporation of public bulk TF ChIP-seq data. Links between regulatory elements and their target genes are established by computing correlations between chromatin accessibility and gene expressions.

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VignetteBuilder knitr

URL https://github.com/xiaosaiyao/epiregulon/

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epiregulon-package

epiregulon: Gene regulatory network inference from single cell epigenomic data

Description

Gene regulatory networks model the underlying gene regulation hierarchies that drive gene expression and observed phenotypes. Epiregulon infers TF activity in single cells by constructing a gene regulatory network (regulons). This is achieved through integration of scATAC-seq and scRNA-seq data and incorporation of public bulk TF ChIP-seq data. Links between regulatory elements and their target genes are established by computing correlations between chromatin accessibility and gene expressions.
Author(s)

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• Shang-Yang Chen <sychen9584@gmail.com>

See Also

Useful links:

• https://github.com/xiaosaiyao/epiregulon/
• Report bugs at https://github.com/xiaosaiyao/epiregulon/issues

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addLogFC

Add log fold changes of gene expression to regulons

Description

Add log fold changes of gene expression to regulons

Usage

addLogFC(
  expMatrix,
  clusters,
  regulon,
  pval.type = c("any", "some", "all"),
  sig_type = c("FDR", "p.value"),
  logFC_condition = NULL,
  logFC_ref = NULL,
  ...
)

Arguments

expMatrix A SingleCellExperiment object or matrix containing gene expression with genes in the rows and cells in the columns
clusters A character or integer vector of cluster or group labels for single cells
regulon A dataframe informing the gene regulatory relationship with the tf column representing transcription factors, idxATAC corresponding to the index in the peak-Matrix and target column representing target genes
pval.type String specifying how p-values are to be combined across pairwise comparisons for a given group/cluster.
addMotifScore

**Description**

Add Motif Scores

<table>
<thead>
<tr>
<th>sig_type</th>
<th>String specifying whether to use &quot;FDR&quot; or &quot;p.value&quot; for sig_cutoff</th>
</tr>
</thead>
<tbody>
<tr>
<td>logFC_condition</td>
<td>A scalar or vector of string indicating the sample names to be compared against</td>
</tr>
<tr>
<td>logFC_ref</td>
<td>A scalar indicating the reference sample used to compute logFC. Default value is rest which is an average of all pairwise comparisons. Users can also specify a reference sample, for example, DMSO.</td>
</tr>
</tbody>
</table>

**Value**

A DataFrame of regulons with additional columns of logFC and significance

**Author(s)**

Xiaosai Yao

**Examples**

```r
# create a mock singleCellExperiment object for gene expression matrix
set.seed(1000)
gene_sce <- scuttle::mockSCE()
gene_sce <- scuttle::logNormCounts(gene_sce)
rownames(gene_sce) <- paste0('Gene_',1:2000)

# create a mock regulon
regulon <- data.frame(tf = c(rep('Gene_1',10), rep('Gene_2',10)),
                      idxATAC = sample(1:100, 20),
target = c(paste0('Gene_', sample(3:2000,10)),
                      paste0('Gene_',sample(3:2000,10))))

# filter regulon
pruned.regulon <- addLogFC(expMatrix = gene_sce, clusters = gene_sce$Treatment,
                           regulon = regulon,
                           sig_type = "p.value")
```
**Usage**

```r
addMotifScore(
  regulon,
  field_name = "motif",
  peaks = NULL,
  pwms = NULL,
  species = c("human", "mouse"),
  genome = c("hg38", "hg19", "mm10"),
  ...
)
```

**Arguments**

- **regulon**
  A DataFrame consisting of tf (regulator) and target in the column names.

- **field_name**
  Character string indicating the column name of the regulon to add the motif information to.

- **peaks**
  A GRanges object indicating the peaks to perform motif annotation on if ArchR project is not provided. The peak indices should match the re column in the regulon.

- **pwms**
  A PWMatrixList for annotation of motifs using 'motifmatchr::matchMotifs'

- **species**
  Character string indicating species. Currently supported species is human or mouse.

- **genome**
  Character string indicating the genomic build.

- **...**
  Additional arguments to pass into motifmatchr::matchMotifs

**Value**

A DataFrame with motif matches added with 1s indicating the presence of motifs and 0s indicating the absence of motifs.

