Package ‘AUCell’

May 29, 2024

Type Package

Title AUCell: Analysis of ‘gene set’ activity in single-cell RNA-seq data (e.g. identify cells with specific gene signatures)

Version 1.26.0

Date 2024-03-09

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Description AUCell allows to identify cells with active gene sets (e.g. signatures, gene modules...) in single-cell RNA-seq data. AUCell uses the ”Area Under the Curve” (AUC) to calculate whether a critical subset of the input gene set is enriched within the expressed genes for each cell. The distribution of AUC scores across all the cells allows exploring the relative expression of the signature. Since the scoring method is ranking-based, AUCell is independent of the gene expression units and the normalization procedure. In addition, since the cells are evaluated individually, it can easily be applied to bigger datasets, subsetting the expression matrix if needed.

URL http://scenic.aertslab.org

Imports DelayedArray,DelayedMatrixStats,data.table,graphics,grDevices,GSEABase,Matrix,methods,mixtools,R.utils,stats,SummarizedExperiment,BiocGenerics,utils

Enhances doMC,doRNG,doParallel,foreach

Suggests Biobase,BiocStyle,doSNOW,dynamicTreeCut,DT,GEOquery,knitr,NMF,plyr,R2HTML,rmarkdown,reshape2,plotly,Rtsne,testthat,zoo

License GPL-3

biocViews SingleCell,GeneSetEnrichment,Transcriptomics,Transcription,GeneExpression,WorkflowStep,Normalization

VignetteBuilder knitr

RoxygenNote 7.2.0

git_url https://git.bioconductor.org/packages/AUCell
Description

This class extends SummarizedExperiment to contain the AUC matrix and cell rankings (as 'assays').

The results are stored in the assays slot, but they can be accessed through the regular methods (i.e. nrow, rownames...)

Types:
- "AUC": The assays contains the AUC for the gene-sets (or region-sets) & cells.
- "ranking": The assays contains the gene rankings for each cell.
Usage

```r
## S4 method for signature 'aucellResults'
show(object)

getAUC(object)

## S4 method for signature 'aucellResults'
getAUC(object)

getRanking(object)

## S4 method for signature 'aucellResults'
getRanking(object)

## S4 method for signature 'aucellResults'
cbind(..., deparse.level = 1)

## S4 method for signature 'aucellResults'
rbind(..., deparse.level = 1)
```

Arguments

- `object`: Results from `AUCell_buildRanking`
- `...` (Only for `cbind`)
  or `AUCell_calcAUC`.
- `deparse.level` (Only for `cbind`)

Value

- `show`: Prints a summary of the object
- `getAUC`: Returns the matrix containing the AUC
- `getRanking`: Returns the matrix containing the rankings
- `cbind`: Combines objects by columns (`cbind` on assays); other other slots are conserved from the first object provided as argument. Both, ranking and AUC are calculated by column (cell or sample). Therefore, it is fine to merge objects as long as they come from equivalent datasets (and keep same genes/genesets, etc...)

Examples

```r
# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

############# Fake run of AUCell #############
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="\n")

dim(exprMatrix)

# Running AUCell
cells_rankings <- AUCell_buildRankings(exprMatrix)

fewGenes <- sample(rownames(exprMatrix), 10)
otherGenes <- sample(rownames(exprMatrix), 5)
geneSets <- list(geneSet1=fewGenes,
                 geneSet2=otherGenes)
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5, nCores=1)

# Exploring the output:
cells_AUC

class(cells_AUC)

# Extracting the AUC matrix:
getAUC(cells_AUC)[,1:5]

# Subsetting and regular manipulation methods are also available:
cells_AUC[1:2,]
cells_AUC[3:4]
dim(cells_AUC)
nrow(cells_AUC)
ncol(cells_AUC)
colnames(cells_AUC)
rownames(cells_AUC)

### Merging 2 objects (ranking or AUC):
sample1 <- cells_AUC[10:20]
sample2 <- cells_AUC[100:140]
cbind(sample1, sample2)

---

**AUCcell_assignCells**

**Description**

Assigns whether the gene sets are considered "active" on each cell based on the given thresholds

**Usage**

\[
\text{AUCcell_assignCells}(\text{cellsAUC}, \text{thresholds}, \text{nCores} = 1)
\]

**Arguments**

- **cellsAUC**: AUC object returned by `AUCcell_calcAUC`.
- **thresholds**: Thresholds selected for each geneset (named vector).
- **nCores**: Number of cores to use for computation.
**AUC_all_assignCells**

**Value**

List with the following elements for each gene-set:

- `'aucThr'` threshold value, in the same format as `AUCCell_exploreThresholds()`
- `'assignment'` List of cells that pass the selected AUC threshold

**See Also**

Previous step in the workflow: `AUCCell_calcAUC` and optionally `AUCCell_exploreThresholds`

See the package vignette for examples and more details: `vignette("AUCell")`

**Examples**

