

Package ‘smartid’

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Title Scoring and Marker Selection Method Based on Modified TF-IDF

Version 1.3.2

Description This package enables automated selection of group specific signature, especially for rare population. The package is developed for generating specific lists of signature genes based on Term Frequency-Inverse Document Frequency (TF-IDF) modified methods. It can also be used as a new gene-set scoring method or data transformation method. Multiple visualization functions are implemented in this package.

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'smartid-package.R' 'score-methods.R' 'select_markers.R'
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cal_score	<i>calculate combined score</i>
-----------	---------------------------------

Description

compute TF (term/feature frequency), IDF (inverse document/cell frequency), IAE (inverse average expression of features) and combine the the final score

Usage

```
cal_score(  
  data,  
  tf = c("logtf", "tf"),  
  idf = "prob",  
  iae = "prob",  
  slot = "counts",  
  new.slot = "score",  
  par.idf = NULL,  
  par.iae = NULL  
)  
  
## S4 method for signature 'AnyMatrix'  
cal_score(  
  data,  
  tf = c("logtf", "tf"),  
  idf = "prob",  
  iae = "prob",  
  par.idf = NULL,  
  par.iae = NULL  
)  
  
## S4 method for signature 'SummarizedExperiment'  
cal_score(  
  data,  
  tf = c("logtf", "tf"),  
  idf = "prob",  
  iae = "prob",  
  slot = "counts",  
  new.slot = "score",  
  par.idf = NULL,  
  par.iae = NULL  
)
```

Arguments

data	an expression object, can be matrix or SummarizedExperiment
tf	a character, specify the TF method to use, can be "tf" or "logtf"

idf	a character, specify the IDF method to use. Available methods can be accessed using <code>idf_iae_methods()</code>
iae	a character, specify the IAE method to use. Available methods can be accessed using <code>idf_iae_methods()</code>
slot	a character, specify which slot to use when data is se object, optional, default 'counts'
new.slot	a character, specify the name of slot to save score in se object, optional, default 'score'
par.idf	other parameters for specified IDF methods
par.iae	other parameters for specified IAE methods

Value

A list of matrices or se object containing combined score

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
cal_score(
  data,
  par.idf = list(label = sample(c("A", "B"), 10, replace = TRUE)),
  par.iae = list(label = sample(c("A", "B"), 10, replace = TRUE))
)
```

cal_score_init	<i>Calculate score for each feature in each cell</i>
----------------	--

Description

Calculate score for each feature in each cell

Usage

```
cal_score_init(
  expr,
  tf = c("logtf", "tf"),
  idf = "prob",
  iae = "prob",
  par.idf = NULL,
  par.iae = NULL
)
```

Arguments

expr	a count matrix, features in row and cells in column
tf	a character, specify the TF method to use, can be "tf" or "logtf"
idf	a character, specify the IDF method to use. Available methods can be accessed using <code>idf_iae_methods()</code>
iae	a character, specify the IAE method to use. Available methods can be accessed using <code>idf_iae_methods()</code>
par.idf	other parameters for specified IDF methods
par.iae	other parameters for specified IAE methods

Value

a list of combined score, tf, idf and iae

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
label <- sample(c("A", "B"), 10, replace = TRUE)
smartid::cal_score_init(data,
  par.idf = list(label = label),
  par.iae = list(label = label)
)
```

gs_score

compute overall score based on the given marker list

Description

compute overall score based on the given marker list

Usage

```
gs_score(data, features = NULL, slot = "score", suffix = "score")

## S4 method for signature 'AnyMatrix,ANY'
gs_score(data, features = NULL)

## S4 method for signature 'AnyMatrix,list'
gs_score(data, features = NULL, suffix = "score")

## S4 method for signature 'SummarizedExperiment,ANY'
gs_score(data, features = NULL, slot = "score", suffix = "score")
```

Arguments

data	an expression object, can be matrix or SummarizedExperiment
features	vector or named list, feature names to compute score
slot	a character, specify which slot to use when data is se object, optional, default 'score'
suffix	a character, specify the name suffix to save score when features is a named list

Value

A vector of overall score for each sample

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(seq_len(10)))
gs_score(data, features = seq_len(3))
```

gs_score_init	<i>Calculate scores of each cell on given features</i>
---------------	--

Description

Calculate scores of each cell on given features

Usage

```
gs_score_init(score, features = NULL)
```

Arguments

score	matrix, features in row and samples in column
features	vector, feature names to compute score

Value

a vector of score

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
gs_score_init(data, 1:5)
```

iae *standard inverse average expression*

Description

standard inverse average expression

Usage

```
iae(expr, features = NULL, thres = 0)
```

Arguments

expr a matrix, features in row and cells in column
 features vector, feature names or indexes to compute
 thres numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IAE}_i = \log\left(1 + \frac{n}{N_{i,j} + 1}\right)$$

where n is the total number of cells, $N_{i,j}$ is the counts of feature i in cell j .

