# Package 'rifiComparative'

May 25, 2024

**Title** 'rifiComparative' compares the output of rifi from two different conditions.

Version 1.5.0

**Description** 'rifiComparative' is a continuation of rifi package. It compares two conditions output of rifi using half-life and mRNA at time 0 segments. As an input for the segmentation, the difference between half-life of both condtions and log2FC of the mRNA at time 0 are used. The package provides segmentation, statistics, summary table, fragments visualization and some additional useful plots for further analysis.

**Depends** R (>= 4.2)

Imports cowplot, doMC, parallel, dplyr, egg, foreach, ggplot2, ggrepel, graphics, grDevices, grid, methods, nnet, rlang, S4Vectors, scales, stats, stringr, tibble, rtracklayer, utils, writexl, DTA, LSD, reshape2, devtools, SummarizedExperiment

Suggests DescTools, knitr, rmarkdown, BiocStyle

VignetteBuilder knitr

**biocViews** RNASeq, DifferentialExpression, GeneRegulation, Transcriptomics, Microarray, Software

BugReports https://github.com/CyanolabFreiburg/rifiComparative

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2 adjusting\_HLToInt

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# Description

'adjusting\_HLToInt' merges HL and intensity segments adapting the positions to each other and combining to the genome annotation. To make HL and intensity segments comparable, log2FC(HL) is used to generate the data frame instead of distance. The fragments should have a significant p\_value from t-test at least from one segmentation, either HL or intensity.

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## Usage

```
adjusting_HLToInt(data, Strand = c("+", "-"), annotation)
```

#### **Arguments**

data data frame: data frame combined data by column

Strand string: either "+" or "-"

annotation data frame: data frame from processed gff3 file.

## **Details**

The functions used are:

1. p\_value\_function extracts and return the p\_values of HL and intensity segments respectively.

2. eliminate\_outlier\_hl eliminates outliers from HL fragments.

3. eliminate\_outlier\_int eliminates outliers from intensity fragments.

 $4. \ mean\_length\_int\ calculates\ the\ mean\ of\ the\ log2FC (intensity)\ fragments\ adapted\ to\ HL\_fragments\ and\ their\ lengths$ 

5. mean\_length\_hl calculates the mean of log2FC(HL) fragments adapted to the intensity fragments and their lengths.

6. calculating\_rate calculates decay rate and log2FC(intensity). Both are used to calculate synthesis rate.

#### Value

The data frame with the corresponding columns:

position: Integer, position of the first fragment

region: String, region annotation covering the fragments

**gene:** String, gene annotation covering the fragments

locus\_tag: String, locus\_tag annotation covering the fragments

**strand:** Boolean. The bin/probe specific strand (+/-)

fragment\_HL: String, HL fragments

fragment\_int: String, intensity fragments

position\_frg\_int: Integer, position of the first fragment and the last position of the last fragment

mean\_HL\_fragment: Integer, mean of the HL of the fragments involved

mean\_int\_fragment: Integer, mean of the intensity of the fragments involved

**log2FC**(decay\_rate): Integer, log2FC(decay(condition1)/ decay(condition2))

log2FC(synthesis\_rate): Integer, sum of log2FC(decay\_rate) and log2FC(intensity)

**intensity\_FC:** Integer, log2FC(mean(intensity(condition1))/mean(intensity(condition2)))

**Log2FC(HL)+Log2FC(int):** Integer, sum of log2FC(decay\_rate) and log2FC(intensity)

**p\_value:** String, indicated by "\*" means at least one fragment either HL fragment or intensity fragment has a significant p\_value

4 annot\_g

#### **Examples**

```
data(stats_df_comb_minimal)
data(annot_g)
df_mean_minimal <- adjusting_HLToInt(data = stats_df_comb_minimal,
annotation = annot_g[[1]])</pre>
```

annot\_g

The result of gff3\_preprocessing of gff3 file A list containing all necessary information from a gff file for adjusting\_HLToInt and visualization.

# **Description**

The result of gff3\_preprocessing of gff3 file A list containing all necessary information from a gff file for adjusting\_HLToInt and visualization.

#### Usage

```
data(annot_g)
```

#### **Format**

A list with 2 items:

data annotation: a data frame with 5853 rows and 6 variables

region: the region from the gff file start: the start of the annotation end: the end of the annotation strand: the strand of the annotation gene: the annotated gene name locus\_tag: the annotated locus tag

genome length: a numeric vector containing the length of the genome

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

data\_combined\_minimal The result of joining\_by\_row for inp\_s and inp\_f example data A data frame containing the output of joining\_by\_row as a data frame

#### **Description**

The result of joining\_by\_row for inp\_s and inp\_f example data A data frame containing the output of joining\_by\_row as a data frame

# Usage

data(data\_combined\_minimal)

#### **Format**

A data frame with 600 rows and 49 variables:

**strand:** The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**FLT:** The bin/probe flag for background level **intensity:** The relative intensity at time point 0

**probe\_TI:** An internal value to determine which fitting model is applied

flag: Information on which fitting model is appliedposition\_segment: The position based segment

**delay:** The delay value of the bin/probe **half life:** The half-life of the bin/probe

**TI\_termination\_factor:** The termination factor of the bin/probe

delay\_fragment: The delay fragment the bin belongs to

velocity\_fragment: The velocity value of the respective delay fragmentintercept: The vintercept of fit through the respective delay fragmentslope: The slope of the fit through the respective delay fragment

**HL\_fragment:** The half-life fragment the bin belongs to

**HL\_mean\_fragment:** The mean half-life value of the respective half-life fragment

intensity\_fragment: The intensity fragment the bin belongs to

intensity\_mean\_fragment: The mean intensity value of the respective intensity fragment

TU: The overarching transcription unit

**TI\_termination\_fragment:** The TI fragment the bin belongs to

TI\_mean\_termination\_factor: The mean termination factor of the respective TI fragment

seg\_ID: The combined ID of the fragment

pausing\_site: presence of pausing site indicated by +/-

iTSS\_I: presence of iTSS\_I indicated by +/-

**ps\_ts\_fragment:** The fragments involved in pausing site or iTSS\_I

event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I

p\_value\_slope: p\_value of the slope

**delay\_frg\_slope:** the slope value of the respective delay fragment **velocity\_ratio:** Integer, ratio of velocity between 2 delay fragments **event\_duration:** Integer, the duration between two delay fragments

event\_position: Integer, the position middle between 2 fragments with an event

**FC\_fragment\_HL:** Integer, the fold change value of 2 intensity fragments

FC\_HL: Integer, the fold change value of 2 HL fragments#'

