# Package 'scFeatureFilter'

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Type Package

**Title** A correlation-based method for quality filtering of single-cell RNAseq data

Version 1.24.0

**Description** An R implementation of the correlation-based method developed in the Joshi laboratory to analyse and filter processed single-cell RNAseq data. It returns a filtered version of the data containing only genes expression values unaffected by systematic noise.

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LazyData TRUE

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bin_	scdata Bin genes by mean expression.

# Description

Divides the genes that were not included in the top window in windows of the same size with decreasing mean expression levels.

# Usage

```
bin_scdata(dataset, window_number = NULL, window_size = NULL,
    verbose = TRUE)
```

# Arguments

dataset	A list, containing the top window generated by extract_top_genes as the first element, and the rest of undivided genes as the second. Usually the output of define_top_genes
window_number	An integer, indicating the number of bins to be used.
window_size	An integer, indicating the number of genes to be included in each window. Ignored if window_size is defined.
verbose	A boolean. Should the function print a message about window size or the number of windows created?

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#### **Details**

Two binning methods are available:

- window\_number: Divides the genes into the number of windows specified.
- window\_size: Divides the genes into windows of the size specified.

This function adds a bin number column to the data frame.

This function is designed to take the list output by the extract\_top\_window function as an argument, operating only on the second element of it. Once the genes in it have been binned, both elements of the list are bound together in a data frame and returned. The output contains a new column bin, which indicates the window number assigned to each gene.

#### Value

A data frame containing the binned genes.

# **Examples**

```
library(magrittr)
expMat <- matrix(
    c(1, 1, 1,
        1, 2, 3,
        0, 1, 2,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)

calculate_cvs(expMat) %>%
    define_top_genes(window_size = 1) %>%
    bin_scdata(window_number = 2)

calculate_cvs(expMat) %>%
    define_top_genes(window_size = 1) %>%
    bin_scdata(window_size = 1)
```

calculate\_cvs

Compute mean expression level, standard deviation and coefficient of variation of each feature.

#### **Description**

Compute mean expression level, standard deviation and coefficient of variation (CV) of each feature (i.e. gene or transcript) in the supplied data. Filter features with high proportion of 0 expression.

#### Usage

```
calculate_cvs(data, max_zeros = 0.75, sce_assay = NULL)
```

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#### **Arguments**

data

sce\_assay

	values for the cells as columns.
max_zeros	A number between 0 and 1 indicating the maximum proportion of zero expression values allowed per row. Features with a higher proportion of 0 will be discarded.

if data is an SingleCellExperiment object, sce\_assay should be one of names(assays(<SingleCellE

A data frame, a matrix or a SingleCellExperiment object. If data frame or matrix, it should contain expression values for each gene as rows, and expression

#### **Details**

Before CV computation, the function removes all rows that have a proportion of zeros above the specified threshold. Genes with many 0s are poorly informative, and would bias the later correlations. Removing them also prevents division by zero when calculating CVs.

The data provided must cell/sample names as column names. Feature name can be given either in the first column or as row names.

In the output, mean, standard deviation and CV are incorporated as new columns in the data frame, named mean, sd and cv.

#### Value

A data frame, containing the filtered data with additional columns: mean, standard deviation and cv values for each row.

#### **Examples**

```
expMat <- matrix(
    c(1, 1, 1,
        1, 2, 3,
        0, 1, 2,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)
calculate_cvs(expMat)
calculate_cvs(expMat, max_zeros = 0.5)</pre>
```

correlate\_windows

Calculate correlations against top window.

#### **Description**

Calculates pairwise correlations between all features each window against all features in the reference window.

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#### Usage

```
correlate_windows(dataset, n_random = 3, ...)
```

#### **Arguments**

A data frame containing all the binned genes. Usually the output of bin\_scdata.

n\_random

Number of top window randomization to serve as a negative control. Default to 3.

Additional arguments to be passed to cor. Default method is pearson which is the fastest.

#### **Details**

This function:

- correlates each feature in each window to each feature in the top window.
- randomize the top window by shuffling expression value, and correlate each gene in each window to the randomized top window. This negative control is repeated as many time as specified by the n\_random parameter.

The input of this function is usually the output of the bin\_scdata function.

#### Value

A tibble containing correlation values.

```
library(magrittr)
expMat <- matrix(
    c(1, 1, 5,
        1, 2, 3,
        0, 1, 4,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)

calculate_cvs(expMat) %>%
    define_top_genes(window_size = 2) %>%
    bin_scdata(window_number = 1) %>%
    correlate_windows
```

```
correlations_to_densities
```

Transform the correlation table to density distributions of correlation values

# **Description**

Takes the output of correlate\_windows and computes density curves of correlation coefficient for each window comparison.