Value

a vector of inverse average expression score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::iae(data)
```

iae_hdb *inverse average expression using hdbscan cluster as label*

Description

inverse average expression using hdbscan cluster as label

Usage

```
iae_hdb(expr, features = NULL, multi = TRUE, thres = 0, minPts = 2, ...)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
multi	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
thres	numeric, cell only counts when <code>expr > threshold</code> , default 0
minPts	integer, minimum size of clusters, default 2. Details in <code>dbscan::hdbscan()</code> .
...	parameters for <code>dbscan::hdbscan()</code>

Details

Details as `iae_prob()`.

Value

a matrix of inverse average expression score

Examples

```
set.seed(123)
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::iae_hdb(data)
```

iae_igm	<i>labeled inverse average expression: IGM</i>
---------	--

Description

labeled inverse average expression: IGM

Usage

```
iae_igm(expr, features = NULL, label, lambda = 7, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
lambda	numeric, hyperparameter for IGM
thres	numeric, cell only counts when <code>expr > threshold</code> , default 0

Details

$$\mathbf{IGM}_i = \log\left(1 + \lambda \frac{\max(\text{mean}(N_{i,j \in D})_k)}{\sum_k^K (\text{mean}(N_{i,j \in D})_k * r_k) + e^{-8}}\right)$$

where λ is the hyper parameter, $N_{i,j \in D}$ is the counts of feature i in cell j within class D , and r_k is the rank of $\text{mean}(N_{i,j \in D})_k$.

Value

a vector of inverse gravity moment score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::iae_igm(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

iae_m	<i>inverse average expression: max</i>
-------	--

Description

inverse average expression: max

Usage

```
iae_m(expr, features = NULL, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IAE}_{i,j} = \log\left(1 + \frac{\max_{\{i' \in j\}}(n_{i'})}{\sum_{j=1}^n \max(0, N_{i,j} - \text{threshold}) + 1}\right)$$

where i is the feature i and i' is the feature except i , $N_{i,j}$ is the counts of feature i in cell j , and $n_{i'}$ is $\sum_{j=1}^n \text{sign}(N_{i,j} > \text{threshold})$.

Value

a matrix of inverse average expression score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid:::iae_m(data)
```

iae_prob

*labeled inverse average expression: probability based***Description**

labeled inverse average expression: probability based

Usage

```
iae_prob(expr, features = NULL, label, multi = TRUE, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
multi	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IAE}_{i,j} = \log\left(1 + \frac{\text{mean}(N_{i,j \in D})}{\max(\text{mean}(N_{i,j \in \hat{D}})) + e^{-8}} * \text{mean}(N_{i,j \in D})\right)$$

where $N_{i,j \in D}$ is the counts of feature i in cell j within class D , and \hat{D} is the class except D .

Value

a matrix of inverse average expression score

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid:::iae_prob(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

iae_rf	<i>labeled inverse average expression: relative frequency</i>
--------	---

Description

labeled inverse average expression: relative frequency

Usage

```
iae_rf(expr, features = NULL, label, multi = TRUE, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
multi	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IAE} = \log\left(1 + \frac{\text{mean}(N_{i,j \in D})}{\text{max}(\text{mean}(N_{i,j \in \hat{D}})) + e^{-8}}\right)$$

where $N_{i,j \in D}$ is the counts of feature i in cell j within class D , and \hat{D} is the class except D .

Value

a matrix of inverse average expression score

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid:::iae_rf(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

iae_sd	<i>inverse average expression using standard deviation (SD)</i>
--------	---

Description

inverse average expression using standard deviation (SD)

Usage

```
iae_sd(expr, features = NULL, log = FALSE, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
log	logical, if to do log-transformation
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IAE} = \log\left(1 + sd(tf_i) * \frac{n}{\sum_{j=1}^n \max(0, N_{i,j}) + 1}\right)$$

where tf_i is the term frequency of feature i , see details in `tf()`, n is the total number of cells and $N_{i,j}$ is the counts of feature i in cell j .