**p\_value\_HL:** p\_value of the fold change of HL fragments

**FC\_intensity:** Integer, the fold change value of 2 intensity fragments

FC\_fragment\_intensity: String, fragments involved in fold change between 2 intensity fragments

**p\_value\_intensity:** p\_value of the fold change of intensity fragments

**FC\_HL\_intensity:** ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_intensity\_fragment:** fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_adapted:** Integer, the fold change of half-life/ fold change of intensity, position of the half-life fragment is adapted to intensity fragment

synthesis\_ratio: Integer, the value corresponding to synthesis rate

synthesis\_ratio\_event: String, the event assigned by synthesis rate either Termination or iTSS

p\_value\_Manova: p\_value of the variance between two fold-changes, HL and intensity

p\_value\_TI: p\_value of TI fragment

**TI\_fragments\_p\_value:** p\_value of 2 TI fragments

**cdt:** The condition assigned to the experiment here cdt2

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

#### Source

https://github.com/CyanolabFreiburg/rifi

df\_comb\_minimal 7

df_comb_minimal	The result of joining_by_column for data_combined_minimal example data A data frame containing the output of joining_by_row as a data
	frame

# **Description**

The result of joining\_by\_column for data\_combined\_minimal example data A data frame containing the output of joining\_by\_row as a data frame

#### Usage

```
data(df_comb_minimal)
```

#### **Format**

A data frame with 300 rows and 18 variables:

**strand:** The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**intensity.cdt1:** The relative intensity at time point 0 for condition 1

position\_segment: The position based segment

half\_life.cdt1: The half-life of the bin/probe condition 1

**TI\_termination\_factor.cdt1:** The termination factor of the bin/probe condition 1

**HL\_fragment.cdt1:** The half-life fragment the bin belongs to condition 1

intensity\_fragment.cdt1: The intensity fragment the bin belongs to condition 1TI termination fragment.cdt1: The TI fragment the bin belongs to condition 1

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

intensity.cdt2: The relative intensity at time point 0 condition 2

half\_life.cdt2: The half-life of the bin/probe condition 2

**TI\_termination\_factor.cdt2:** The termination factor of the bin/probe condition 2

**HL\_fragment.cdt2:** The half-life fragment the bin belongs to condition 2

intensity\_fragment.cdt2: The intensity fragment the bin belongs to condition 2

TI\_termination\_fragment.cdt2: The TI fragment the bin belongs to condition 2

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

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df_mean_minimal	The result of adjusting_HLToInt for stats_df_comb_minimal and an-
	notation example data A data frame containing the output of adjusting_HLToInt as a data frame

# Description

The result of adjusting\_HLToInt for stats\_df\_comb\_minimal and annotation example data A data frame containing the output of adjusting\_HLToInt as a data frame

#### Usage

```
data(df_mean_minimal)
```

#### **Format**

A data frame with 52 rows and 15 variables:

position: The bin/probe specific positionregion: the region from the gff filegene: the annotated gene namelocus\_tag: the annotated locus tag

strand: The strand specific

fragment\_HL: The half-life fragment the bin belongs
fragment\_int: The intensity fragment the bin belongs

position\_frg\_int: The position of the first fragment and the last position of the last fragment

mean\_HL\_fragment: The mean half-life value of the respective half-life fragments mean\_int\_fragment: The mean intensity value of the respective intensity fragments

log2FC(decay\_rate): log2FC(decay(condition1)/decay(condition2))

**Log2FC(HL)-Log2FC(int):** log2FC(decay\_rate/intensity)

log2FC(synthesis\_rate): log2FC(decay\_rate) + log2FC(intensity)

**intensity\_FC:** log2FC(mean(intensity(condition1))/mean(intensity(condition2)))

**p\_value:** indicated by "\*" means at least one fragment either HL fragment or intensity fragment has a significant p\_value

## Source

https://github.com/CyanolabFreiburg/rifiComparative

differential\_expression

differential\_expression

An example data frame from Synechosystis PCC 6803 differential probes expression obtained from limma package and only interesting variables were selected. The data frame was used entirely.

# Description

An example data frame from Synechosystis PCC 6803 differential probes expression obtained from limma package and only interesting variables were selected. The data frame was used entirely.

# Usage

```
data(differential_expression)
```

#### **Format**

A data frame of differential\_expression with 55508 rows and 4 variables:

position: The bin/probe specific position

**strand:** The strand specific

**logFC\_int:** The bin/probe differential expression

**P.Value:** The bin/probe p\_value adjusted

#### **Source**

https://github.com/CyanolabFreiburg/rifiComparative

figures\_fun

'figures\_fun': generates several plots

# **Description**

'figures\_fun' plots at one the density of HL, the HL category as histogram, log2FC of decay and synthesis rate and their heatscatter. Scatter plot of HL and volcano plot. The function uses the four output generated previously.

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#### Usage

```
figures_fun(
   data.1,
   data.2,
   input.1,
   input.2,
   cdt1,
   cdt2,
   y = 30,
   x = 30,
   limits = c(0, 20)
```

# **Arguments**

data.1	data frame output of statistic
data.2	data frame joining two outputs from rifi_stats by row
input.1	data frame joining two outputs from rifi_stats by column
input.2	data frame of differential gene expression at time 0
cdt1	string for the first condition
cdt2	string for the second condition
У	integer to break the scaling in scatter plot for y_axis
X	integer to break the scaling in scatter plot for x_axis
limits	vector to limit the scaling in scatter plot for both axis

# **Details**

The functions used are:

plot\_decay\_synt: plots the changes in RNA decay rates versus the changes in RNA synthesis rates plot\_heatscatter: plots the changes in RNA decay rates versus the changes in RNA synthesis rates with density.

plot\_volcano: plots statistical significance versus magnitude of change .

plot\_histogram: plot a histogram of probe/bin half-life categories from 2 to 20 minutes in both conditions.

plot\_density: plots the probe/bin half-life density in both conditions.

plot\_scatter: plots of the bin/probe half-life in one condition1 vs. condition2.

extract the object of rifi\_statistics from both conditions. The differential gene expression at time 0 needs to be run separately. The columns log2FC, p\_value adjusted, position and strand are extracted and saved to a data frame. loading\_fun\_fig joins the differential gene expression table and the output from rifi statistics into one data frame.

