

# Package: SCent (via r-universe)

June 25, 2024

**Title** Single Cell Entropy Analysis of Gene Heterogeneity in Cell Populations

**Version** 0.1.0

**Description** Analyse single cell RNA sequencing data using entropy to calculate heterogeneity and homogeneity of genes amongst the cell population. From the work of Michael J. Casey, Ruben J. Sanchez-Garcia and Ben D. MacArthur (2020) [doi:10.1101/2020.10.01.322255](https://doi.org/10.1101/2020.10.01.322255).

**License** GPL (>= 3)

**Encoding** UTF-8

**Roxygen** list(markdown = TRUE)

**RoxygenNote** 7.1.2

**Imports** entropy, tibble

**Suggests** rmarkdown, knitr, testthat (>= 3.0.0)

**VignetteBuilder** knitr

**Config/testthat/edition** 3

**Repository** <https://hwarden162.r-universe.dev>

**RemoteUrl** <https://github.com/hwarden162/scent>

**RemoteRef** HEAD

**RemoteSha** f99eb8ea89a9c652b59857b56b74b9442e6e232e

## Contents

gene_counts . . . . .	2
gene_het . . . . .	2
gene_hom . . . . .	3
normalise . . . . .	4
rm_low_counts . . . . .	5
rm_low_counts_tidy . . . . .	6
scent_select . . . . .	7
scent_select_tidy . . . . .	8

<b>Index</b>	<b>10</b>
--------------	-----------

---

`gene_counts`*Find the Number of Times Each Gene Has Been Expressed*

---

**Description**

Find the Number of Times Each Gene Has Been Expressed

**Usage**

```
gene_counts(expr, transpose = FALSE)
```

**Arguments**

<code>expr</code>	A matrix of gene expressions with cells as rows and genes as columns
<code>transpose</code>	A logical value indicating whether the matrix should be transposed before operations are carried out

**Value**

A vector of counts of expression for each gene

**Examples**

```
# Creating Data
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
genes <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(genes) <- paste0("cell", 1:7)
colnames(genes) <- paste0("gene", 1:5)

#Calculating Gene Counts
gene_counts(genes)
```

---

`gene_het`*Find the Heterogeneity of a Gene Within a Population*

---

**Description**

Find the Heterogeneity of a Gene Within a Population

**Usage**

```
gene_het(expr, unit = "log2", normalise = TRUE, transpose = FALSE)
```

**Arguments**

expr	A vector or matrix of gene expressions. For the matrix, genes should be represented as rows and cells as columns.
unit	The units to be parsed to the entropy function.
normalise	A logical value representing whether the gene frequencies should be normalised into a distribution.
transpose	A logical value representing whether the matrix should be transposed before any calculations are performed.

**Value**

A vector of the information gained from the gene distribution compared to the uniform distribution. The higher the value more heterogeneous the cell is within the population.

**Examples**

```
# Creating Data
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)

# Calculating Heterogeneity For Each Gene
gene_het(gene1)
gene_het(gene2)
gene_het(gene3)
gene_het(gene4)
gene_het(gene5)

# Calculating Heterogeneity For a Matrix
gene_het(gene_counts)
```

---

gene\_hom

*Find the Homogeneity of a Gene Within a Population*


---

**Description**

Find the Homogeneity of a Gene Within a Population

**Usage**

```
gene_hom(expr, unit = "log2", normalise = TRUE, transpose = FALSE)
```

**Arguments**

expr	A vector or matrix of gene expressions. For the matrix, genes should be represented as rows and cells as columns.
unit	The units to be parsed to the entropy function.
normalise	A logical value representing whether the gene frequencies should be normalised into a distribution.
transpose	A logical value representing whether the matrix should be transposed before any calculations are performed.

**Value**

A vector of the information contained in the distribution of each gene. The higher this is, the more homogeneous the gene is within the cell population.

**Examples**

```
# Creating Data
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)

# Calculating Homogeneity For Each Gene
gene_hom(gene1)
gene_hom(gene2)
gene_hom(gene3)
gene_hom(gene4)
gene_hom(gene5)

# Calculating Homogeneity For a Matrix
gene_hom(gene_counts)
```

---

normalise

*Normalise Counts into a Distribution*


---

**Description**

A function that takes frequency count data and normalises it into a probability distribution. Only available internally within SCEnt.

**Usage**

```
normalise(dist)
```

**Arguments**

dist                    A vector of a frequency distribution.

**Value**

A vector of a probability distribution relative to the frequencies.

---

rm_low_counts	<i>Remove Lowly Expressed Genes From Expression Data</i>
---------------	--

---

**Description**

Remove Lowly Expressed Genes From Expression Data

**Usage**

```
rm_low_counts(
  expr,
  count_threshold = NULL,
  perc_threshold = NULL,
  transpose = FALSE
)
```

**Arguments**

expr                    A matrix of gene expression with cells as rows and genes as columns

count\_threshold        A threshold for the number of counts a gene must have to be included. Only one threshold may be used at a time.

perc\_threshold        A threshold for what percentile the gene counts should be cut off at. Only one threshold may be used at a time.

transpose              A logical value indicating whether the expression matrix should be transposed before any operations are carried out.