Value

a vector of inverse average expression score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid:::iae_sd(data)
```

idf	<i>standard inverse cell frequency</i>
-----	--

Description

standard inverse cell frequency

Usage

```
idf(expr, features = NULL, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IDF}_i = \log\left(1 + \frac{n}{n_i + 1}\right)$$

where n is the total number of cells, n_i is the number of cells containing feature i .

Value

a vector of inverse cell frequency score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf(data)
```

idf_hdb	<i>inverse document frequency using hdbscan cluster as label</i>
---------	--

Description

inverse document frequency using hdbscan cluster as label

Usage

```
idf_hdb(expr, features = NULL, multi = TRUE, thres = 0, minPts = 2, ...)
```

Arguments

<code>expr</code>	a matrix, features in row and cells in column
<code>features</code>	vector, feature names or indexes to compute
<code>multi</code>	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
<code>thres</code>	numeric, cell only counts when <code>expr > threshold</code> , default 0
<code>minPts</code>	integer, minimum size of clusters, default 2. Details in <code>dbscan::hdbscan()</code> .
<code>...</code>	parameters for <code>dbscan::hdbscan()</code>

Details

Details as `idf_prob()`.

Value

a matrix of inverse cell frequency score

Examples

```
set.seed(123)
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_hdb(data)
```

idf_iae_methods

Get names of available IDF and IAE methods

Description

Returns a named vector of IDF/IAE methods

Usage

```
idf_iae_methods()
```

Value

names of methods implemented

Examples

```
idf_iae_methods()
```

idf_igm	<i>labeled inverse cell frequency: IGM</i>
---------	--

Description

labeled inverse cell frequency: IGM

Usage

```
idf_igm(expr, features = NULL, label, lambda = 7, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
lambda	numeric, hyperparameter for IGM
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IGM}_i = \log\left(1 + \lambda \frac{\max(n_{i,j \in D})_k}{\sum_k ((n_{i,j \in D})_k * r_k) + e^{-8}}\right)$$

where λ is the hyper parameter, $n_{i,j \in D}$ is the number of cells containing feature i in class D , r_k is the rank of $n_{i,j \in D}$.

Value

a vector of inverse gravity moment score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_igm(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

idf_m	<i>inverse cell frequency: max</i>
-------	------------------------------------

Description

inverse cell frequency: max

Usage

```
idf_m(expr, features = NULL, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IDF}_{i,j} = \log\left(\frac{\max_{\{i' \in j\}}(n_{i'})}{n_i + 1}\right)$$

where i is the feature i and i' is the feature except i , n_i is the number of cells containing feature i , and $n_{i'}$ is the number of cells containing feature i' .

Value

a matrix of inverse cell frequency score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_m(data)
```

idf_prob	<i>labeled inverse cell frequency: probability based</i>
----------	--

Description

labeled inverse cell frequency: probability based

Usage

```
idf_prob(expr, features = NULL, label, multi = TRUE, thres = 0)
```


Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
multi	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IDF}_{i,j} = \log\left(1 + \frac{\frac{n_{i,j \in D}}{n_{j \in D}}}{\max\left(\frac{n_{i,j \in \hat{D}}}{n_{j \in \hat{D}}}\right) + e^{-8}} \frac{n_{i,j \in D}}{n_{j \in D}}\right)$$

where $n_{i,j \in D}$ is the number of cells containing feature i in class D , $n_{j \in D}$ is the total number of cells in class D , \hat{D} is the class except D .

Value

a matrix of inverse cell frequency score

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_prob(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

idf_rf	<i>labeled inverse cell frequency: relative frequency</i>
--------	---

Description

labeled inverse cell frequency: relative frequency

Usage

```
idf_rf(expr, features = NULL, label, multi = TRUE, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
label	vector, group label of each cell
multi	logical, if to compute based on binary (FALSE) or multi-class (TRUE)
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IDF}_{i,j} = \log\left(1 + \frac{\frac{n_{i,j \in D}}{n_{j \in D}}}{\max\left(\frac{n_{i,j \in \hat{D}}}{n_{j \in \hat{D}}}\right) + e^{-8}}\right)$$

where $n_{i,j \in D}$ is the number of cells containing feature i in class D , $n_{j \in D}$ is the total number of cells in class D , \hat{D} is the class except D .

Value

a matrix of inverse cell frequency score

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_rf(data, label = sample(c("A", "B"), 10, replace = TRUE))
```

idf_sd	<i>inverse cell frequency using standard deviation (SD)</i>
--------	---

Description

inverse cell frequency using standard deviation (SD)