**Examples**

```r
regulon <- S4Vectors::DataFrame(tf = c('AR', 'AR', 'AR', 'ESR1', 'ESR1', 'NKX2-1'),
idxATAC = 1:6)
peaks <- GenomicRanges::GRanges(seqnames = c('chr12', 'chr19', 'chr19', 'chr11', 'chr6', 'chr1'),
ranges = IRanges::IRanges(start = c(124914563, 50850845, 50850844, 101034172, 151616327, 1000),
end = c(124914662, 50850929, 50850929, 101034277, 151616394, 2000)))
regulon <- addMotifScore(regulon, peaks=peaks)
```

---

### AddTFMotifInfo

**Add TF binding motif occupancy information to the peak2gene object**

**Description**

Add TF binding motif occupancy information to the peak2gene object.
addTFMotifInfo

Usage

addTFMotifInfo(p2g, grl, peakMatrix = NULL)

Arguments

p2g  A Peak2Gene dataframe created by ArchR or getP2Glinks() function
grl  GRangeList object containing reference TF binding information. We recommend retrieving grl from getTFMotifInfo which contains TF occupancy data derived from public and ENCODE ChIP-seq peaks. Alternatively, if the users would like to provide a GRangeList of motif annotations. This can be derived using motifmatchr::matchMotifs. See details
peakMatrix A matrix of scATAC-seq peak regions with peak ids as rows

Details

This function annotates each regulatory element with possible transcription factors. We can either provide a GRangeList of known ChIP-seq binding sites (TF occupancy) or positions of TF motifs (TF motifs). While public ChIP-seq data may not fully align with the ground truth TF occupancy in users' data (due to technical challenges of ChIP-seq or cell type nature of TF occupancy), it does offer a few important advantages over TF motif information:

1. TF occupancy allows co-activators to be included. Co-activators are chromatin modifiers that do not directly bind to DNA but nonetheless play an important role in gene regulation
2. TF occupancy can distinguish between members of the same class that may share similar motifs but that may have drastically different binding sites

If multiple ChIP-seq are available for the same TF, we merge the ChIP-seq data to represent an universal set of possible binding sites. The predicted TF occupancy is further refined by pruneRegulon.

If the users prefer to use TF motifs instead of TF occupancy, the users can create a GRangeList of motif annotation using motifmatchr::matchMotifs. Here, we demonstrate how to annotate peaks with cisbp motif database

```r
library(motifmatchr)
library(chromVARmotifs)
data("human_pwms_v1")
peaks <- GRanges(seqnames = c("chr1", "chr2", "chr2"),
                 ranges = IRanges(start = c(76585873, 42772928, 100183786),
                                 width = 500))
eh <- AnnotationHub::query(ExperimentHub::ExperimentHub(),
                           pattern = c("scMultiome", "TF motifs", "human"))
pwms <- readRDS(eh[[eh$ah_id]]))
grl <- matchMotifs(pwms, peaks, genome = "hg38", out = "positions")
retain only TF symbols. TF symbols need to be consistent with gene names in regulon
names(grl) <- sapply(strsplit(names(grl), "_"), "[",3)
```

Value

A data frame containing overlapping ids of scATAC-seq peak regions and reference TF binding regions
addWeights

Calculate weights for the regulons by computing co-association between TF and target gene expression

Description

Calculate weights for the regulons by computing co-association between TF and target gene expression

Usage

addWeights(
  regulon,
  expMatrix = NULL,
  peakMatrix = NULL,
  exp_assay = "logcounts",
  peak_assay = "PeakMatrix",
  method = c("wilcoxon", "corr", "MI"),
)
clusters = NULL,
exp_cutoff = 1,
peak_cutoff = 0,
block_factor = NULL,
aggregation_function = mean,
min_targets = 10,
tf_re.merge = FALSE,
aggregateCells = FALSE,
useDim = "IterativeLSI_ATAC",
cellNum = 10,
BPPARAM = BiocParallel::SerialParam(progressbar = TRUE)
)