```r
# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

########### Fake expression matrix ###########
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))
dim(exprMatrix)

########### Previous steps in the workflow ###########
# Step 1.
cells_rankings <- AUCCell_buildRankings(exprMatrix, plotStats=FALSE)
# Step 2.
# (Gene sets: random genes)
geneSets <- list(geneSet1=sample(rownames(exprMatrix), 10),
geneSet2=sample(rownames(exprMatrix), 5))
cells_AUC <- AUCCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5)

############# Step 3: Assign cells ##############
# 1. Plot histograms and obtain some pre-computed thresholds
# (this example is only meant to show the interface/arguments of the function,
# see the vignette for meaningful examples)
set.seed(123)
par(mfrow=c(1,2)) # Plot is divided into one row and two columns
thresholds <- AUCCell_exploreThresholds(cells_AUC, plotHist=TRUE)
thresholds$geneSet1$aucThr

# 2. Obtain cells over a given threshold:
names(which(getAUC(cells_AUC)["geneSet1",] > 0.19))

# Alternative 1: assign cells according to the 'automatic' threshold
cells_assignment <- AUCCell_exploreThresholds(cells_AUC,}
```
# Cells assigned:
getAssignments(cells_assignment)

# Threshold applied:
getThresholdSelected(cells_assignment)

# Alternative 2: choose a threshold manually and assign cells
newThresholds = getThresholdSelected(cells_assignment)
newThresholds['geneSet1'] = 0.8
newAssignments = AUCell_assignCells(cells_AUC, newThresholds)
getAssignments(newAssignments)

---

**AUCell_buildRankings**  
*Build gene expression rankings for each cell*

**Description**

Builds the "rankings" for each cell: expression-based ranking for all the genes in each cell. The genes with same expression value are shuffled. Therefore, genes with expression '0' are randomly sorted at the end of the ranking.

These "rankings" can be seen as a new representation of the original dataset. Once they are calculated, they can be saved for future analyses.

**Usage**

```r
AUCell_buildRankings(
  exprMat,
  featureType = "genes",
  plotStats = TRUE,
  splitByBlocks = FALSE,
  BPPARAM = NULL,
  keepZeroesAsNA = FALSE,
  verbose = TRUE,
  nCores = NULL,
  mctype = NULL,
  ...
)
```