Usage

```
idf_sd(expr, features = NULL, log = FALSE, thres = 0)
```

Arguments

expr	a matrix, features in row and cells in column
features	vector, feature names or indexes to compute
log	logical, if to do log-transformation
thres	numeric, cell only counts when expr > threshold, default 0

Details

$$\mathbf{IDF}_i = \log\left(1 + sd(tf_i) * \frac{n}{n_i + 1}\right)$$

where tf_i is the term frequency of feature i , see details in `tf()`, n is the total number of cells and n_i is the number of cells containing feature i .

Value

a vector of inverse cell frequency score for each feature

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid::idf_sd(data)
```

markers_hdbscan	<i>select markers using HDBSCAN method</i>
-----------------	--

Description

select markers using HDBSCAN method

Usage

```
markers_hdbscan(
  top_markers,
  column = ".dot",
  s_thres = NULL,
  method = c("max.one", "remove.min"),
  minPts = 5,
  plot = FALSE,
  ...
)
```

Arguments

top_markers	output of top_markers()
column	character, specify which column used as group label
s_thres	NULL or numeric, only features with score > threshold will be returned, if NULL will use 2 * average probability as threshold
method	can be "max.one" or "remove.min", if to only keep features in 1st component or return features not in the last component
minPts	integer, minimum size of clusters for dbscan::hdbscan()
plot	logical, if to plot mixture density and hist
...	other params for dbscan::hdbscan()

Value

a list of markers for each group

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
top_n <- top_markers(data, label = rep(c("A", "B"), 5))
markers_hdbscan(top_n, minPts = 2)
```

markers_mclust *select markers using mclust EM method*

Description

select markers using mclust EM method

Usage

```
markers_mclust(  
  top_markers,  
  column = ".dot",  
  prob = 0.99,  
  s_thres = NULL,  
  method = c("max.one", "remove.min"),  
  plot = FALSE,  
  ...  
)
```

Arguments

top_markers	output of top_markers()
column	character, specify which column used as group label
prob	numeric, probability cutoff for 1st component classification
s_thres	NULL or numeric, only features with score > threshold will be returned, if NULL will use 2 * average probability as threshold
method	can be "max.one" or "remove.min", if to only keep features in 1st component or return features not in the last component
plot	logical, if to plot mixture density and hist
...	other params for mclust::densityMclust()

Value

a list of markers for each group

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))  
top_n <- top_markers(data, label = rep(c("A", "B"), 5))  
markers_mclust(top_n)
```

markers_mixmdl	<i>select markers using mixtools EM method</i>
----------------	--

Description

select markers using mixtools EM method

Usage

```
markers_mixmdl(
  top_markers,
  column = ".dot",
  prob = 0.99,
  k = 3,
  ratio = 2,
  dist = c("norm", "gamma"),
  maxit = 1e+05,
  plot = FALSE,
  ...
)
```

Arguments

top_markers	output of top_markers()
column	character, specify which column used as group label
prob	numeric, probability cutoff for 1st component classification
k	integer, number of components of mixtures
ratio	numeric, ratio cutoff of 1st component mu to 2nd component mu, only when ratio > cutoff will return markers for the group
dist	can be one of "norm" and "gamma", specify if to use mixtools::normalmixEM() or mixtools::gammamixEM()
maxit	integer, maximum number of iterations for EM
plot	logical, if to plot mixture density and hist
...	other params for mixtools::normalmixEM() or mixtools::gammamixEM()

Value

a list of markers for each group

Examples

```
set.seed(1000)
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
top_n <- top_markers(data, label = rep(c("A", "B"), 5))
markers_mixmdl(top_n, k = 3)
```

ova_score_boxplot	<i>boxplot of features overall score</i>
-------------------	--

Description

boxplot of features overall score

Usage