### Value

several plots

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## **Examples**

```
data(data_combined_minimal)
data(df_comb_minimal)
data(differential_expression)
data(df_mean_minimal)
figures_fun(data.1 = df_mean_minimal, data.2 = data_combined_minimal,
input.1 = df_comb_minimal, input.2 = differential_expression, cdt1 = "sc",
cdt2 = "fe")
```

fragmentation

fragmentation: Conveniently wraps all fragmentation steps

# Description

fragmentation fragments the half-life and intensity into segments using the penalties output.

# Usage

```
fragmentation(data, pen_HL, pen_int, cores = 2)
```

# **Arguments**

data	data frame: data frame combined data by column
pen_HL	list: list of the penalties set optimal for the fragmentation for half-life
pen_int	list: list of the penalties set optimal for the fragmentation for intensity
cores	integer: the number of assigned cores for the task. It needs to be increased in case of big data.

## Value

Two data frames with half-life and intensity fragments and the mean of the coefficient fragment based.

# **Examples**

```
data(penalties_df)
data(pen_HL)
data(pen_int)
df_comb_minimal <- fragmentation(data = penalties_df, pen_HL,
pen_int)</pre>
```

12 fragment\_int

fragment_int	The result of fragmentation for df_comb_minimal example data A data
	frame containing the output of fragmentation as a data frame

#### **Description**

The result of fragmentation for df\_comb\_minimal example data A data frame containing the output of fragmentation as a data frame

#### Usage

```
data(fragment_int)
```

#### **Format**

A data frame with 500 rows and 24 variables:

**strand:** The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**intensity.cdt1:** The relative intensity at time point 0 for condition 1

position\_segment: The position based segment

half\_life.cdt1: The half-life of the bin/probe condition 1

**TI\_termination\_factor.cdt1:** The termination factor of the bin/probe condition 1

**HL\_fragment.cdt1:** The half-life fragment the bin belongs to condition 1

**intensity\_fragment.cdt1:** The intensity fragment the bin belongs to condition 1 **TI\_termination\_fragment.cdt1:** The TI fragment the bin belongs to condition 1

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

**intensity.cdt2:** The relative intensity at time point 0 condition 2

half\_life.cdt2: The half-life of the bin/probe condition 2

**TI\_termination\_factor.cdt2:** The termination factor of the bin/probe condition 2

**HL\_fragment.cdt2:** The half-life fragment the bin belongs to condition 2

**intensity\_fragment.cdt2:** The intensity fragment the bin belongs to condition 2

**TI\_termination\_fragment.cdt2:** The TI fragment the bin belongs to condition 2

distance\_HL: The bin/probe difference of half-life from both conditions

distance\_int: The bin/probe log2 fold change of intensity at time 0

**HL\_comb\_fragment:** The half-life fragment the bin belongs to both conditions

**HL** mean comb fragment: The half-life mean of the fragment the bin belongs to both conditions

intensity comb fragment: The intensity fragment the bin belongs to both conditions

**intensity\_mean\_comb\_fragment:** The intensity mean of the fragment the bin belongs to both conditions

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#### **Source**

https://github.com/CyanolabFreiburg/rifiComparative

gff3\_preprocess

gff3\_preprocess processes gff3 file from database

# **Description**

gff3\_preprocess processes the gff3 file extracting gene names and locus\_tag from all coding regions (CDS). UTRs/ncRNA/asRNA if available, are also extracted. The resulting dataframe contains region, positions, strand, gene and locus\_tag.

# Usage

```
gff3_preprocess(path)
```

# **Arguments**

path

path: path to the directory containing the gff3 file.

# Value

A list with 2 items:

data annotation: region: String, the region from the gff file

**start:** Integer, the start of the annotation **end:** Integer, the end of the annotation

strand: Boolean, the strand of the annotation

**gene:** String, the annotated gene name **locus\_tag:** String, the annotated locus tag

genome length: a numeric vector containing the length of the genome

# Examples

```
gff3_preprocess(
path = gzfile(system.file("extdata", "gff_synechocystis_6803.gff.gz",
package = "rifiComparative"))
)
```

inp\_f

inp_f	The result of loading_fun for stats_se_cdt2 example data Two data frame containing the output of loading_fun as second element of a
	list.

#### **Description**

The result of loading\_fun for stats\_se\_cdt2 example data Two data frame containing the output of loading\_fun as second element of a list.

# Usage

```
data(inp_f)
```

#### **Format**

A data frame with 500 rows and 49 variables:

strand: The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**FLT:** The bin/probe flag for background level **intensity:** The relative intensity at time point 0

probe\_TI: An internal value to determine which fitting model is applied

flag: Information on which fitting model is appliedposition\_segment: The position based segment

**delay:** The delay value of the bin/probe **half\_life:** The half-life of the bin/probe

**TI\_termination\_factor:** The termination factor of the bin/probe

delay\_fragment: The delay fragment the bin belongs to

**velocity\_fragment:** The velocity value of the respective delay fragment **intercept:** The vintercept of fit through the respective delay fragment

**slope:** The slope of the fit through the respective delay fragment

**HL\_fragment:** The half-life fragment the bin belongs to

**HL\_mean\_fragment:** The mean half-life value of the respective half-life fragment

intensity\_fragment: The intensity fragment the bin belongs to

intensity\_mean\_fragment: The mean intensity value of the respective intensity fragment