```r
## S4 method for signature 'dgCMatrix'
AUCell_buildRankings(
  exprMat,
  featureType = "genes",
  plotStats = TRUE,
  splitByBlocks = TRUE,
  ...
)
```
AUCell_buildRankings

BPPARAM = NULL,
keepZeroesAsNA = FALSE,
verbose = TRUE,
nCores = NULL,
mctype = NULL
)

## S4 method for signature 'matrix'
AUCell_buildRankings(
  exprMat,
  featureType = "genes",
  plotStats = TRUE,
  splitByBlocks = FALSE,
  BPPARAM = NULL,
  keepZeroesAsNA = FALSE,
  verbose = TRUE,
  nCores = NULL,
  mctype = NULL
)

## S4 method for signature 'ExpressionSet'
AUCell_buildRankings(
  exprMat,
  featureType = "genes",
  plotStats = TRUE,
  splitByBlocks = FALSE,
  BPPARAM = NULL,
  keepZeroesAsNA = FALSE,
  verbose = TRUE,
  nCores = NULL,
  mctype = NULL
)

## S4 method for signature 'SummarizedExperiment'
AUCell_buildRankings(
  exprMat,
  featureType = "genes",
  plotStats = TRUE,
  splitByBlocks = FALSE,
  BPPARAM = NULL,
  keepZeroesAsNA = FALSE,
  verbose = TRUE,
  assayName = NULL,
  nCores = NULL,
  mctype = NULL
)
Arguments

exprMat  
Expression matrix (genes as rows, cells as columns) The expression matrix can also be provided as one of the R/Bioconductor classes:
  - dgCMatrix-class: Sparse matrix
  - RangedSummarizedExperiment and derived classes (e.g. SingleCellExperiment): The matrix will be obtained through assay(exprMatrix), which will extract the first assay (usually the counts) or the assay name given in \'assayName\'
  - ExpressionSet: The matrix will be obtained through exprs(exprMatrix)

featureType  
Name for the rows (e.g. "genes"). Only for naming the rankings, not used internally.

plotStats  
Should the function plot the expression boxplots/histograms? (TRUE / FALSE). These plots can also be produced with the function \texttt{plotGeneCount}.

splitByBlocks  
Whether to split the matrix by blocks in the ranking calculation. Allows using multiple cores. FALSE by default. If using sparse matrices it is automatically set to TRUE.

BPPARAM  
Set to use multiple cores. Only used if \'splitByBlocks=TRUE\'

keepZeroesAsNA  
Convert zeroes to NA instead of locating randomly at the end of the ranking.

verbose  
Should the function show progress messages? (TRUE / FALSE)

nCores  
Deprecated

mctype  
Deprecated

...  
Other arguments

assayName  
Name of the assay containing the expression matrix (e.g. in \texttt{SingleCellExperiment} objects)

Details

It is important to check that most cells have at least the number of expressed/detected genes that are going to be used to calculate the AUC (\'aucMaxRank\' in \texttt{calcAUC()}). The histogram provided by \texttt{\textquote{AUCCell\_buildRankings()}} allows to quickly check this distribution. \texttt{\textquote{plotGeneCount(exprMatrix)}} allows to obtain only the plot before building the rankings.

Value

Ranking of the feature within the cell (features as rows, cells as columns)

See Also

Next step in the workflow: \texttt{AUCCell\_calcAUC}.

See the package vignette for examples and more details: \texttt{vignette("AUCCell")}
Examples

# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

############# Fake expression matrix #############
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))

##################################################
cells_rankings <- AUCell_buildRankings(exprMatrix, plotStats=TRUE)
cells_rankings

AUCell_calcAUC  Calculate AUC

Description

Calculates the 'AUC' for each gene-set in each cell.

Usage

AUCell_calcAUC(
geneSets,
rankings,
nCores = 1,
normAUC = TRUE,
aucMaxRank = ceiling(0.05 * nrow(rankings)),
verbose = TRUE
)

## S4 method for signature 'list'
AUCell_calcAUC(
geneSets,
rankings,
nCores = 1,
normAUC = TRUE,
aucMaxRank = ceiling(0.05 * nrow(rankings)),
verbose = TRUE
)

## S4 method for signature 'character'
AUCell_calcAUC(
geneSets,
rankings,
nCores = 1,
normAUC = TRUE,
aucMaxRank = ceiling(0.05 * nrow(rankings)),
verbose = TRUE
)

## S4 method for signature 'GeneSet'
AUCell_calcAUC(
  geneSets,
  rankings,
  nCores = 1,
  normAUC = TRUE,
  aucMaxRank = ceiling(0.05 * nrow(rankings)),
  verbose = TRUE
)

## S4 method for signature 'GeneSetCollection'
AUCell_calcAUC(
  geneSets,
  rankings,
  nCores = 1,
  normAUC = TRUE,
  aucMaxRank = ceiling(0.05 * nrow(rankings)),
  verbose = TRUE
)

Arguments

- **geneSets**: List of gene-sets (or signatures) to test in the cells. The gene-sets should be provided as `GeneSet, GeneSetCollection` or character list (see examples).
- **rankings**: 'Rankings' created for this dataset with `AUCell_buildRankings`.
- **nCores**: Number of cores to use for computation.
- **normAUC**: Wether to normalize the maximum possible AUC to 1 (Default: TRUE).
- **aucMaxRank**: Threshold to calculate the AUC (see 'details' section)
- **verbose**: Should the function show progress messages? (TRUE / FALSE)

Details

In a simplified way, the AUC value represents the fraction of genes, within the top X genes in the ranking, that are included in the signature. The parameter 'aucMaxRank' allows to modify the number of genes (maximum ranking) that is used to perform this computation. By default, it is set to 5% of the total number of genes in the rankings. Common values may range from 1 to 20%.

Value

Matrix with the AUC values (gene-sets as rows, cells as columns).
**See Also**

Previous step in the workflow: `AUCell_buildRankings`. Next step in the workflow: `AUCell_exploreThresholds`. See the package vignette for examples and more details: `vignette("AUCell")`

**Examples**

# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

```
#---------------- Fake expression matrix ----------------#
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))

#---------------- Previous step in the workflow ----------------#
# Step 1.
cells_rankings <- AUCell_buildRankings(exprMatrix)

#---------------- Step 2: Calculate AUC ----------------#
# In this example we use two gene sets: 10 and 5 random genes
# (see other formatting examples at the end)
fewGenes <- sample(rownames(exprMatrix), 10)
otherGenes <- sample(rownames(exprMatrix), 5)
geneSets <- list(geneSet1=fewGenes,
                  geneSet2=otherGenes)
geneSets

# Calculate AUC with the rankings from Step 1.
# To be able to run this fake example (which contain only 20 genes),
# we use aucMaxRank=5 (top 25% of the genes in the ranking)
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5, nCores=1)

cells_AUC

deAUC(cells_AUC)[1:2,]
deAUC(cells_AUC)[3:4]
getAUC(cells_AUC)[,1:5]