Arguments

regulon A DataFrame object consisting of tf (regulator) and target in the column names.
expMatrix A SingleCellExperiment object containing gene expression information
peakMatrix A SingleCellExperiment object or matrix containing peak accessibility with
peaks in the rows and cells in the columns
exp_assay String specifying the name of the assay to be retrieved from the SingleCellEx-
periment object
peak_assay String indicating the name of the assay in peakMatrix for chromatin accessibility
method String specifying the method of weights calculation. Four options are available: corr, MI, lmfit, wilcoxon and logFC.
clusters A vector corresponding to the cluster labels of the cells
exp_cutoff A scalar indicating the minimum gene expression for transcription factor above
which cell is considered as having expressed transcription factor.
peak_cutoff A scalar indicating the minimum peak accessibility above which peak is consid-
ered open.
block_factor String specifying the field in the colData of the SingleCellExperiment object to
be used as blocking factor (such as batch)
aggregation_function Function being used for summarizing weights from the transcription factor-
target gene pair with many regulatory elements.
min_targets Integer specifying the minimum number of targets for each tf in the regulon with
10 targets as the default
tf_re.merge A logical to indicate whether to consider both TF expression and chromatin
accessibility. See details.
aggregateCells A logical to indicate whether to aggregate cells into groups determined by cell-
Num. This option can be used to overcome data sparsity when using wilcoxon.
useDim String indicating the name of the dimensionality reduction matrix in expMatrix
used for cell aggregation
cellNum An integer specifying the number of cells per cluster for cell aggregation. De-
fault is 10.
**addWeights**

A BiocParallelParam object specifying whether summation should be parallelized. Use BiocParallel::SerialParam() for serial evaluation and use BiocParallel::MulticoreParam() for parallel evaluation.

**Details**

This function estimates the regulatory potential of transcription factor on its target genes, or in other words, the magnitude of gene expression changes induced by transcription factor activity, using one of the four methods:

- **corr** - correlation between TF and target gene expression
- **MI** - mutual information between the TF and target gene expression
- **wilcoxon** - effect size of the Wilcoxon test between target gene expression in cells jointly expressing all 3 elements vs cells that do not

Two measures (corr and wilcoxon) give both the magnitude and directionality of changes whereas MI always outputs positive weights. The correlation and mutual information statistics are computed on the pseudobulked gene expression or accessibility matrices, whereas the Wilcoxon method groups cells based on the joint expression of TF, RE and TG in each single cell.

When using the corr method, the default practice is to compute weights by correlating the pseudobulk target gene expression vs the pseudobulk TF gene expression. However, often times, an inhibitor of TF does not alter the gene expression of the TF. In rare cases, cells may even compensate by increasing the expression of the TF. In this case, the activity of the TF, if computed by TF-TG correlation, may show a spurious increase in its activity. As an alternative to gene expression, we may correlate the product of TF and RE against TG. When tf_re.merge is TRUE, we take the product of the gene expression and chromatin accessibility.

**Value**

A DataFrame with columns of corr and/or MI added to the regulon. TFs not found in the expression matrix and regulons not meeting the minimal number of targets were filtered out.

**Author(s)**

Xiaosai Yao, Shang-yang Chen, Tomasz Wlodarczyk

**Examples**

```r
dataMatrix <- scuttle::mockSCE() dataMatrix <- scuttle::logNormCounts(dataMatrix) dataMatrix$cluster <- sample(LETTERS[1:5], ncol(dataMatrix), replace=TRUE)

# create a mock singleCellExperiment object for peak matrix
peakMatrix <- scuttle::mockSCE() rownames(peakMatrix) <- 1:2000

# create a mock regulon
regulon <- S4Vectors::DataFrame(tf=c(rep('Gene_0001',5), rep('Gene_0002',10)),
       idxATAC=1:15,
```
aggregateAcrossCells

Aggregate expression across cells

Description
Aggregate expression values across cells based on one or more grouping factors. This is primarily used to create pseudo-bulk profiles for each cluster/sample combination.

Usage
aggregateAcrossCells(x, factors, num.threads = 1)

Arguments
- **x**: Any matrix-like object. Expression values are typically expected to be counts.
- **factors**: A list or data frame containing one or more grouping factors. Each entry should be a factor of the same length as the number of cells in x.
- **num.threads**: Integer specifying the number of threads to be used for aggregation.

Value
A list containing:
- **sums**: a numeric matrix where each row corresponds to a gene and each column corresponds to a unique combination of grouping levels. Each entry contains the summed expression across all cells with that combination.
- **detected**: an integer matrix where each row corresponds to a gene and each column corresponds to a unique combination of grouping levels. Each entry contains the number of cells with detected expression in that combination.
- **combinations**: a data frame describing the levels for each unique combination. Rows of this data frame correspond to columns of sums and detected, while columns correspond to the factors in factors.
- **counts**: the number of cells associated with each combination. Each entry corresponds to a row of combinations.
• index, an integer vector of length equal to the number of cells in \( x \). This specifies the combination in combinations to which each cell was assigned.