TU: The overarching transcription unit

**TI\_termination\_fragment:** The TI fragment the bin belongs to

TI\_mean\_termination\_factor: The mean termination factor of the respective TI fragment

seg\_ID: The combined ID of the fragment

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pausing\_site: presence of pausing site indicated by +/-

iTSS\_I: presence of iTSS\_I indicated by +/-

ps\_ts\_fragment: The fragments involved in pausing site or iTSS\_I

event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I

**p\_value\_slope:** p\_value of the slope

**delay\_frg\_slope:** the slope value of the respective delay fragment **velocity\_ratio:** Integer, ratio of velocity between 2 delay fragments **event\_duration:** Integer, the duration between two delay fragments

event\_position: Integer, the position middle between 2 fragments with an event

**FC\_HL:** Integer, the fold change value of 2 HL fragments

**FC\_fragment\_HL:** Integer, the fold change value of 2 intensity fragments

**p\_value\_HL:** p\_value of the fold change of HL fragments

**FC\_intensity:** Integer, the fold change value of 2 intensity fragments

FC\_fragment\_intensity: String, fragments involved in fold change between 2 intensity fragments

**p\_value\_intensity:** p\_value of the fold change of intensity fragments

**FC\_HL\_intensity:** ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_intensity\_fragment:** fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_adapted:** Integer, the fold change of half-life/ fold change of intensity, position of the half-life fragment is adapted to intensity fragment

synthesis\_ratio: Integer, the value corresponding to synthesis rate

synthesis\_ratio\_event: String, the event assigned by synthesis rate either Termination or iTSS

p\_value\_Manova: p\_value of the variance between two fold-changes, HL and intensity

p\_value\_TI: p\_value of TI fragment

**TI\_fragments\_p\_value:** p\_value of 2 TI fragments

**cdt:** The condition assigned to the experiment here cdt2

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

inp\_s

inp\_s

The result of loading\_fun for stats\_se\_cdt1 example data Two data frame containing the output of loading\_fun as first element of a list.

#### **Description**

The result of loading\_fun for stats\_se\_cdt1 example data Two data frame containing the output of loading\_fun as first element of a list.

# Usage

```
data(inp_s)
```

#### **Format**

A data frame with 500 rows and 49 variables:

strand: The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**FLT:** The bin/probe flag for background level **intensity:** The relative intensity at time point 0

**probe\_TI:** An internal value to determine which fitting model is applied

flag: Information on which fitting model is appliedposition\_segment: The position based segment

**delay:** The delay value of the bin/probe **half life:** The half-life of the bin/probe

**TI\_termination\_factor:** The termination factor of the bin/probe

delay\_fragment: The delay fragment the bin belongs to

velocity\_fragment: The velocity value of the respective delay fragmentintercept: The vintercept of fit through the respective delay fragmentslope: The slope of the fit through the respective delay fragment

**HL\_fragment:** The half-life fragment the bin belongs to

**HL\_mean\_fragment:** The mean half-life value of the respective half-life fragment

intensity\_fragment: The intensity fragment the bin belongs to

intensity\_mean\_fragment: The mean intensity value of the respective intensity fragment

TU: The overarching transcription unit

TI\_termination\_fragment: The TI fragment the bin belongs to

TI\_mean\_termination\_factor: The mean termination factor of the respective TI fragment

seg\_ID: The combined ID of the fragment

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pausing\_site: presence of pausing site indicated by +/-

iTSS\_I: presence of iTSS\_I indicated by +/-

ps\_ts\_fragment: The fragments involved in pausing site or iTSS\_I

event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I

p\_value\_slope: p\_value of the slope

**delay\_frg\_slope:** the slope value of the respective delay fragment **velocity\_ratio:** Integer, ratio of velocity between 2 delay fragments **event\_duration:** Integer, the duration between two delay fragments

event\_position: Integer, the position middle between 2 fragments with an event

**FC\_HL:** Integer, the fold change value of 2 HL fragments

FC\_fragment\_HL: Integer, the fold change value of 2 intensity fragments

**p\_value\_HL:** p\_value of the fold change of HL fragments

**FC\_intensity:** Integer, the fold change value of 2 intensity fragments

FC\_fragment\_intensity: String, fragments involved in fold change between 2 intensity fragments

**p\_value\_intensity:** p\_value of the fold change of intensity fragments

**FC\_HL\_intensity:** ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_intensity\_fragment:** fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_adapted:** Integer, the fold change of half-life/ fold change of intensity, position of the half-life fragment is adapted to intensity fragment

synthesis\_ratio: Integer, the value corresponding to synthesis rate

synthesis\_ratio\_event: String, the event assigned by synthesis rate either Termination or iTSS

p\_value\_Manova: p\_value of the variance between two fold-changes, HL and intensity

p\_value\_TI: p\_value of TI fragment

**TI\_fragments\_p\_value:** p\_value of 2 TI fragments

cdt: The condition assigned to the experiment here cdt1

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

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joining\_data\_column joining\_data\_column joins two data frames by column

#### **Description**

'joining\_data\_column': joins two data frames from different conditions by column.

# Usage

```
joining_data_column(data)
```

## **Arguments**

data

data frame with joined columns from both conditions

#### Value

The data frame with joined columns from both conditions with the corresponding columns: strand, position, ID, intensity.cdt1, position\_segment, half\_life.cdt1, TI\_termination\_factor.cdt1", HL\_fragment.cdt1, intensity\_fragment.cdt1, TI\_termination\_fragment.cdt1, logFC\_int, P.Value, intensity.cdt2, half\_life.cdt2, TI\_termination\_factor.cdt2, HL\_fragment.cdt2, intensity\_fragment.cdt2, TI\_termination\_fragment.cdt2. cdt1: first condition, cdt2: second condition.

#### **Examples**

```
data(data_combined_minimal)
df_comb_minimal <- joining_data_column(data = data_combined_minimal)</pre>
```

joining\_data\_row

joining\_data\_row joins two data frames by row

# **Description**

joining\_data\_row joins two data frames from different conditions by row.

# Usage

```
joining_data_row(input1, input2)
```

#### **Arguments**

input1 data frame from SummarizedExperiment output of rifi\_stats from rifi package

of the first condition.

input2 data frame from SummarizedExperiment output of rifi\_stats from rifi package

of the second condition.