# These methods are also available:
dim(cells_AUC)
nrow(cells_AUC)
ncol(cells_AUC)
colnames(cells_AUC)[1:4]
```
rownames(cells_AUC)

# Alternatives for the input of gene sets:

# a) Character vector (i.e. only one gene-set)
# It will take the default name 'geneSet'
rowGenes <- AUCell_calcAUC(fewGenes, cells_rankings, aucMaxRank=5)

# b) List
geneSets <- list(geneSet1=fewGenes,
              geneSet2=otherGenes)
geneSets <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5)

# c) GeneSet object (from GSEABase)
library(GSEABase)
geneSetOne <- GeneSet(fewGenes, setName="geneSetOne")
geneSetOne <- AUCell_calcAUC(geneSetOne, cells_rankings, aucMaxRank=5)

# d) GeneSetCollection object (from GSEABase)
geneSetTwo <- GeneSet(otherGenes, setName="geneSetTwo")
geneSets <- GeneSetCollection(geneSetOne, geneSetTwo)
geneSets <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5)

**Usage**

```r
AUCell_exploreThresholds(
cellsAUC,
  thrP = 0.01,
  nCores = 1,
  smallestPopPercent = 0.25,
  plotHist = TRUE,
  densAdjust = 2,
  assignCells = FALSE,
  nBreaks = 100,
)```
AUCell_exploreThresholds

verbose = TRUE

getThresholdSelected(aucellThresholds)

getAssignments(aucellThresholds)

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>cellsAUC</td>
<td>AUC object returned by AUCell_calcAUC.</td>
</tr>
<tr>
<td>thrP</td>
<td>Probability to determine outliers in some of the distributions (see 'details' section). By default it is set to 1% (thrP): if there are 3000 cells in the dataset, it is expected that approximately 30 cells are over this threshold if the AUC is normally distributed.</td>
</tr>
<tr>
<td>nCores</td>
<td>Number of cores to use for computation.</td>
</tr>
<tr>
<td>smallestPopPercent</td>
<td>Size (percentage) of the smallest population of cells expected. Used to calculate some of the thresholds.</td>
</tr>
<tr>
<td>plotHist</td>
<td>Whether to plot the AUC histograms. (TRUE / FALSE)</td>
</tr>
<tr>
<td>densAdjust</td>
<td>Parameter for the density curve. (See density for details).</td>
</tr>
<tr>
<td>assignCells</td>
<td>Return the list of cells that pass the automatically selected threshold? (TRUE/FALSE)</td>
</tr>
<tr>
<td>nBreaks</td>
<td>Number of bars to plot in the histograms.</td>
</tr>
<tr>
<td>verbose</td>
<td>Should the function show progress messages? (TRUE / FALSE)</td>
</tr>
<tr>
<td>aucellThresholds</td>
<td>For aux functions: Output from AUCell_exploreThresholds</td>
</tr>
</tbody>
</table>

**Details**

To ease the selection of an assignment threshold, this function adjusts the AUCs of each gene-set to several distributions and calculates possible thresholds:

- **minimumDens** (plot in Blue): Inflection point of the density curve. This is usually a good option for the ideal situation with bimodal distributions. To avoid false positives, by default this threshold will not be chosen if the second distribution is higher (i.e. the majority of cells have the gene-set "active").
- **L_k2** (plot in Red): Left distribution, after adjusting the AUC to a mixture of two distributions. The threshold is set to the right (prob: 1-(thrP/nCells)). Only available if 'mixtools' package is installed.
- **R_k3** (plot in Pink): Right distribution, after adjusting the AUC to a mixture of three distributions. The threshold is set to the left (prob: thrP). Only available if 'mixtools' package is installed.
- **Global_k1** (plot in Grey): "global" distribution (i.e. mean and standard deviations of all cells). The threshold is set to the right (prob: 1-(thrP/nCells)). The threshold based on the global distribution is ignored from the automatic selection unless the mixed models are overlapping.
Note: If assignCells=TRUE, the highest threshold is used to select cells. However, keep in mind that this function is only meant to ease the selection of the threshold, and we highly recommend to look at the AUC histograms and adjust the threshold manually if needed. We recommend to be specially aware on gene-sets with few genes (10-15) and thresholds that are set extremely low.

Value

List with the following elements for each gene-set:

- `'aucThr'` Thresholds calculated with each method (see 'details' section), and the number of cells that would be assigned using that threshold.
  
  If assignCells=TRUE, the threshold selected automatically is the highest value (in most cases, excluding the global distribution).

- `'assignment'` List of cells that pass the selected AUC threshold (if assignCells=TRUE)

If plotHist=TRUE the AUC histogram is also plot, including the distributions calculated and the corresponding thresholds in the same color (dashed vertical lines). The threshold that is automatically selected is shown as a thicker non-dashed vertical line.

See Also

Previous step in the workflow: `AUCell_calcAUC`.