```
ova_score_boxplot(data, features, ref.group, label, method = "t.test")
```

Arguments

data	matrix, features in row and samples in column
features	vector, feature names to plot
ref.group	character, reference group name
label	vector, group labels
method	character, statistical test to use, details in ggpubr::stat_compare_means()

Value

ggplot object

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
ova_score_boxplot(data, 1:5, ref.group = "A", label = rep(c("A", "B"), 5))
```

scale_mgm	<i>scale by mean of group mean for imbalanced data</i>
-----------	--

Description

scale by mean of group mean for imbalanced data

Usage

```
scale_mgm(expr, label, pooled.sd = FALSE)
```

Arguments

expr	matrix
label	a vector of group label
pooled.sd	logical, if to use pooled SD for scaling

Details

$$z = \frac{x - \frac{\sum_k^{n_D} (\mu_k)}{n_D}}{s}$$

where μ_k is the mean of x in k^{th} class, and n_D is the number of classes, s is the standard deviation of x , when pooled. `.sd` is set to be `TRUE`, s will be replaced with s_{pooled} , $s_{pooled} = \sqrt{\frac{\sum_k^{n_D} (n_k - 1) s_k^2}{\sum_k^{n_D} n_k - k}}$

Value

scaled matrix

Examples

```
scale_mgm(matrix(rnorm(100), 10), label = rep(letters[1:2], 5))
```

score_barplot	<i>barplot of processed score</i>
---------------	-----------------------------------

Description

barplot of processed score

Usage

```
score_barplot(top_markers, column = ".dot", f_list, n = 30)
```

Arguments

top_markers	output of <code>top_markers()</code>
column	character, specify which column used as group label
f_list	a named list of markers
n	numeric, number of returned top genes for each group

Value

ggplot object

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
top_n <- top_markers(data, label = rep(c("A", "B"), 5))
score_barplot(top_n)
```

sim_sce_test	<i>scRNA-seq test data of 4 groups simulated by splatter.</i>
--------------	---

Description

A SingleCellExperiment object containing 4 groups with each group up-regulated DEGs saved in metadata.

Usage

```
data(sim_sce_test)
```

Format

A SingleCellExperiment object of 100genes * 400 cells.

Value

SingleCellExperiment

Source

[splatter::splatSimulate\(\)](#)

sin_score_boxplot	<i>boxplot of split single feature score</i>
-------------------	--

Description

boxplot of split single feature score

Usage

```
sin_score_boxplot(data, features = NULL, ref.group, label, method = "t.test")
```

Arguments

data	matrix, features in row and samples in column
features	vector, feature names to plot
ref.group	character, reference group name
label	vector, group labels
method	character, statistical test to use, details in ggpubr::stat_compare_means()

Value

faceted ggplot object

Examples

```
data <- matrix(rnorm(100), 10, dimnames = list(1:10))
sin_score_boxplot(data, 1:2, ref.group = "A", label = rep(c("A", "B"), 5))
```

 smartid_Package

Scoring and Marker Selection method based on modified TF-IDF

Description

smartid This package enables automated selection of group specific signature, especially for rare population. The package is developed for generating specific lists of signature genes based on TF-IDF modified methods. It can also be used as a new gene-set scoring method or data transformation method. Multiple visualization functions are implemented in this package.

Value

Marker list and scores

Author(s)

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See Also

Useful links:

- <https://davislaboratory.github.io/smartid>
- Report bugs at <https://github.com/DavisLaboratory/smartid/issues>

 tf

compute term/feature frequency within each cell

Description

compute term/feature frequency within each cell

Usage

```
tf(expr, log = FALSE)
```

Arguments

expr	a count matrix, features in row and cells in column
log	logical, if to do log-transformation

Details

$$\mathbf{TF}_{i,j} = \frac{N_{i,j}}{\sum_j N_{i,j}}$$

where $N_{i,j}$ is the counts of feature i in cell j .

Value

a matrix of term/gene frequency

Examples

```
data <- matrix(rpois(100, 2), 10, dimnames = list(1:10))
smartid:::tf(data)
```

top_markers	<i>scale score and return top markers</i>
-------------	---

Description

scale and transform score and output top markers for groups

Usage

```
top_markers(
  data,
  label,
  n = 10,
  use.glm = TRUE,
  batch = NULL,
  scale = TRUE,
  use.mgm = TRUE,
  softmax = TRUE,
  slot = "score",
  ...
)

## S4 method for signature 'AnyMatrix'
top_markers(
  data,
  label,
  n = 10,
  use.glm = TRUE,
  batch = NULL,
  scale = TRUE,
  use.mgm = TRUE,
  softmax = TRUE,
```

```

    slot = "score",
    ...
  )