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#### Value

The data frame with joined rows from both conditions with the corresponding columns: ID with position, strand, intensity, probe\_TI, flag, position\_segment, delay, half\_life, TI\_termination\_factor, delay\_fragment, velocity\_fragment, intercept, slope, HL\_fragment, HL\_mean\_fragment, intensity\_fragment, intensity\_mean\_fragment, TU, TI\_termination\_fragment, TI\_mean\_termination\_factor, seg\_ID, pausing\_site, iTSS\_I, ps\_ts\_fragment, event\_ps\_itss\_p\_value\_Ttest, p\_value\_slope, delay\_frg\_slope, velocity\_ratio, event\_duration, event\_position, FC\_HL, FC\_fragment\_HL, p\_value\_HL, FC\_intensity, FC\_fragment\_intensity, p\_value\_intensity, FC\_HL\_intensity, FC\_HL\_intensity\_fragment, FC\_HL\_adapted, synthesis\_ratio, synthesis\_ratio\_event, p\_value\_Manova, p\_value\_TI, cdt (condition), logFC\_int (log2FC(intensity)), P.Value of log2FC(intensity)

#### **Examples**

```
data(inp_s)
data(inp_f)
data_combined_minimal <-
joining_data_row(input1 = inp_s, input2 = inp_f)</pre>
```

loading\_fun

loading fun loads the data of all conditions

## **Description**

loading\_fun extract outputs from rifi\_stats of all conditions and join each data to the differential expression table. The differential gene expression at time 0 needs to be run separately. The columns log2FC, p\_value adjusted, position and strand are extracted and saved to a data frame. loading\_fun joins the differential gene expression table and the output from rifi statistics into one data frame.

#### **Usage**

```
loading_fun(data1, data2, data3)
```

## **Arguments**

data1	data frame from rifi_stats of one condition
data2	data frame from rifi_stats of other condition
data3	data frame from differential gene expression at time 0

#### Value

A list of two data frames with joined columns from differential expression and output of rifi\_stats with the corresponding columns: ID with position, strand, intensity, probe\_TI, flag, position\_segment, delay, half\_life, TI\_termination\_factor, delay\_fragment, velocity\_fragment, intercept, slope, HL\_fragment, HL\_mean\_fragment, intensity\_mean\_fragment, TU, TI\_termination\_fragment, TI\_mean\_termination\_factor, seg\_ID, pausing\_site, iTSS\_I, ps\_ts\_fragment, event\_ps\_itss\_p\_value\_Ttest, p\_value\_slope, delay\_frg\_slope, velocity\_ratio, event\_duration, event\_position, FC\_HL, FC\_fragment\_HL, p\_value\_HL, FC\_intensity, FC\_fragment\_intensity, p\_value\_intensity, FC\_HL\_intensity, FC\_HL\_intensity\_fragment,

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FC\_HL\_adapted, synthesis\_ratio, synthesis\_ratio\_event, p\_value\_Manova, p\_value\_TI, cdt (condition), logFC\_int (log2FC(intensity)), P.Value of log2FC(intensity).

# **Examples**

```
data(stats_se_cdt1)
data(stats_se_cdt2)
data(differential_expression)
inp_s <-
loading_fun(stats_se_cdt1, stats_se_cdt2, differential_expression)[[1]]
inp_f <-
loading_fun(stats_se_cdt1, stats_se_cdt2, differential_expression)[[2]]</pre>
```

make\_pen

make\_pen assigns automatically penalties

## **Description**

make\_pen calls one of four available penalty functions to automatically assign penalties for the dynamic programming. The two functions to be called are:

- 1. fragment\_HL\_pen
- 2. fragment\_inty\_pen

# Usage

```
make_pen(
   probe,
   FUN,
   cores = 1,
   logs,
   dpt = 1,
   smpl_min = 10,
   smpl_max = 100,
   sta_pen = 0.5,
   end_pen = 4.5,
   rez_pen = 9,
   sta_pen_out = 0.5,
   end_pen_out = 3.5,
   rez_pen_out = 7)
```

#### **Arguments**

probe data frame: data frame combined data by column

FUN function: one of the four bottom level functions (see details)

cores integer: the number of assigned cores for the task

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logs numeric vector: the logbook vector. integer: the number of times a full iteration cycle is repeated with a more narrow dpt range based on the previous cycle. integer: the smaller end of the sampling size. smpl\_min smpl\_max integer: the larger end of the sampling size. numeric: the lower starting penalty. sta\_pen end\_pen numeric: the higher starting penalty. numeric: the number of penalties iterated within the penalty range. rez\_pen numeric: the lower starting outlier penalty. sta\_pen\_out end\_pen\_out numeric: the higher starting outlier penalty. numeric: the number of outlier penalties iterated within the outlier penalty rez\_pen\_out

#### **Details**

The two functions called return the amount of statistically correct and statistically wrong splits at a specific pair of penalties. 'make\_pen' iterates over many penalty pairs and picks the most suitable pair based on the difference between wrong and correct splits. The sample size, penalty range and resolution as well as the number of cycles can be customized. The primary start parameters create a matrix with n = rez\_pen rows and n = rez\_pen\_out columns with values between sta\_pen/sta\_pen\_out and end\_pen/end\_pen\_out. The best penalty pair is picked. If dept is bigger than 1 the same process is repeated with a new matrix of the same size based on the result of the previous cycle. Only position segments with length within the sample size range are considered for the penalties to increase run time.

#### Value

A list with 4 items:

**logbook:** The logbook vector containing all penalty information **penalties:** a vector with the respective penalty and outlier penalty

correct: a matrix of the correct splitswrong: a matrix of the incorrect splits

range.

# Examples

```
data(df_comb_minimal)

df_comb_minimal$distance_HL <- df_comb_minimal$half_life.cdt1 -

df_comb_minimal$half_life.cdt2

df_comb_minimal$distance_int <- df_comb_minimal$logFC_int

pen_HL <- make_pen(
    probe = df_comb_minimal, FUN = rifiComparative:::fragment_HL_pen,
    cores = 2, logs = as.numeric(rep(NA, 8)), dpt = 1, smpl_min = 10,
    smpl_max = 50, sta_pen = 0.5, end_pen = 4.5, rez_pen = 9, sta_pen_out = 0.5,</pre>
```

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```
end_pen_out = 3.5, rez_pen_out = 7
)

pen_int <- make_pen(
    probe = df_comb_minimal, FUN = rifiComparative:::fragment_inty_pen,
    cores = 2, logs = as.numeric(rep(NA, 8)), dpt = 1, smpl_min = 10,
    smpl_max = 50, sta_pen = 0.5, end_pen = 4.5, rez_pen = 9, sta_pen_out = 0.5,
    end_pen_out = 3.5, rez_pen_out = 7
)</pre>
```

penalties

penalties wraps conveniently all penalty steps

#### **Description**

penalties finds the best set of penalties for half-life and intensity fragmentation using dynamic programming. The segmentation of the HL uses the difference between 2 conditions and the segmentation of the intensity uses the log2FC.

#### Usage