See the package vignette for examples and more details: vignette("AUCell")

Examples

```r
# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

#################### Fake expression matrix ######################
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))
dim(exprMatrix)

########################################################################

########## Previous steps in the workflow ##########

# Step 1.
cells_rankings <- AUCell_buildRankings(exprMatrix, plotStats=FALSE)

# Step 2.
# (Gene sets: random genes)
geneSets <- list(geneSet1=sample(rownames(exprMatrix), 10),
geneSet2=sample(rownames(exprMatrix), 5))
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5)

########################################################################

########## Step 3: Assign cells ##########
```
# 1. Plot histograms and obtain some pre-computed thresholds
# (this example is only meant to show the interface/arguments of the function,
# see the vignette for meaningful examples)
set.seed(123)
par(mfrow=c(1,2))  # Plot is divided into one row and two columns
thresholds <- AUCell_exploreThresholds(cells_AUC, plotHist=TRUE)
thresholds$geneSet1$aucThr

# 2. Obtain cells over a given threshold:
names(which(getAUC(cells_AUC)["geneSet1",] > 0.19))

# Alternative 1: assign cells according to the 'automatic' threshold
cells_assignment <- AUCell_exploreThresholds(cells_AUC,
                                           plotHist=FALSE, assignCells=TRUE)

# Cells assigned:
getAssignments(cells_assignment)
# Threshold applied:
getThresholdSelected(cells_assignment)

# Alternative 2: choose a threshold manually and assign cells
newThresholds = getThresholdSelected(cells_assignment)
newThresholds['geneSet1'] = 0.8
newAssignments = AUCell_assignCells(cells_AUC, newThresholds)
getAssignments(newAssignments)

AUCell_plotHist  
Plot AUC histogram

**Description**

Plots the distribution of AUC across the cells (for each gene-set) as an histogram.

**Usage**

```r
AUCell_plotHist(
  cellsAUC,
  aucThr = max(cellsAUC),
  nBreaks = 100,
  onColor = "dodgerblue4",
  offColor = "slategray2",
  ...
)
```

**Arguments**

- `cellsAUC`  
  Subset of the object returned by `AUCell_calcAUC` (i.e. including only the gene-sets to plot)

aucThr  AUC value planned to use as threshold (to make sure the X axis includes it), if any. Otherwise, the X axis extends to cover only the AUC values plotted.
nBreaks  Number of 'bars' to plot (breaks argument for hist function).
onColor  Color for the bars that pass the AUC threshold
offColor  Color for the bars that do not pass the AUC threshold
...  Other arguments to pass to hist function.

Value
List of histogram objects (invisible).

See Also
See the package vignette for examples and more details: vignette("AUCell")

Examples

# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

########### Fake expression matrix #############
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))
dim(exprMatrix)

##################################################

############# Begining of the workflow #############
# Step 1.
cells_rankings <- AUCell_buildRankings(exprMatrix, plotStats=FALSE)

# Step 2.
# (Gene set: 10 random genes)
genes <- sample(rownames(exprMatrix), 10)
geneSets <- list(geneSet1=genes)
# (aucMaxRank=5 to run with this fake example, it will return 'high' AUC values)
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5)

##################################################

# Plot histogram:
AUCell_plotHist(cells_AUC["geneSet1",], nBreaks=10)
**Description**

Plots the AUC histogram and t-SNE coloured by AUC, binary activity and TF expression

**Usage**

```r
AUCell_plotTSNE(
  tSNE, 
  exprMat = NULL, 
  cellsAUC = NULL, 
  thresholds = NULL, 
  reorderGeneSets = FALSE, 
  cex = 1, 
  alphaOn = 1, 
  alphaOff = 0.2, 
  borderColor = adjustcolor("lightgray", alpha.f = 0.1), 
  offColor = "lightgray", 
  plots = c("histogram", "binaryAUC", "AUC", "expression"), 
  exprCols = c("goldenrod1", "darkorange", "brown"), 
  asPNG = FALSE, 
  ... 
)
```

**Arguments**

- `tSNE`  
  t-SNE coordinates (e.g. `tSNE$Y`)
- `exprMat`  
  Expression matrix
- `cellsAUC`  
  AUC (as returned by calcAUC)
- `thresholds`  
  Thresholds returned by AUCell
- `reorderGeneSets`  
  Whether to reorder the gene sets based on AUC similarity
- `cex`  
  Scaling factor for the dots in the scatterplot
- `alphaOn`  
  Transparency for the dots representing "active" cells
- `alphaOff`  
  Transparency for the dots representing "inactive" cells
- `borderColor`  
  Border color for the dots (scatterplot)
- `offColor`  
  Color for the dots representing "inactive" cells
- `plots`  
  Which plots to generate? Select one or multiple: `plots=c("histogram", "binaryAUC", "AUC", "expression")`
- `exprCols`  
  Color scale for the expression
- `asPNG`  
  Output each individual plot in a .png file? (can also be a directory)
- `...`  
  Other arguments to pass to `hist` function.
Details
To avoid calculating thresholds, set thresholds to FALSE

Value
Returns invisible: cells_trhAssignment

See Also
List of vignettes included in the package: vignette(package="AUCell")