**Author(s)**

Aaron Lun

**Examples**

```r
# Mocking a matrix:
library(Matrix)
y <- round(abs(rsparsematrix(1000, 100, 0.1) * 100))

# Simple aggregation:
clusters <- sample(LETTERS, 100, replace=TRUE)
agg <- aggregateAcrossCells(y, list(cluster=clusters))
str(agg)

# Multi-factor aggregation
samples <- sample(1:5, 100, replace=TRUE)
agg2 <- aggregateAcrossCells(y, list(cluster=clusters, sample=samples))
str(agg2)
```

---

**aggregateAcrossCellsFast**

*Aggregate cells in SingleCellExperiment*

**Description**

Aggregate expression values across cells in SingleCellExperiment based on a grouping factor. This is primarily used to create pseudo-bulk profiles for each cluster/sample combination. It is wrapped around aggregateAcrossCells, which relies on the C++ code.

**Usage**

```r
aggregateAcrossCellsFast(
    sce,
    clusters,
    assay.name = "counts",
    fun_name = c("mean", "sum"),
    num.threads = 1,
    aggregateColData = TRUE
)
```
Arguments

sce     A SingleCellExperiment object
clusters A vector used as a grouping variable. The length should be equal to the number of cells.
assay.name A character indicating the name of the assay containing the values to be aggregated.
fun_name A character indicating the function used to aggregate data. The selection is restricted to "mean" or "sum".
num.threads Integer specifying the number of threads to be used for aggregation.
aggregateColData A logical specifying if the columns in the colData should be included in the output object. Only those columns are selected which can be decomposed by grouping variable into the vectors whose all elements are the same.

Value

A SingleCellExperiment object containing aggregated cells.

Examples

# create a mock singleCellExperiment object for gene expression matrix
set.seed(1000)
example_sce <- scuttle::mockSCE()
ids <- sample(LETTERS[1:5], ncol(example_sce), replace=TRUE)
out <- aggregateAcrossCellsFast(example_sce, ids)

calculateActivity

Calculate the per cell activity of master regulators based on a regulon

Description

Calculate the per cell activity of master regulators based on a regulon

Usage

calculateActivity(
    expMatrix = NULL,
    exp_assay = "logcounts",
    regulon = NULL,
    normalize = FALSE,
    mode = "weight",
    method = c("weightedmean", "aucell"),
    genesets = NULL,
    clusters = NULL,
    FUN = c("mean", "sum"),
    ncore = 1,
    BPPARAM = BiocParallel::SerialParam()
  )
**Arguments**

- `expMatrix`: A `SingleCellExperiment` object containing gene expression information with rows representing genes and columns represent cells. Row names (either gene symbols or geneID) must be consistent with the naming convention in the regulon.
- `exp_assay`: String specifying the name of the assay to be retrieved from the `SingleCellExperiment` object. Set to 'logcounts' as the default.
- `regulon`: A DataFrame object consisting of tf (regulator) and target in the column names, with additional columns indicating degree of association between tf and target such as 'mor' or 'corr' obtained from `addWeights`.
- `normalize`: Logical indicating whether row means should be subtracted from expression matrix. Default is FALSE.
- `mode`: String indicating the name of column to be used as the weights.
- `method`: String indicating the method for calculating activity. Available methods are `weightedMean` or `aucell`.
- `genesets`: A list of genesets. Each list element can be a dataframe with the first column indicating the genes and second column indicating the weights. Alternatively, each list element is a character vector corresponding to the genes in the geneset. A feature set collection in the form of `CompressedSplitDataFrameList` that contains genes in the first column and weights in the second column. See details.
- `clusters`: A vector indicating cluster assignment.
- `FUN`: Function to aggregate the weights.
- `ncore`: Integer specifying the number of cores to be used in `AUCell`.
- `BPPARAM`: A `BiocParallelParam` object specifying whether summation should be parallelized. Use `BiocParallel::SerialParam()` for serial evaluation and use `BiocParallel::MulticoreParam()` for parallel evaluation.