```
penalties(data, cores = 2)
```

# **Arguments**

data frame with the joined columns from differential expression and output of

rifi\_stats.

cores integer: the number of assigned cores for the task. It needs to be increased in

case of big data.

# Details

```
The function uses 4 functions:
```

```
score_fun_ave.r
make_pen.r
fragment_HL_pen.r
fragment_inty_pen.r
```

#### Value

A list of data frame and penalties, The first element is data frame with 2 more variables, second and third are HL and intensity penalties respectively.

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## **Examples**

```
data(df_comb_minimal)
penalties_df <- penalties(df_comb_minimal)[[1]]
pen_HL <- penalties(df_comb_minimal)[[2]]
pen_int <- penalties(df_comb_minimal)[[3]]</pre>
```

penalties\_df

The result of penalties for df\_comb\_minimal example data A data frame containing the output of penalties as a data frame

#### **Description**

The result of penalties for df\_comb\_minimal example data A data frame containing the output of penalties as a data frame

#### Usage

```
data(penalties_df)
```

#### **Format**

A data frame with 300 rows and 20 variables:

**strand:** The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

intensity.cdt1: The relative intensity at time point 0 for condition 1

position\_segment: The position based segment

half life.cdt1: The half-life of the bin/probe condition 1

**TI termination factor.cdt1:** The termination factor of the bin/probe condition 1

**HL\_fragment.cdt1:** The half-life fragment the bin belongs to condition 1

**intensity\_fragment.cdt1:** The intensity fragment the bin belongs to condition 1 **TI termination fragment.cdt1:** The TI fragment the bin belongs to condition 1

logFC\_int: The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

intensity.cdt2: The relative intensity at time point 0 condition 2

half\_life.cdt2: The half-life of the bin/probe condition 2

**TI termination factor.cdt2:** The termination factor of the bin/probe condition 2

**HL\_fragment.cdt2:** The half-life fragment the bin belongs to condition 2

**intensity\_fragment.cdt2:** The intensity fragment the bin belongs to condition 2

TI\_termination\_fragment.cdt2: The TI fragment the bin belongs to condition 2

**distance HL:** The bin/probe difference of half-life from both conditions

**distance\_int:** The bin/probe log2 fold change of intensity at time 0

pen\_int

#### **Source**

https://github.com/CyanolabFreiburg/rifiComparative

pen_HL	The result of penalties for df_comb_minimal example data. A list containing the output from penalties including the logbook and two penalty objects.

# Description

The result of penalties for df\_comb\_minimal example data. A list containing the output from penalties including the logbook and two penalty objects.

# Usage

```
data(pen_HL)
```

## **Format**

A list with 5 items:

pen\_obj\_HL: A list with 4 items:

**logbook:** The logbook vector containing half-life penalty information **HL\_penalties:** a vetor with the half-life penalty and half-life outlier penalty

**correct:** a matrix of the correct splits **wrong:** a matrix of the incorrect splits

# Source

```
https://github.com/CyanolabFreiburg/rifi
```

pen_int	The result of penalties for df_comb_minimal example data. A list containing the output from penalties including the logbook and two
	penalty objects.

# **Description**

The result of penalties for df\_comb\_minimal example data. A list containing the output from penalties including the logbook and two penalty objects.

## Usage

```
data(pen_int)
```

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#### **Format**

A list with 5 items:

pen\_int: A list with 4 items:

logbook: The logbook vector containing intensity penalty information

int\_penalties: a vector with the intensity penalty and intensity outlier penalty

**correct:** a matrix of the correct splits **wrong:** a matrix of the incorrect splits

#### **Source**

https://github.com/CyanolabFreiburg/rifi

rifiComparative

%

rifiComparative

rifiComparative a successor package of rifi. It compares 2 rifi outputs from 2 different conditions of the same organism.

# Description

rifiComparative was developed to compare 2 rifi outputs from 2 conditions. The rifi output may differ significantly from 2 conditions. The complexity of the segments number, position, length and the events make the comparison between 2 conditions nearly impossible. rifiComparative uses a simple strategy to generate single segments for both conditions, extract the features and make them comparable.

## **Details**

Five major steps ate described in rifiComparative:

- 1. Joining data
- 2. penalties
- 3. fragmentation
- 4. statistics
- 5. visualization

# Author(s)

```
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```

```
rifi_visualization_comparison

rifi_visualization_comparison plots the segments and genome annotation
```

# Description

rifi\_visualization\_comparison plots the genome annotation, half-life difference (HL), log2FC(intensity) fragments. It uses several functions to plot TUs and genes including small-RNAs. Additionally it plots the statistical t-test between the neighboring fragment, significant p-values from t-test are assigned with '\*'.

#### Usage

```
rifi_visualization_comparison(
  data,
  data_c,
  genomeLength = annot_g[[2]],
  annot = annot_g[[1]],
  condition = c("cdt1", "cdt2"),
  Strand = c("+", "-"),
  region = c("CDS", "asRNA", "5'UTR", "ncRNA", "3'UTR", "tRNA"),
 color_region = c("grey0", "red", "blue", "orange", "yellow", "green", "white",
    "darkseagreen1", "grey50", "black"),
  color_TU = c("cyan", "yellow", "orange"),
  scaling_TU = c(0, 3.4, 6.6),
  color_text.1 = "grey0",
  color_text.2 = "black",
  Alpha = 0.5,
  size_tu = 1.6,
  size_locusTag = 1.6,
  size\_gene = 1.6,
  Limit = 10,
  shape = 22,
  face = "bold",
  tick_length = 0.3,
  arrow.color = "darkseagreen1",
  col_above20 = "#00FFFF",
  fontface = "plain",
  shape_above20 = 14,
  axis_text_y_size = 3,
  axis_title_y_size = 6,
  iTSS_threshold = 1.2,
  p_value_manova = 0.05,
  termination_threshold = 0.8
)
```

## **Arguments**

data dataframe: the probe based dataframe with joined columns.

data\_c dataframe: the probe based dataframe with joined rows.

genomeLength integer: genome length output of gff3\_preprocess function.

annot dataframe: the annotation file.

condition string: assigned as cdt1 (condition 1) and cdt2 (condition2), it could be adapted

to any name.