# Usage

```
correlations_to_densities(df, n = 64, absolute_cc = TRUE)
```

# **Arguments**

A data frame, usually the output of correlate\_windows.

Resolution of the correlation density curve. Default to 64.

absolute\_cc Should the function use the absolute value of correlation coefficients? Default

to TRUE to simplify plots and avoid annoying, non-symmetrical, near 0, shifts of

distributions.

#### Value

A tibble with columns bin, window, cor\_coef and density.

```
library(magrittr)
expMat <- matrix(
    c(1, 1, 5,
        1, 2, 3,
        0, 1, 4,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)

calculate_cvs(expMat) %>%
    define_top_genes(window_size = 2) %>%
    bin_scdata(window_number = 1) %>%
    correlate_windows %>%
    correlations_to_densities
```

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del me_cop_genes Define the reference withdow using the most highly expressed features.	define_top_genes	Define the reference window using the most highly expressed features.
---	------------------	---

#### **Description**

Define the group of features in the dataset that will be considered as reference, the top window, by specifying either a number of features or an expression threshold.

# Usage

```
define_top_genes(dataset, window_size = NULL, mean_expression = NULL,
    min_expression = NULL)
```

#### **Arguments**

dataset A data frame, containing features as rows and cells as columns, and where the

mean expression value for each gene has been added as a column. Usually the

output of calculate\_cvs.

window\_size Number of features in the defined top window. Recommended to 100 features.

mean\_expression

A number. Genes with a mean expression across cells higher than the value will

be selected. Ignored if window\_size is defined.

min\_expression A number. Genes with a minimum expression across all cells higher than the

value will be selected. Ignored if window\_size or mean\_expression is defined.

#### **Details**

There are three selection methods available:

- window\_size: features are ranked by mean expression across cells, and the top slice of the specified size is selected.
- mean\_expression: the mean column is checked, and all features with mean expression above the threshold indicated are selected.
- min\_expression: features where all expression values are above the expression threshold indicated are selected.

In general, it is advisable to avoid generating top windows larger than 250 features (100 features is the recommended value), to prevent excessively long computation time as well as to preserve the quality of the analysis, as the top window should only include a subset of reliable values.

#### Value

A list with two elements, both data frames: the defined top window, and the rest of the genes.

determine\_bin\_cutoff

#### **Examples**

```
library(magrittr)
expMat <- matrix(
    c(1, 1, 1,
        1, 2, 3,
        0, 1, 2,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)

calculate_cvs(expMat) %>%
    define_top_genes(window_size = 2)

calculate_cvs(expMat) %>%
    define_top_genes(mean_expression = 1.5)
```

#### **Description**

Takes the output of get\_mean\_median and decide until which window to keep based on background level and a threshold.

#### Usage

```
determine_bin_cutoff(metric_table, threshold = 2,
  selected_metric = c("mean", "median", "score"),
  random_function_summarisation = mean)
```

# **Arguments**

metric\_table A data frame, usually the output of get\_mean\_median
threshold How many time higher than the background should the last bin be? Default to 2.
selected\_metric
Which metric to use (i.e. which column from metric\_table to work with). Default to mean.
random\_function\_summarisation

A function used to aggregate the randomised control across bin. Default to mean.

#### **Details**

Background level is estimated by averaging correlation coefficient obtained from the top window randomisations.

Bins (or windows) of features are kept until the mean (or median) correlation coefficient falls under a threshold value threshold x background level.

filter\_expression\_table

# Value

A number, the first bin of features to discard.

#### See Also

```
get_mean_median, plot_metric
```

#### **Examples**

```
myData <- tibble::tibble(
    bin = rep(c(1, 2, 3), each = 3),
    window = rep(c("top_window", "shuffled_top_window_1", "shuffled_top_window_2"), 3),
    mean = c(0.8, 0.1, 0.11, 0.14, 0.12, 0.09, 0.10, 0.13, 0.08)
)
determine_bin_cutoff(myData)</pre>
```

```
filter_expression_table
```

Filter binned expression matrix

#### **Description**

Takes a binned expression table (the output of bin\_scdata), a bin number (usually the output of determine\_bin\_cutoff) and returned a filtered expression table or matrix.

#### **Usage**

```
filter_expression_table(bined_table, bin_cutoff, as_matrix = FALSE)
```

#### **Arguments**

```
bined_table A tibble, usually the output of bin_scdata.

bin_cutoff the number of the first bin to be filtered out. Can be the output of determine_bin_cutoff).

A boolean. Should the return be a tibble (FALSE, the default) or a matrix (TRUE).
```

#### Value

A tibble or a matrix depending on the value of as\_matrix

#### See Also

```
bin_scdata, determine_bin_cutoff
```

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#### **Examples**

```
myData <- tibble::data_frame(
    bin = rep(c(1, 2, 3), each = 3),
    mean = 9:1,
    sd = runif(9),
    cv = runif(9),
    cell1 = 8:0 + runif(9),
    cell2 = 8:0 + runif(9)
)
filter_expression_table(myData, bin_cutoff = 2)
filter_expression_table(myData, bin_cutoff = 3)</pre>
```

get\_mean\_median

Extract mean and median correlation coefficient values

# **Description**

Takes the output of correlate\_windows and extract the mean and the median correlation value for each window comparison.