**Details**

This function calculates activity score from a regulon that is a DataFrame consisting of a tf column, a target column and a weight column. Alternatively, instead of a regulon, this function also accepts weighted signature sets where each gene set or signature is a data frame or unweighted signature sets where each gene set is a character vector. The user has the option of computing signature score by weighted mean of target gene expression or the relative ranking of the target genes computed by `AUCell`.

**Value**

A matrix of inferred transcription factor (row) activities in single cells (columns).

**Author(s)**

Xiaosai Yao, Shang-yang Chen
Examples

# create a mock singleCellExperiment object for gene expression matrix
set.seed(1000)
gene_sce <- scuttle::mockSCE()
gene_sce <- scuttle::logNormCounts(gene_sce)
rownames(gene_sce) <- paste0('Gene_', 1:2000)

# create a mock singleCellExperiment object for peak matrix
peak_gr <- GRanges(seqnames = 'chr1',
                  ranges = IRanges(start = seq(from = 1, to = 10000, by = 100), width = 100))
peak_counts <- matrix(sample(x = 0:4, size = ncol(gene_sce)*length(peak_gr), replace = TRUE),
                       nrow = length(peak_gr), ncol=ncol(gene_sce))
peak_sce <- SingleCellExperiment(list(counts = peak_counts), colData = colData(gene_sce))
rownames(peak_sce) <- paste0('Peak_', 1:100)

# create a mock regulon
regulon <- data.frame(tf = c(rep('Gene_1',10), rep('Gene_2',10)),
                      idxATAC = sample(1:100, 20),
                      target = c(paste0('Gene_', sample(3:2000,10)),
                                     paste0('Gene_',sample(3:2000,10))))

# # prune regulon
pruned.regulon <- pruneRegulon(expMatrix = gene_sce,
                               exp_assay = 'logcounts',
                               peakMatrix = peak_sce,
                               peak_assay = 'counts',
                               regulon = regulon,
                               clusters = gene_sce$Treatment,
                               regulon_cutoff = 0.5,
                               p_adj = TRUE)

regulon.w <- addWeights(regulon = regulon,
                         expMatrix = gene_sce,
                         clusters = gene_sce$Treatment,
                         exp_assay = 'logcounts',
                         min_targets = 5,
                         method = 'corr')

# calculate activity
activity <- calculateActivity(expMatrix = gene_sce,
                               regulon = regulon.w,
                               exp_assay = 'logcounts')

# calculate cluster-specific activity if cluster-specific weights are supplied
regulon.w$weight <- matrix(rnorm(ncol(gene_sce)*2, 0, 1), nrow(regulon.w),2)
colnames(regulon.w$weight) <- c('treat1','treat2')
activity.cluster <- calculateActivity(gene_sce,
                                      regulon = regulon.w, clusters = gene_sce$Treatment,
                                      exp_assay = 'logcounts', FUN = 'mean')

# compute signature scores from weighted genesets
weighted_genesets <- list(set1 = data.frame(genes = c('Gene_1', 'Gene_2', 'Gene_3'), weights = c(1,2,3)), set2 = data.frame(genes = c('Gene_4', 'Gene_5', 'Gene_6'), weights = c(4,5,6)))

activity <- calculateActivity(gene_sce, genesets = weighted_genesets)

# compute signature scores from unweighted genesets
unweighted_genesets <- list(set1 = c('Gene_1', 'Gene_2', 'Gene_3'), set2 = c('Gene_4', 'Gene_5', 'Gene_6'))
activity <- calculateActivity(gene_sce, genesets = unweighted_genesets)

calculateP2G

Establish peak to gene links based on correlations between ATAC-seq peaks and RNA-seq genes

description
Establish peak to gene links based on correlations between ATAC-seq peaks and RNA-seq genes

Usage

calculateP2G(
  peakMatrix = NULL,
  expMatrix = NULL,
  reducedDim = NULL,
  useDim = "IterativeLSI",
  maxDist = 250000,
  cor_cutoff = 0.5,
  cellNum = 100,
  exp_assay = "logcounts",
  peak_assay = "counts",
  gene_symbol = "name",
  clusters = NULL,
  cor_method = c("pearson", "kendall","spearman"),
  assignment_method = c("correlation", "nearest"),
  frac_RNA = 0,
  frac_ATAC = 0,
  BPPARAM = BiocParallel::SerialParam()
)