Strand string: either ("+" or "-").

region dataframe: gff3 features of the genome.

color\_region string vector: vector of colors.

color\_TU string. TU color

scaling\_TU vector: values to adjusted termination and iTSSs to TUs.

color\_text.1 string: TU color text
color\_text.2 string: genes color text

Alpha integer: color transparency degree.

size\_tu integer: TU size

size\_locusTag integer: locus\_tag size

size\_gene integer: font size for gene annotation.

Limit integer: value for y-axis limit.

shape integer: value for shape.

face string: label font.

tick\_length integer: value for ticks.
arrow.color string: arrows color.

col\_above20 string: color for probes/bin above value 20.

fontface integer: font type

shape\_above20 integer: shape for probes/bins above value 20.

axis\_text\_y\_size

integer: text size for y-axis.

axis\_title\_y\_size

integer: title size for y-axis.

iTSS\_threshold integer: threshold for iTSS\_II selected to plot, default 1.2.

p\_value\_manova integer: p\_value of manova test fragments to plot, default 0.05.

termination\_threshold

integer: threshold for termination to plot, default .8.

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#### **Details**

The functions used are:

strand\_selection: plots HL, intensity fragments from both strands.

splitGenome\_function: splits the genome into fragments.

annotation\_plot\_comp: plots the corresponding annotation.

indice\_function: assign a new column to the data to distinguish between fragments, outliers from delay or HL or intensity.

empty\_data\_positive: plots empty boxes in case no data is available for positive strand

empty\_data\_negative: plots empty boxes in case no data is available for negative strand

TU\_annotation: designs the segments border for the genes and TUs annotation.

gene\_annot\_function: it requires gff3 file, returns a dataframe adjusting each fragment according to its annotation. It allows as well the plot of genes and TUs shared into two pages.

secondaryAxis: adjusts the half-life or delay to 20 in case of the dataframe row numbers is equal to 1 and the half-life or delay exceed the limit, they are plotted with different shape and color.

my\_arrow: creates an arrow for the annotation.

arrange\_byGroup: selects the last row for each segment and add 40 nucleotides in case of negative strand for a nice plot.

my\_segment\_T: plots terminals and pausing sites labels.

#### Value

The plot.

## **Examples**

```
data(data_combined_minimal)
data(stats_df_comb_minimal)
data(annot_g)
rifi_visualization_comparison(
    data = data_combined_minimal,
    data_c = stats_df_comb_minimal,
    genomeLength = annot_g[[2]],
    annot = annot_g[[1]]
    )
```

statistics

statistics check segments significance using statistical test

#### **Description**

statistics uses t-test to check HL and intensity segments significance. The function returns the data frame with p\_value and p\_value adjusted. The function used is t\_test\_function.

## Usage

```
statistics(data)
```

#### **Arguments**

data

data frame: data frame output of fragmentation

#### Value

A list of two data frames, the first one contains all segments with p\_value and p\_value adjusted. The second one removes the duplicated segments from intensity and could be saved as an excel file.

# **Examples**

```
data(fragment_int)
stats_df_comb_minimal <- statistics(data= fragment_int)[[1]]
df_comb_uniq_minimal <- statistics(data= fragment_int)[[2]]</pre>
```

stats\_df\_comb\_minimal The result of statistics for fragment\_int example data A data frame containing the output of statistics as a data frame

# Description

The result of statistics for fragment\_int example data A data frame containing the output of statistics as a data frame

#### Usage

```
data(stats_df_comb_minimal)
```

### Format

A data frame with 500 rows and 26 variables:

strand: The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

intensity.cdt1: The relative intensity at time point 0 for condition 1

position\_segment: The position based segment

half\_life.cdt1: The half-life of the bin/probe condition 1

**TI\_termination\_factor.cdt1:** The termination factor of the bin/probe condition 1

**HL\_fragment.cdt1:** The half-life fragment the bin belongs to condition 1

intensity\_fragment.cdt1: The intensity fragment the bin belongs to condition 1

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TI\_termination\_fragment.cdt1: The TI fragment the bin belongs to condition 1

**logFC\_int:** The bin/probe log2 fold change of intensity at time 0

**P.Value:** The bin/probe p\_value adjusted

intensity.cdt2: The relative intensity at time point 0 condition 2

half\_life.cdt2: The half-life of the bin/probe condition 2

**TI\_termination\_factor.cdt2:** The termination factor of the bin/probe condition 2

**HL\_fragment.cdt2:** The half-life fragment the bin belongs to condition 2

 $intensity\_fragment.cdt2$ : The intensity fragment the bin belongs to condition 2

**TI\_termination\_fragment.cdt2:** The TI fragment the bin belongs to condition 2

distance\_HL: The bin/probe difference of half-life from both conditions

**distance\_int:** The bin/probe log2 fold change of intensity at time 0

**HL\_comb\_fragment:** The half-life fragment the bin belongs to both conditions

**HL\_mean\_comb\_fragment:** The half-life mean of the fragment the bin belongs to both conditions

intensity\_comb\_fragment: The intensity fragment the bin belongs to both conditions

**intensity\_mean\_comb\_fragment:** The intensity mean of the fragment the bin belongs to both conditions

p\_value\_distance\_HL: The p\_value adjusted of the half-life fragment the bin belongs to both conditions

p\_value\_distance\_intensity: The p\_value adjusted of the intensity fragment the bin belongs to both conditions

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

stats_se_cdt1	An example SummarizedExperiment from Synechosystis PCC 6803
	first condition obtained from rifi_statistics and used as input for ri- fiComparative

# Description

An example SummarizedExperiment from Synechosystis PCC 6803 first condition obtained from rifi\_statistics and used as input for rifiComparative