**Value**

A matrix of gene expressions with the low count genes, as specified by the user, removed.

**Examples**

```
# Creating Data
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
```

```
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)

# Removing Low Count Genes
rm_low_counts(gene_counts, count_threshold = 7)
rm_low_counts(gene_counts, perc_threshold = 0.1)
```

---

rm\_low\_counts\_tidy      *Tidy Wrapper To Remove Lowly Expressed Genes From Expression Data*

---

## Description

Tidy Wrapper To Remove Lowly Expressed Genes From Expression Data

## Usage

```
rm_low_counts_tidy(
  expr,
  count_threshold = NULL,
  perc_threshold = NULL,
  transpose = FALSE
)
```

## Arguments

**expr**                    A tibble of gene expression with cells as rows and genes as columns

**count\_threshold**        A threshold for the number of counts a gene must have to be included. Only one threshold may be used at a time.

**perc\_threshold**        A threshold for what percentile the gene counts should be cut off at. Only one threshold may be used at a time.

**transpose**             A logical value indicating whether the expression matrix should be transposed before any operations are carried out.

## Value

A tibble of gene expressions with the low count genes, as specified by the user, removed.

## Examples

```
# Creating Data
library(tibble)
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
```

```

gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)
gene_counts <- as_tibble(gene_counts)

# Removing Low Count Genes
rm_low_counts_tidy(gene_counts, count_threshold = 7)
rm_low_counts_tidy(gene_counts, perc_threshold = 0.1)

```

---

scent\_select

*Feature Selection by Gene Heterogeneity*


---

## Description

Feature Selection by Gene Heterogeneity

## Usage

```

scent_select(
  expr,
  bit_threshold = NULL,
  count_threshold = NULL,
  perc_threshold = NULL,
  unit = "log2",
  normalise = TRUE,
  transpose = FALSE
)

```

## Arguments

expr	A matrix of gene expression data. Cells should be represented as rows and genes should be represented as columns.
bit_threshold	The threshold for the amount of bits of information a gene must add to be selected as a feature. Only one threshold can be used at a time.
count_threshold	A number represented how many of the most heterogeneous cells should be selected. Only one threshold can be used at a time.
perc_threshold	The percentile of the heterogeneity distribution above which a gene should be to be selected as a feature.
unit	The units to be used when calculating entropy.
normalise	A logical value representing whether the gene counts should be normalised into a probability distribution.
transpose	A logical value representing whether the matrix should be transposed before having any operations computed on it.

**Value**

A matrix of gene expression values where genes with low heterogeneity have been removed.

**Examples**

```
# Creating Data
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)

# Performing Feature Selection
scent_select(gene_counts, bit_threshold = 0.85)
scent_select(gene_counts, count_threshold = 2)
scent_select(gene_counts, perc_threshold = 0.25)
```

---

scent\_select\_tidy      *A Tidy Wrapper for Feature Selection by Heterogeneity*

---

**Description**

A Tidy Wrapper for Feature Selection by Heterogeneity

**Usage**

```
scent_select_tidy(
  expr,
  bit_threshold = NULL,
  count_threshold = NULL,
  perc_threshold = NULL,
  unit = "log2",
  normalise = TRUE,
  transpose = FALSE
)
```

**Arguments**

expr	A tibble of gene expression data. Cells should be represented as rows and genes should be represented as columns.
bit_threshold	The threshold for the amount of bits of information a gene must add to be selected as a feature. Only one threshold can be used at a time.
count_threshold	A number represented how many of the most heterogeneous cells should be selected. Only one threshold can be used at a time.



perc_threshold	The percentile of the heterogeneity distribution above which a gene should be to be selected as a feature.
unit	The units to be used when calculating entropy.
normalise	A logical value representing whether the gene counts should be normalised into a probability distribution.
transpose	A logical value representing whether the matrix should be transposed before having any operations computed on it.

### Value

A tibble of gene expression values where genes with low heterogeneity have been removed.

### Examples

```
# Creating Data
library(tibble)
gene1 <- c(0, 0, 0, 0, 1, 2, 3)
gene2 <- c(5, 5, 3, 2, 0, 0, 0)
gene3 <- c(2, 0, 2, 1, 3, 0, 1)
gene4 <- c(3, 3, 3, 3, 3, 3, 3)
gene5 <- c(0, 0, 0, 0, 5, 0, 0)
gene_counts <- matrix(c(gene1, gene2, gene3, gene4, gene5), ncol = 5)
rownames(gene_counts) <- paste0("cell", 1:7)
colnames(gene_counts) <- paste0("gene", 1:5)
gene_counts <- as_tibble(gene_counts)

# Performing Feature Selection
scent_select_tidy(gene_counts, bit_threshold = 0.85)
scent_select_tidy(gene_counts, count_threshold = 2)
scent_select_tidy(gene_counts, perc_threshold = 0.25)
```

# Index

gene\_counts, [2](#)

gene\_het, [2](#)

gene\_hom, [3](#)

normalise, [4](#)

rm\_low\_counts, [5](#)

rm\_low\_counts\_tidy, [6](#)

scent\_select, [7](#)

scent\_select\_tidy, [8](#)