Arguments

peakMatrix A SingleCellExperiment object containing counts of chromatin accessibility at each peak region or genomic bin from scATAC-seq. rowRanges should contain genomic positions of the peaks in the form of GRanges.

exMatrix A SingleCellExperiment object containing gene expression counts from scRNA-seq. rowRanges should contain genomic positions of the genes in the form of GRanges. rowData should contain a column of gene symbols with column name matching the gene_symbol argument.
calculateP2G

reducedDim A matrix of dimension reduced values
useDim String specifying the name of the reduced dimension matrix supplied by the user
maxDist An integer to specify the base pair extension from transcription start start for overlap with peak regions
cor_cutoff A numeric scalar to specify the correlation cutoff between ATAC-seq peaks and RNA-seq genes to assign peak to gene links. Default correlation cutoff is 0.5.
cellNum An integer to specify the number of cells to include in each K-means cluster
exp_assay String indicating the name of the assay in expMatrix for gene expression
peak_assay String indicating the name of the assay in peakMatrix for chromatin accessibility
gene_symbol String indicating the column name in the rowData of expMatrix that corresponds to gene symbol
clusters A vector corresponding to the cluster labels for calculation of correlations within each cluster. If left NULL, correlation is calculated across all clusters. See details for the use of clusters
cor_method String indicating which correlation coefficient is to be computed. One of 'pearson' (default), 'kendall', or 'spearman'.
assignment_method String indicating the method used to assign target genes to regulatory elements. 'Correlation' is based on correlation between ATAC and RNA above a correlation threshold set by cor_cutoff. 'Nearest' assigns the closest expressed gene to regulatory element meeting a correlation threshold set by cor_cutoff. Set cor_cutoff to 0 if wishing to assign the closest expressed gene without any correlation cutoff
frac_RNA An integer to indicate the fraction of cells expressing a gene. It is used to filter the gene expression matrix for expressed genes
frac_ATAC An integer to indication the fraction of cells showing chromatin accessibility. It is used to filter the peak Matrix for open regions
BPPARAM A BiocParallelParam object specifying whether summation should be parallelized. Use BiocParallel::SerialParam() for serial evaluation and use BiocParallel::MulticoreParam() for parallel evaluation

Details
Cluster information is sometimes helpful to avoid the Simpson's paradox in which baseline differences between cell lines or cell types can create artificial or even inverse correlations between peak accessibility and gene expression. If Cluster information is provided, correlation is performed within cell aggregates of each cluster.

Value
A DataFrame of Peak to Gene correlation

Author(s)
Xiaosai Yao, Shang-yang Chen
getRegulon

Combine the TF binding info and peak to gene correlations to generate regulons

Examples

```r
# create a mock singleCellExperiment object for gene expression matrix
set.seed(1000)
gene_sce <- scuttle::mockSCE()
gene_sce <- scuttle::logNormCounts(gene_sce)
gene_gr <- GRanges(seqnames = Rle(c('chr1', 'chr2', 'chr3', 'chr4'), nrow(gene_sce)/4),
                  ranges = IRanges(start = seq(from = 1, length.out=nrow(gene_sce), by = 1000),
                                 width = 100))
rownames(gene_sce) <- rownames(gene_sce)
gene_gr$name <- rownames(gene_sce)
rowRanges(gene_sce) <- gene_gr

# create a mock singleCellExperiment object for peak matrix
peak_gr <- GRanges(seqnames = 'chr1',
                   ranges = IRanges(start = seq(from = 1, to = 10000, by = 1000),
                                   width = 100))
peak_counts <- matrix(sample(x = 0:4, size = ncol(gene_sce)*length(peak_gr), replace = TRUE),
                       nrow = length(peak_gr), ncol=ncol(gene_sce))
peak_sce <- SingleCellExperiment(list(counts = peak_counts), colData = colData(gene_sce))
rowRanges(peak_sce) <- peak_gr
rownames(peak_sce) <- paste0('peak',1:10)

# create a mock reducedDim matrix
reducedDim_mat <- matrix(runif(ncol(gene_sce)*50, min = 0, max = 1), nrow = ncol(gene_sce), 50)
p2g <- calculateP2G(peakMatrix = peak_sce, expMatrix = gene_sce, reducedDim = reducedDim_mat,
                     cellNum = 20, clusters = gene_sce$Treatment)
```