Examples

```
####
# Fake run of AUCell
set.seed(123)
eexprMatrix <- matrix(
  data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
  nrow=20,
  dimnames=list(paste("Gene", 1:20, sep=""),
               paste("Cell", 1:500, sep=""))
geneSets <- list(geneSet1=sample(rownames(eexprMatrix), 10),
                 geneSet2=sample(rownames(eexprMatrix), 5))

cells_rankings <- AUCell_buildRankings(eexprMatrix, plotStats = FALSE)
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5, nCores=1)
selectedThresholds <- rowMeans(getAUC(cells_AUC))
cellsTsne<- Rtsne::Rtsne(t(exprMatrix),max_iter = 10)$Y
rownames(cellsTsne) <- colnames(exprMatrix)
####

par(mfrow=c(2,3))
ths <- AUCell_plotTSNE(tSNE=cellsTsne, exprMat=NULL,
cellsAUC=cells_AUC, thresholds=selectedThresholds,
plots = c("histogram", "binaryAUC", "AUC"))

####
# Color based on the known phenodata:
cellInfo <- data.frame(cellType1=sample(LETTERS[1:3],ncol(exprMatrix), replace=TRUE),
cellType2=sample(letters[5:7],ncol(exprMatrix), replace=TRUE),
nGenes=abs(rnorm(ncol(exprMatrix))),
row.names=colnames(exprMatrix))
colVars <- list(cellType2=setNames(c("skyblue", "magenta", "darkorange"),letters[5:7]))
# dev.off()
plotTsnecellProps(cellsTsne, cellInfo, colVars=colVars)
```
Run AUCell

Description

Runs AUCell (calculates the ranking + score genesets)

Usage

AUCell_run(
  exprMat,
  geneSets,
  featureType = "genes",
  keepZeroesAsNA = FALSE,
  normAUC = TRUE,
  aucMaxRank = ceiling(0.05 * nrow(exprMat)),
  BPPARAM = NULL,
  ...
)

## S4 method for signature 'dgCMatrix'
AUCell_run(
  exprMat,
  geneSets,
  featureType = "genes",
  keepZeroesAsNA = FALSE,
  normAUC = TRUE,
  aucMaxRank = ceiling(0.05 * nrow(exprMat)),
  BPPARAM = NULL
)

## S4 method for signature 'matrix'
AUCell_run(
  exprMat,
  geneSets,
  featureType = "genes",
  keepZeroesAsNA = FALSE,
  normAUC = TRUE,
  aucMaxRank = ceiling(0.05 * nrow(exprMat)),
  BPPARAM = NULL
)

## S4 method for signature 'SummarizedExperiment'
AUCell_run(
  exprMat,
  geneSets,
  featureType = "genes",
  keepZeroesAsNA = FALSE,
Arguments

exprMat  Expression matrix (genes/regions as rows, cells as columns) The expression matrix can also be provided as one of the R/Bioconductor classes:
  • dgCMatrix-class: Sparse matrix
  • RangedSummarizedExperiment and derived classes (e.g. SingleCellExperiment): The matrix will be obtained through assay(exprMatrix), -which will extract the first assay (usually the counts)- or the assay name given in 'assayName'

geneSets  List of gene-sets (or signatures) to test in the cells. The gene-sets should be provided as GeneSet, GeneSetCollection or character list (see examples).

featureType  Name for the rows (e.g. "genes"). Only for naming the rankings, not used internally.

keepZeroesAsNA  Convert zeroes to NA instead of locating randomly at the end of the ranking.

normAUC  Whether to normalize the maximum possible AUC to 1 (Default: TRUE).

aucMaxRank  Threshold to calculate the AUC (see 'details' section)

BPPARAM  Set to use multiple cores. Only used if 'splitByBlocks=TRUE'

...  Other arguments

assayName  Name of the assay containing the expression matrix (e.g. in SingleCellExperiment objects)

Details

In a simplified way, the AUC value represents the fraction of genes, within the top X genes in the ranking, that are included in the signature. The parameter 'aucMaxRank' allows to modify the number of genes (maximum ranking) that is used to perform this computation. By default, it is set to 5% of the total number of genes in the rankings. Common values may range from 1 to 20%.

Value

Matrix with the AUC values (gene-sets as rows, cells as columns).

See Also

Includes AUCell_buildRankings and AUCell_calcAUC. Next step in the workflow: AUCell_exploreThresholds. See the package vignette for examples and more details: vignette("AUCell")
Examples

# This example is run using a fake expression matrix.
# Therefore, the output will be meaningless.

############# Fake expression matrix #############
set.seed(123)
exprMatrix <- matrix(data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
paste("Cell", 1:500, sep="")))
exprMatrix <- as(exprMatrix, "dgCMatrix")