### Usage

```
data(stats_se_cdt1)
```

stats\_se\_cdt1 31

#### **Format**

A rowRanges of SummarizedExperiment with 500 rows and 50 variables:

seqnames: The sequence name chromosome

start: The bin/probe start positionend: The bin/probe end positionwidth: The bin/probe lengthstrand: The strand specific

position: The bin/probe specific position

**ID:** The bin/probe specific ID

**FLT:** The bin/probe flag for background level **intensity:** The relative intensity at time point 0

probe\_TI: An internal value to determine which fitting model is applied

flag: Information on which fitting model is applied position segment: The position based segment

**delay:** The delay value of the bin/probe **half\_life:** The half-life of the bin/probe

**TI\_termination\_factor:** The termination factor of the bin/probe

delay\_fragment: The delay fragment the bin belongs to

velocity\_fragment: The velocity value of the respective delay fragmentintercept: The vintercept of fit through the respective delay fragmentslope: The slope of the fit through the respective delay fragment

**HL\_fragment:** The half-life fragment the bin belongs to

**HL\_mean\_fragment:** The mean half-life value of the respective half-life fragment

intensity\_fragment: The intensity fragment the bin belongs to

intensity mean fragment: The mean intensity value of the respective intensity fragment

TU: The overarching transcription unit

**TI termination fragment:** The TI fragment the bin belongs to

TI mean termination factor: The mean termination factor of the respective TI fragment

seg\_ID: The combined ID of the fragment

pausing\_site: presence of pausing site indicated by +/-

iTSS\_I: presence of iTSS\_I indicated by +/-

ps\_ts\_fragment: The fragments involved in pausing site or iTSS\_I
event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I#'
delay\_frg\_slope: the slope value of the respective delay fragment

**p\_value\_slope:** p\_value of the slope

velocity\_ratio: Integer, ratio of velocity between 2 delay fragmentsevent\_duration: Integer, the duration between two delay fragments

stats\_se\_cdt2

event\_position: Integer, the position middle between 2 fragments with an event

**FC\_HL:** Integer, the fold change value of 2 HL fragments

**FC\_fragment\_HL:** Integer, the fold change value of 2 intensity fragments

**p\_value\_HL:** p\_value of the fold change of HL fragments

**FC\_intensity:** Integer, the fold change value of 2 intensity fragments

FC\_fragment\_intensity: String, fragments involved in fold change between 2 intensity fragments

p\_value\_intensity: p\_value of the fold change of intensity fragments

**FC\_HL\_intensity:** ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_intensity\_fragment:** fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_adapted:** Integer, the fold change of half-life/ fold change of intensity, position of the half-life fragment is adapted to intensity fragment

synthesis\_ratio: Integer, the value corresponding to synthesis rate

synthesis\_ratio\_event: String, the event assigned by synthesis rate either Termination or iTSS

p\_value\_Manova: p\_value of the variance between two fold-changes, HL and intensity

**p\_value\_TI: p\_value** of TI fragment

**TI\_fragments\_p\_value:** p\_value of 2 TI fragments

#### Source

https://github.com/CyanolabFreiburg/rifiComparative

stats_se_cdt2	An example SummarizedExperiment from Synechosystis PCC 6803 second condition obtained from rifi_statistics and used as input for rifiComparative
	rijiComparative

#### **Description**

An example SummarizedExperiment from Synechosystis PCC 6803 second condition obtained from rifi\_statistics and used as input for rifiComparative

#### **Usage**

data(stats\_se\_cdt2)

stats\_se\_cdt2 33

#### **Format**

A rowRanges of SummarizedExperiment with 500 rows and 50 variables:

seqnames: The sequence name chromosome

start: The bin/probe start positionend: The bin/probe end positionwidth: The bin/probe lengthstrand: The strand specific

**position:** The bin/probe specific position

**ID:** The bin/probe specific ID

**FLT:** The bin/probe flag for background level **intensity:** The relative intensity at time point 0

probe\_TI: An internal value to determine which fitting model is applied

flag: Information on which fitting model is applied position segment: The position based segment

**delay:** The delay value of the bin/probe **half\_life:** The half-life of the bin/probe

**TI\_termination\_factor:** The termination factor of the bin/probe

delay\_fragment: The delay fragment the bin belongs to

velocity\_fragment: The velocity value of the respective delay fragmentintercept: The vintercept of fit through the respective delay fragmentslope: The slope of the fit through the respective delay fragment

**HL\_fragment:** The half-life fragment the bin belongs to

**HL\_mean\_fragment:** The mean half-life value of the respective half-life fragment

intensity\_fragment: The intensity fragment the bin belongs to

intensity mean fragment: The mean intensity value of the respective intensity fragment

TU: The overarching transcription unit

**TI termination fragment:** The TI fragment the bin belongs to

TI mean termination factor: The mean termination factor of the respective TI fragment

seg\_ID: The combined ID of the fragment

pausing\_site: presence of pausing site indicated by +/-

iTSS\_I: presence of iTSS\_I indicated by +/-

ps\_ts\_fragment: The fragments involved in pausing site or iTSS\_I
event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I

**p\_value\_slope:** p\_value of the slope

**delay\_frg\_slope:** the slope value of the respective delay fragment **velocity\_ratio:** Integer, ratio of velocity between 2 delay fragments **event\_duration:** Integer, the duration between two delay fragments

stats\_se\_cdt2

event\_position: Integer, the position middle between 2 fragments with an event

FC\_HL: Integer, the fold change value of 2 HL fragments

FC\_fragment\_HL: Integer, the fold change value of 2 intensity fragments

**p\_value\_HL:** p\_value of the fold change of HL fragments

**FC\_intensity:** Integer, the fold change value of 2 intensity fragments

FC\_fragment\_intensity: String, fragments involved in fold change between 2 intensity fragments

**p\_value\_intensity:** p\_value of the fold change of intensity fragments

**FC\_HL\_intensity:** ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_intensity\_fragment:** fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments

**FC\_HL\_adapted:** Integer, the fold change of half-life/ fold change of intensity, position of the half-life fragment is adapted to intensity fragment

synthesis\_ratio: Integer, the value corresponding to synthesis rate

synthesis\_ratio\_event: String, the event assigned by synthesis rate either Termination or iTSS

p\_value\_Manova: p\_value of the variance between two fold-changes, HL and intensity

p\_value\_TI: p\_value of TI fragment

TI fragments p value: p value of 2 TI fragments

#### **Source**

https://github.com/CyanolabFreiburg/rifiComparative

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