Description

Combine the TF binding info and peak to gene correlations to generate regulons

Usage

```
getRegulon(p2g, overlap, aggregate = FALSE, FUN = "mean")
```

Arguments

- **p2g**: A Peak2Gene data frame created by ArchR or getP2Glinks() function
- **overlap**: A data frame storing overlaps between the regions of the peak matrix with the bulk TF ChIP-seq binding sites computed from addTFMotifInfo
- **aggregate**: logical to specify whether regulatory elements are aggregated across the same TF-target pairs
- **FUN**: function to aggregate TF-target sharing different regulatory elements
getTFMotifInfo

Retrieve TF binding sites or motif positions

**Value**

A DataFrame consisting of tf(regulator), target and a column indicating degree of association between TF and target such as 'mor' or 'corr'.

**Author(s)**

Xiaosai Yao, Shang-yang Chen

**Examples**

```r
set.seed(1)
# create a mock peak-to-gene matrix
p2g <- data.frame(idxATAC = c(rep(1,5), rep(2,5)), Chrom = 'chr1', idxRNA = 1:10, target = paste0('Gene_', 1:10), Correlation = runif(10, 0, 1))

# create a Granges list of TF binding sites
grl <- GRangesList('TF1' = GRanges(seqnames = 'chr1', ranges = IRanges(start = c(50,1050), width = 100)), 'TF2' = GRanges(seqnames = 'chr1', ranges = IRanges(start = c(1050), width = 100))
)

# Create a mock peak matrix
peak_gr <- GRanges(seqnames = 'chr1',
   ranges = IRanges(start = seq(from = 1, to = 10000, by = 1000), width = 100))
peak_counts <- matrix(sample(x = 0:4, size = 100*length(peak_gr), replace = TRUE), nrow = length(peak_gr),ncol = 100)
peak_sce <- SingleCellExperiment(list(counts = peak_counts))
rowRanges(peak_sce) <- peak_gr
rownames(peak_sce) <- paste0('peak', 1:10)

# create overlaps between p2g matrix, TF binding sites and peak matrix
overlap <- addTFMotifInfo(p2g, grl, peakMatrix = peak_sce)
utils::head(overlap)

# aggregate gene expression if the gene is bound by the same TF at regulatory elements
regulon <- getRegulon(p2g, overlap, aggregate = FALSE)
```

**Description**

Combined transcription factor ChIP-seq data from ChIP-Atlas and ENCODE or from CistromeDB and ENCODE.
getTFMotifInfo

Usage

getTFMotifInfo(
  genome = c("hg38", "hg19", "mm10"),
  source = c("atlas", "cistrome", "encode.sample", "atlas.sample", "atlas.tissue"),
  metadata = FALSE,
  mode = c("occupancy", "motif"),
  peaks = NULL
)

Arguments

genome character string specifying the genomic build
source character string specifying the ChIP-seq data source
metadata logical flag specifying whether to return data or metadata only
mode a string indicating whether to download a GRangelist of TF binding sites (‘occupancy’) or motif matches (‘motif’). TF binding information is retrieved from scMultiome::tfBinding. The motif information was obtained from chromVAR-motifs (human_pwms_v2 and mouse_pwms_v2, version 0.2 with filtering of cisBP motifs) and is also hosted on scMultiome.
peaks A GRanges object indicating the peaks to perform motif annotation on if ArchR project is not provided. The peak indices should match the re column in the regulon

Value

A list of TF binding sites as a GrangesList object.

References


ENCODE: https://www.encodeproject.org/


Examples

```r
# retrieve TF binding info
getTFMotifInfo('mm10', 'atlas')
```

# retrieve motif info
```r
peaks <- GRanges(seqnames = c('chr12','chr19','chr19','chr11','chr6'),
                 ranges = IRanges(start = c(124914563,50850845, 50850844, 101034172, 151616327),
                                end = c(124914662,50850929, 50850929, 101034277, 151616394)))
grl <- getTFMotifInfo(genome = 'hg38', mode = 'motif', peaks=peaks)
```

---

**pruneRegulon**

Prune regulons for true transcription factor - regulatory elements - target genes relationships

---

**Description**

Prune regulons for true transcription factor - regulatory elements - target genes relationships