# In this example we use two gene sets: 10 and 5 random genes
# (see other formatting examples at the end)
fewGenes <- sample(rownames(exprMatrix), 10)
otherGenes <- sample(rownames(exprMatrix), 5)

geneSets <- list(geneSet1=fewGenes,
geneSet2=otherGenes)
geneSets

# Calculate AUCell score for the genes in the sets
# To be able to run this fake example (which contain only 20 genes),
# we use aucMaxRank=5 (top 25% of the genes in the ranking)
cells_AUC <- AUCell_run(exprMatrix, geneSets, aucMaxRank=5)

## To run in parallel:
# cells_AUC <- AUCell_run(exprMatrix, geneSets, aucMaxRank=5,
# BPPARAM=BiocParallel::MulticoreParam(5))

# Format of the output:
cells_AUC

# To subset & access the AUC slot (as matrix):
cells_AUC[1:2,]
cells_AUC[,3:4]
getAUC(cells_AUC)[,1:5]

# These methods are also available:
dim(cells_AUC)
nrow(cells_AUC)
ncol(cells_AUC)
colnames(cells_AUC)[1:4]
rownames(cells_AUC)

#########################################################
# Alternatives for the input of gene sets:
# a) Character vector (i.e. only one gene-set)
# It will take the default name 'geneSet'
fewGenes
test <- AUCell_run(exprMatrix, fewGenes, aucMaxRank=5)

# b) List
geneSets <- list(geneSet1=fewGenes,
geneSet2=otherGenes)
geneSets
test <- AUCell_run(exprMatrix, fewGenes, aucMaxRank=5)

# c) GeneSet object (from GSEABase)
library(GSEABase)
geneSetOne <- GeneSet(fewGenes, setName="geneSetOne")
geneSetOne
test <- AUCell_run(exprMatrix, fewGenes, aucMaxRank=5)

# d) GeneSetCollection object (from GSEABase)
geneSetTwo <- GeneSet(otherGenes, setName="geneSetTwo")
geneSets <- GeneSetCollection(geneSetOne, geneSetTwo)
geneSets
test <- AUCell_run(exprMatrix, fewGenes, aucMaxRank=5)

getSetNames

## Description

Get gene set name (excluding number of genes/regions after space)

## Usage

getSetNames(aucMat, patterns, startChr = "^", endChr = "\_")

## Arguments

- `aucMat`: AUC matrix
- `patterns`: patterns
- `startChr`: Character to indicate the start (typically "^")
- `endChr`: Character at the end of the gene set name, i.e. space or "\_" after a transcription factor name

## Value

Returns the gene set name (i.e. selects the given pattern)
### Description

Functions to manipulate `GeneSet` and `GeneSetCollection` objects (from package GSEABase)

### Usage

```r
nGenes(geneSet)

## S4 method for signature 'GeneSet'
nGenes(geneSet)

## S4 method for signature 'GeneSetCollection'
nGenes(geneSet)

subsetGeneSets(geneSets, geneNames)

## S4 method for signature 'GeneSetCollection'
subsetGeneSets(geneSets, geneNames)

setGeneSetNames(geneSets, newNames)

## S4 method for signature 'GeneSetCollection'
setGeneSetNames(geneSets, newNames)
```

### Arguments

- `geneSet` One gene-set (`GeneSet`)
- `geneSets` Gene-set collection (`GeneSetCollection`)
- `geneNames` Gene names (for subset)
- `newNames` New names (to assign to the gene sets)

### Value

- **nGenes()**: provides the number of genes in the gene-set, or each of the gene-sets in a collection.
- **subsetGeneSets()**: Subsets each of the gene-sets in a collection to contain only the genes in the given list. Equivalent to `intersect()`, but keeping the original gene-set name.
- **setGeneSetNames()**: Modifies the name of each gene-set in a collection.
Examples

```r
library(GSEABase)
genes_1 <- GeneSet(paste("Gene", 1:20, sep=""), setName="geneSet1")
genes_2 <- GeneSet(paste("Gene", 18:22, sep=""), setName="geneSet2")
geneSets <- GeneSetCollection(genes_1, genes_2)

nGenes(genes_1)
nGenes(geneSets)

subsetGeneSets(geneSets, paste("Gene", 15:20, sep=""))

geneSets_newNames <- setGeneSetNames(geneSets, c("one", "two"))
names(geneSets_newNames)
```

orderAUC

Description

Reorder the gene-sets based on AUC similarity

Usage

```r
orderAUC(auc)
```

Arguments

- **auc**: AUC (as returned by calcAUC)

Value

Gene-set names in the suggested order

Examples

```r
# cellsAUC <- cellsAUC[orderAUC(cellsAUC),]
```
Description

Colors the embeddings (t-SNE/Umap) based on the activity of 3 (groups of) geneSets

Usage

plotEmb_rgb(
  aucMat,
  embedding,
  geneSetsByCol,
  aucType = "AUC",
  aucMaxContrast = 0.8,
  offColor = "#c0c0c030",
  showPlot = TRUE,
  showLegend = TRUE,
  ...
)

Arguments

- **aucMat**: AUC matrix (continuous or binary)
- **embedding**: AUC matrix (continuous or binary)
- **geneSetsByCol**: Gene sets to plot
- **aucType**: "AUC" or "Binary"
- **aucMaxContrast**: To increase the AUC contrast decrease the value.
- **offColor**: Color por the cells completely off. To deactivate (color as black), set to NULL.
- **showPlot**: Whether to plot the coloured embeddings.
- **showLegend**: Whether to plot add a legend to the plot.
- **...**: Other arguments to pass to the plot function.