#### Usage

```
get_mean_median(df, absolute_cc = TRUE)
```

#### **Arguments**

df

A data frame, usually the output of correlate\_windows.

absolute\_cc

Should the function work of absolute value of correlation coefficients? Default to TRUE to simplify plots and avoid annoying, non-symmetrical, near 0, shifts of distributions.

#### Value

A data\_frame with columns bin, window, mean and median.

```
library(magrittr)
expMat <- matrix(
    c(1, 1, 5,
        1, 2, 3,
        0, 1, 4,
        0, 0, 2),
    ncol = 3, byrow = TRUE, dimnames = list(paste("gene", 1:4), paste("cell", 1:3))
)
calculate_cvs(expMat) %>%
    define_top_genes(window_size = 2) %>%
```

```
bin_scdata(window_number = 1) %>%
correlate_windows(n_random = 2) %>%
get_mean_median
```

```
plot_correlations_distributions
```

Produce a density plot of correlation values for each window of feature

# **Description**

Feature by feature correlation values between every windows and the reference to window of features are visualized as density lines, one facet per comparison. Two density lines are drown in each facets:

- A thin colored line, the correlations between the bin and the reference top bin of features
- A thicker blue line with grey error area, the correlations between the bin and the **randomized** top bin of features. The lines are not shown if n\_random = 0 in correlate\_windows.

# Usage

```
plot_correlations_distributions(df, metrics = NULL, vlines = c("mean",
    "median"), facet_ncol = 4)
```

# **Arguments**

df	A tibble, usually the output of correlations_to_densities.
metrics	Optional. The output of get_mean_median. Dashed line will represent mean or median of the correlation coefficient distributions.
vlines	A string, either "mean" or "median". Should the dashed line represent the mean or the median of the correlation coefficient distributions? Ignored if metrics is NULL.
facet_ncol	The number of columns to arrange the plots.

#### Value

A ggplot2 plot.

#### See Also

```
correlations_to_densities, get_mean_median
```

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# **Examples**

```
library(magrittr)
myData <- scData_hESC %>%
calculate_cvs %>%
    define_top_genes(window_size = 100) %>%
    bin_scdata(window_size = 1000)

corDistrib <- correlate_windows(myData, n_random = 3)

corDens <- correlations_to_densities(corDistrib)

plot_correlations_distributions(corDens)

metrics <- get_mean_median(corDistrib)

plot_correlations_distributions(corDens, metrics = metrics)</pre>
```

plot\_mean\_variance

*Produce a mean expression x coefficient of variation scatter plot.* 

# **Description**

Use the output of calculate\_cvs or bin\_scdata and plot a feature mean expression x coefficient of variation scatter plot. Mean expression is represented as log10(mean + 1). Each dot represents a feature. Means and coefficient of variations were obtained across single cells. Optionally, colours each dot according to the defined bins of features. Optionally, adds a density2d geom.

#### Usage

```
plot_mean_variance(df, density = TRUE, colourByBin = TRUE,
  density_color = "blue", ...)
```

# **Arguments**

df A tibble, usually the output of calculate\_cvs or bin\_scdata.

density A boolean. Should a density2d geom be added to the plot?

colourByBin A boolean. Should feature be coloured by bin? Need a bin column in df (i.e. the output of bin\_scdata).

density\_color Colour of the density2d curves.

Further arguments are passed to geom\_point such as size.

Value

A ggplot2 plot.

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#### See Also

```
calculate_cvs, bin_scdata
```

#### **Examples**

```
library(magrittr)
scData_hESC %>%
    calculate_cvs %>%
    plot_mean_variance(colourByBin = FALSE)

scData_hESC %>%
    calculate_cvs %>%
    define_top_genes(window_size = 100) %>%
    bin_scdata(window_size = 1000) %>%
    plot_mean_variance
```

plot\_metric

Produce a bar chart of mean (or median) correlation coefficient per bin of feature.

# **Description**

Use the output of get\_mean\_median and produce a bar chart of mean (or median) correlation coefficient per bin of features. Correlations against the randomised top window are shown as dot-and-whiskers, and are used to estimate a background level.

