# Package 'scUtils'

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Title Utility Functions for Single-Cell RNA Sequencing Data
Version 0.1.0
Description Analysis of single-cell RNA sequencing data can be simple and clear with the right utility functions. This package collects such functions, aiming to fulfill the following criteria: code clarity over performance (i.e. plain R code instead of C code), most important analysis steps over completeness (analysis 'by hand', not automated integration etc.), emphasis on quantitative visualization (intensity-coded color scale, etc.).
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closed\_breaks\_log2

closed\_breaks\_log2

Closed breaks for log scale

#### **Description**

Finds breaks that are powers of 2, and forces inclusion of upper and lower limits (displaying the closed interval). Including limits specifically is particularly useful for ggplot2's color/fill, as it emphasizes the meaning of maximal/minimal color intensities (see examples).

## Usage

```
closed_breaks_log2(lims)
```

#### **Arguments**

lims

Vector with lower and upper limits (in that order) of the data that you want breaks for.

#### **Details**

The feat function uses closed\_breaks\_log2 to color by gene expression, where the maximal expression gives valuable intuition for a gene's overall expression strength. For x- or y-axis (scale\_\*\_log10), I still recommend breaks\_log from the scales package.

#### Value

Numeric vector with breaks.

#### See Also

```
closed_labels
```

#### **Examples**

```
# closed breaks include maximum, breaks_log do not:
closed_breaks_log2(lims = c(.01, 977.1))
scales::breaks_log()(c(.01, 977.1))
```

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closed_labels	Human-readable labels for closed breaks
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## Description

Complements the closed\_breaks\_log2 function.

## Usage

```
closed_labels(x, min_is_zero = FALSE)
```

## Arguments

Х	Vector of breaks for which to produce labels. Typically, this is the output of closed_breaks_log2.
min_is_zero	Should the smallest break be displayed as zero (TRUE) or as the actual value (FALSE). Default: FALSE

#### **Details**

This is a helper for the feat function. feat replaces numeric zeros with the next-smallest expression value to avoid taking the logarithm of zero. min\_is\_zero can be used to display the lowest break of the color scale as zero in these cases.

#### Value

Character vector with labels, used by feat function.

#### See Also

```
label_scientific label_number_auto
```

## **Examples**

```
# human readable output:
closed_labels(c(.001111,.122, 0.5, 10, 100, 1800))
```

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colVars\_spm

Variance computation for sparse matrices

#### **Description**

Compute variance for each column / each row of a dgCMatrix (from Matrix package).

#### Usage

```
colVars_spm(spm)
rowVars_spm(spm)
```

#### **Arguments**

spm

A sparse matrix of class dgCMatrix from the Matrix package.

#### **Details**

The only supported format currently is dgCMatrix. While the Matrix package has other formats, this one is used for scRNAseq raw count data. Function code written by Simon Anders.

#### Value

Vector with variances.

#### See Also

```
vignette("Intro2Matrix", package="Matrix") CsparseMatrix-class
```

## **Examples**

```
library(Matrix)
mat <- as(matrix(rpois(900,1), ncol=3), "dgCMatrix")
colVars_spm(mat)</pre>
```

feat

Feature Plot

#### **Description**

Highlight gene expression data in a 2D-embedding (UMAP, tSNE, etc.).

#### Usage

```
feat(embedding, expression, legend_name = "Expression")
```

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#### **Arguments**

A matrix/data.frame/tibble/... with exactly two columns. If colnames are missing, the axis will be named "Dim1" and "Dim2". Other classes than matrix/data.frame/tibble are possible, as long as data.frame(embedding)) produces a numeric data.frame.

Numeric vector with expression values of the gene of interest. Order has to correspond to the row order in embedding. Typically, expression is normalized gene expression and we recommend k/s/mean(1/s), where k are UMI counts for the gene of interest and s are totalUMI of the cell (aka library size).

legend\_name Text displayed above the legend. Most commonly the name of the displayed

gene.

#### **Details**

This function discourages customization on purpose, because it bundles geoms, themes and settings that I found important for visualizing gene expression in scRNAseq data:

- · coord\_fixed, to avoid distortion of embeddings
- geom\_point with size=.4, to ameliorate overplotting
- No background grid, because distances and axis units in embeddings do not carry meaning for most dimensionality reduction techniques.
- Intensity-coded color scales (viridis) displayed with log2-transformation. Makes visualization independent of colorblindness and appropriate for gene expression data (which is usually Log Normal distributed).
- Color scale breaks are displayed as 'closed interval', i.e. max(expression) and min(expression) are the most extreme breaks. Rounding makes them human-readable. This functionality is provided by closed\_breaks\_log2 and closed\_labels.

If you insist on customizing, think of this function as a great starting point, you can simply copypaste the code after typing feat into your console.

#### Value

A ggplot2 object storing a colored scatter plot.

#### See Also

```
ggplot, closed_labels, closed_breaks_log2
```

#### **Examples**

```
# expression goes from 0 to 22:
set.seed(100)
feat(matrix(rnorm(2000, c(.1, 3)), ncol=2), rpois(1000, c(.1, 11)))
# expression goes from 2 to 52:
set.seed(100)
feat(matrix(rnorm(2000, c(.1, 3)), ncol=2), rpois(1000, c(10, 31)))
```

is\_wholenumber

is_wholenumber	Check if number(s) is/are integers. In contrast to is.integer, is_wholenumber does not check the class but accepts all numbers that are integers with reasonable precision.
	are integers with reasonable precision.

## Description

Check if number(s) is/are integers. In contrast to is.integer, is\_wholenumber does not check the class but accepts all numbers that are integers with reasonable precision.

## Usage

```
is_wholenumber(x, tol = .Machine$double.eps^0.5)
```

## Arguments

x Number to test

tol tolerance for testing

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