## S4 method for signature 'SummarizedExperiment'
top_markers(
  data,
  label,
  n = 10,
  use.glm = TRUE,
  batch = NULL,
  scale = TRUE,
  use.mgm = TRUE,
  softmax = TRUE,
  slot = "score",
  ...
)

```

Arguments

data	an expression object, can be matrix or SummarizedExperiment
label	a vector of group label
n	integer, number of returned top genes for each group
use.glm	logical, if to use <code>stats::glm()</code> to compute group mean score, if TRUE, also compute mean score difference as output
batch	a vector of batch labels, default NULL
scale	logical, if to scale data by row
use.mgm	logical, if to scale data using <code>scale_mgm()</code>
softmax	logical, if to apply softmax transformation on output
slot	a character, specify which slot to use when data is se object, optional, default 'score'
...	params for <code>top_markers_abs()</code> or <code>top_markers_glm()</code>

Value

A tibble with top n feature names, group labels and ordered scores

Examples

```

data <- matrix(rgamma(100, 2), 10, dimnames = list(1:10))
top_markers(data, label = rep(c("A", "B"), 5))

```

top_markers_abs	<i>calculate group median, MAD or mean score and order genes based on scores</i>
-----------------	--

Description

calculate group median, MAD or mean score and order genes based on scores

Usage

```
top_markers_abs(
  data,
  label,
  n = 10,
  pooled.sd = FALSE,
  method = c("median", "mad", "mean"),
  scale = TRUE,
  use.mgm = TRUE,
  softmax = TRUE,
  tau = 1
)
```

Arguments

data	matrix, features in row and samples in column
label	a vector of group label
n	integer, number of returned top genes for each group
pooled.sd	logical, if to use pooled SD for scaling
method	character, specify metric to compute, can be one of "median", "mad", "mean"
scale	logical, if to scale data by row
use.mgm	logical, if to scale data using scale_mgm()
softmax	logical, if to apply softmax transformation on output
tau	numeric, hyper parameter for softmax

Value

a tibble with feature names, group labels and ordered processed scores

Examples

```
data <- matrix(rgamma(100, 2), 10, dimnames = list(1:10))
top_markers_abs(data, label = rep(c("A", "B"), 5))
```

top_markers_glm	<i>calculate group mean score using glm and order genes based on scores difference</i>
-----------------	--

Description

calculate group mean score using glm and order genes based on scores difference

Usage

```
top_markers_glm(  
  data,  
  label,  
  n = 10,  
  family = gaussian(),  
  batch = NULL,  
  scale = TRUE,  
  use.mgm = TRUE,  
  pooled.sd = FALSE,  
  softmax = TRUE,  
  tau = 1  
)
```

Arguments

data	matrix, features in row and samples in column
label	a vector of group label
n	integer, number of returned top genes for each group
family	family for glm, details in stats::glm()
batch	a vector of batch labels, default NULL
scale	logical, if to scale data by row
use.mgm	logical, if to scale data using scale_mgm()
pooled.sd	logical, if to use pooled SD for scaling
softmax	logical, if to apply softmax transformation on output
tau	numeric, hyper parameter for softmax

Value

a tibble with feature names, group labels and ordered processed scores

Examples

```
data <- matrix(rgamma(100, 2), 10, dimnames = list(1:10))  
top_markers_glm(data, label = rep(c("A", "B"), 5))
```

top_markers_init	<i>compute group summarized score and order genes based on processed scores</i>
------------------	---

Description

compute group summarized score and order genes based on processed scores

Usage

```
top_markers_init(
  data,
  label,
  n = 10,
  use.glm = TRUE,
  batch = NULL,
  scale = TRUE,
  use.mgm = TRUE,
  softmax = TRUE,
  ...
)
```

Arguments

data	matrix, features in row and samples in column
label	a vector of group label
n	integer, number of returned top genes for each group
use.glm	logical, if to use <code>stats::glm()</code> to compute group mean score, if TRUE, also compute mean score difference as output
batch	a vector of batch labels, default NULL
scale	logical, if to scale data by row
use.mgm	logical, if to scale data using <code>scale_mgm()</code>
softmax	logical, if to apply softmax transformation on output
...	params for <code>top_markers_abs()</code> or <code>top_markers_glm()</code>

Value

a tibble with feature names, group labels and ordered processed scores

Examples

```
data <- matrix(rgamma(100, 2), 10, dimnames = list(1:10))
top_markers_init(data, label = rep(c("A", "B"), 5))
```

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