#### Usage

```
plot_metric(metric_table, selected_metric = c("mean", "median", "score"),
    show_ctrl = TRUE, control_color = "blue", show_threshold = TRUE,
    threshold = 2, threshold_color = "red", line_size = 1,
    annotate_lines = TRUE)
```

#### **Arguments**

```
metric_table
                  A tibble, usually the output of get_mean_median.
selected_metric
                  Which column in metricsTable to use? Default to mean.
show_ctrl
                  A boolean. Should a dashed line indicate the estimated background level?
control_color
                 The colour of the background dashed line (default to blue).
show_threshold A boolean. Should a dashed line indicate the estimated threshold level?
threshold
                  How many times the background level should be multiplies do determine a
                  threshold? Default to 2. The higher the more stringent.
threshold_color
                  The colour of the threshold dashed line (default to blue).
line size
                  Thickness of the dashed lines.
annotate_lines A boolean. Should the dashed lines be annotated?
```

# Value

A ggplot2 plot.

#### See Also

```
get_mean_median
```

#### **Examples**

```
library(magrittr)
scData_hESC %>%
    calculate_cvs %>%
    define_top_genes(window_size = 100) %>%
    bin_scdata(window_size = 1000) %>%
    correlate_windows(n_random = 3) %>%
    get_mean_median %>%
    plot_metric
```

plot\_top\_window\_autocor

Utility plot to choose a top\_window size

# **Description**

Plot mean autocorrelation value of the features of the top window depending on increasing top window size.

#### Usage

```
plot_top_window_autocor(sc_data, from = 10, to = 400, by = 2, ...)
```

# **Arguments**

 $sc\_data \hspace{1cm} A \ tibble, \ usually \ the \ output \ of \ link\{calculate\_cvs\}.$ 

from Minimum size of the top window. to Maximum size of the top window.

by Size of the steps to walk form from to to. See seq.

... Arguments to be passed to cor, for example method = "spearman"

#### Value

A ggplot2 plot.

```
plot_top_window_autocor(calculate_cvs(scData_hESC))
```

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scData_hESC	Expression data from 32 human embryonic stem cells	

#### **Description**

Expression of 60,468 Gencode gene (in FPKM) from 32 single cell RNAseq of human embryonic stem cells

# Usage

```
scData_hESC
```

#### **Format**

```
A tibble with 60,468 rows (genes) and 33 columns (cells): 
tracking_id The Gencode human encode gene id
next 33 columns Single embryonic stem cells
```

#### Value

A tibble.

#### Source

The example dataset was processed by Mantsoki et al. (2016) https://www.ncbi.nlm.nih.gov/pubmed/26951854, data from human Embryonic Stem cells was generated by Yan et al. (2013)https://www.ncbi.nlm.nih.gov/pubmed/23934149.

```
sc_feature_filter Filter scRNA-seq expression matrix to keep only highly informative features. Integrated pipeline.
```

# Description

This pipeline function takes an expression matrix as an input and select the features (genes, transcripts) with an estimated technical noise level lower that biological variation in the data. This is achieved by binning the data and calculating the correlation for each bin with highly expressed (lowest noise) gene set (see the vignette for details on the method).

#### Usage

```
sc_feature_filter(sc_data, print_plots = FALSE, max_zeros = 0.75,
  threshold = 2, top_window_size = 100, other_window_size = 1000,
  n_random = 3, sce_assay = NULL)
```

sc\_feature\_filter

#### **Arguments**

sc\_data A data frame, a matrix or a SingleCellExperiment object. If data frame or matrix, it should contain expression values for each gene as rows, and expression values for the cells as columns. print\_plots A boolean. Should the function produce three plots as a side effect? Plots are the output of  $plot_mean_variance$ ,  $plot_correlations_distributions$  and plot\_metric. A number between 0 and 1. Maximum proportion of cells with 0 expression for max\_zeros a feature to be kept. threshold A number higher than 1. The higher the more stringent the feature selection will be. See determine\_bin\_cutoff. top\_window\_size Size of the reference bin. See define\_top\_genes other\_window\_size Size of the other bins of feature. See bin\_scdata Number of control windows generated by shuffling the top bin of features. n\_random if sc\_data is an SingleCellExperiment object, sce\_assay should be one of sce\_assay names(assays(<SingleCellExperiment>)).

#### **Details**

The function can optionally produce three plots of print\_plots is TRUE. It is recommended to open a graphical device (i.e. through pdf or png), to call scFeatureFilter, and then to close the device with dev.off.

#### Value

A matrix or a tibble, depending on the type of sc\_data, containing only the top expressed features.

```
sc_feature_filter(scData_hESC)

# with plots
## Not run:
pdf("diagnostic.pdf")
sc_feature_filter(sc_data, print_plots = TRUE)
dev.off()

## End(Not run)
```

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