**Usage**

```r
pruneRegulon(
    regulon, expMatrix = NULL, peakMatrix = NULL, exp_assay = "logcounts",
    peak_assay = "PeakMatrix", test = c("chi.sq", "binom"), clusters = NULL,
    exp_cutoff = 1, peak_cutoff = 0, regulon_cutoff = 0.05, p_adj = TRUE,
    prune_value = "pval", aggregateCells = FALSE, useDim = "IterativeLSI_ATAC",
    cellNum = 10, BPPARAM = BiocParallel::SerialParam(progressbar = TRUE)
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>regulon</td>
<td>A dataframe informing the gene regulatory relationship with the tf column representing transcription factors, idxATAC corresponding to the index in the peak- Matrix and target column representing target genes</td>
</tr>
</tbody>
</table>
The function prunes the network by performing tests of independence on the observed number of cells jointly expressing transcription factor (TF), regulatory element (RE) and target gene (TG) vs the expected number of cells if TF/RE and TG are independently expressed.

In other words, if no regulatory relationship exists, the expected probability of cells expressing all three elements is \( P(\text{TF, RE}) \times P(\text{TG}) \), that is, the product of (1) proportion of cells both expressing transcription factor and having accessible corresponding regulatory element, and (2) proportion of cells expressing target gene. The expected number of cells expressing all three elements is therefore \( n \times P(\text{TF, RE}) \times P(\text{TG}) \), where \( n \) is the total number of cells. However, if a TF-RE-TG relationship exists, we expect the observed number of cells jointly having all three elements (TF, RE, TG) to deviate from the expected number of cells predicted from an independent relationship.

If the user provides cluster assignment, the tests of independence are performed on a per-cluster basis in addition to providing all cells statistics. This enables pruning by cluster, and thus yields cluster-specific gene regulatory relationships.

We implement two tests, the binomial test and the chi-square test.
In the binomial test, the expected probability is $P(TF, RE) \times P(TG)$, and the number of trials is the number of cells, and the observed successes is the number of cells jointly expressing all three elements.

In the chi-square test, the expected probability for having all 3 elements active is also $P(TF, RE) \times P(TG)$ and the probability otherwise is $1 - P(TF, RE) \times P(TG)$. The observed cell count for the active category is the number of cells jointly expressing all three elements, and the cell count for the inactive category is $n - n_{\text{triple}}$.

**Value**

A DataFrame of pruned regulons with p-values indicating the probability of independence either for all cells or for individual clusters, z-score statistics for binomial tests or chi-square statistics for chi-square test and q-adjusted values.

**Author(s)**

Xiaosai Yao, Tomasz Wlodarczyk

**Examples**

```r
# create a mock singleCellExperiment object for gene expression matrix
set.seed(1000)
gene_sce <- scuttle::mockSCE()
gene_sce <- scuttle::logNormCounts(gene_sce)
rownames(gene_sce) <- paste0('Gene_', 1:2000)

# create a mock singleCellExperiment object for peak matrix
peak_gr <- GRanges(seqnames = 'chr1',
    ranges = IRanges(start = seq(from = 1, to = 10000, by = 100), width = 100))
peak_counts <- matrix(sample(x = 0:4, size = ncol(gene_sce) * length(peak_gr), replace = TRUE),
    nrow = length(peak_gr), ncol = ncol(gene_sce))
peak_sce <- SingleCellExperiment(list(counts = peak_counts), colData = colData(gene_sce))
rownames(peak_sce) <- paste0('Peak_', 1:100)

# create a mock regulon
regulon <- data.frame(tf = c(rep('Gene_1', 10), rep('Gene_2', 10)),
    idxATAC = sample(1:100, 20),
    target = c(paste0('Gene_', sample(3:2000, 10)),
        paste0('Gene_', sample(3:2000, 10))))

# prune regulon
pruned.regulon <- pruneRegulon(expMatrix = gene_sce,
    exp_assay = 'logcounts', peakMatrix = peak_sce, peak_assay = 'counts',
    regulon = regulon, clusters = gene_sce$Treatment, regulon_cutoff = 0.5)

# add weights with cell aggregation
gene_sce <- scater::runPCA(gene_sce)
pruned.regulon <- pruneRegulon(expMatrix = gene_sce, exp_assay = 'logcounts',
    peakMatrix = peak_sce, peak_assay = 'counts', regulon = regulon,
    clusters = gene_sce$Treatment, regulon_cutoff = 0.5,
    aggregateCells=TRUE, cellNum=3, useDim = 'PCA')
```
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