Value

The cell colors (invisible)
plotGeneCount

Description

Plots a histogram and boxplot for the number of genes detected in each cell.

Usage

plotGeneCount(exprMat, plotStats = TRUE, verbose = TRUE)

Arguments

exprMat Expression matrix (genes as rows, cells as columns)
plotStats Logical. If true, it plots the histogram, otherwise only calculates the percentages of genes detected.
verbose Should the function show progress messages? (TRUE / FALSE)

Details

It is important to check that most cells have at least the number of expressed/detected genes that are going to be used to calculate the AUC ('aucMaxRank' in 'calcAUC()'). The histogram provided by 'AUCell_buildRankings()' allows to quickly check this distribution. 'plotGeneCount(exprMatrix)' allows to obtain only the plot before building the rankings.

Value

Quantiles with the number of genes detected by cell (invisible). This result is also printed if verbose=TRUE.

See Also

See the package vignette for more details: vignette("AUCell")

Examples

### (Fake expression matrix)
exprMatrix <- matrix(sample(c(rep(0, 500), sample(1:3, 500, replace=TRUE))), nrow=20)
rownames(exprMatrix) <- paste("Gene", 1:20, sep="")
colnames(exprMatrix) <- paste("Sample", 1:50, sep="")
###
plotGeneCount(exprMatrix)
title(sub="Fake expression matrix")
Description

Plots the t-SNE coloured based on the known cell properties

Usage

```r
plotTsne_cellProps(
  tSNE,
  cellInfo,
  colVars = NULL,
  cex = 1,
  sub = "",
  gradientCols = c("yellow", "orange", "red"),
  showLegend = TRUE
)
```

Arguments

- `tSNE`: t-SNE coordinates (e.g. `tSNE$Y`)
- `cellInfo`: Dataframe with cell phenodata
- `colVars`: Colors for the cell properties (optional)
- `cex`: Scaling factor for the dots in the scatterplot
- `sub`: Subtitle (e.g. tSNE type)
- `gradientCols`: Gradient colors for numerical variables
- `showLegend`: Whether to show the legend

Value

Plots the t-SNE

Examples

```r
# Fake run of AUCell
set.seed(123)
exprMatrix <- matrix(
data=sample(c(rep(0, 5000), sample(1:3, 5000, replace=TRUE))),
nrow=20,
dimnames=list(paste("Gene", 1:20, sep=""),
            paste("Cell", 1:500, sep="")))
geneSets <- list(geneSet1=sample(rownames(exprMatrix), 10),
                 geneSet2=sample(rownames(exprMatrix), 5))
```
cells_rankings <- AUCell_buildRankings(exprMatrix, plotStats = FALSE)
cells_AUC <- AUCell_calcAUC(geneSets, cells_rankings, aucMaxRank=5, nCores=1)
selectedThresholds <- rowMeans(getAUC(cells_AUC))
cellsTsne<- Rtsne::Rtsne(t(exprMatrix),max_iter = 10)$Y
# cellsTsne<- tsne::tsne(t(exprMatrix),max_iter = 10)
rownames(cellsTsne) <- colnames(exprMatrix)

par(mfrow=c(2,3))
ths <- AUCell_plotTSNE(tSNE=cellsTsne, exprMat=NULL, cellsAUC=cells_AUC, thresholds=selectedThresholds, plots = c("histogram", "binaryAUC", "AUC"))

# Color based on the known phenodata:
cellInfo <- data.frame(cellType1=sample(LETTERS[1:3],ncol(exprMatrix), replace=TRUE),
cellType2=sample(letters[5:7],ncol(exprMatrix), replace=TRUE),
nGenes=abs(rnorm(ncol(exprMatrix))),
row.names=colnames(exprMatrix))
colVars <- list(cellType2=setNames(c("skyblue", "magenta", "darkorange"),letters[5:7]))
# dev.off()
plotTsne_cellProps(cellsTsne, cellInfo, colVars=colVars)

dev.off()

updateAucellResults

updateAucellResults  Update AUCell results

Description
Updates the AUC scores provided by AUCell from a previous version.

Usage
updateAucellResults(oldAucObject, objectType = "AUC")

Arguments
oldAucObject  Object to update
objectType  Either "AUC" or "ranking" indicating the object type

Value
Updated version of the object as aucellResults.
Examples

```r
oldAuc <- matrix(data=1:2000, nrow=50, ncol=40)
updateAucellResults(oldAuc)
```
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