# Package 'rifi'

# January 2, 2025

**Title** 'rifi' analyses data from rifampicin time series created by microarray or RNAseq

#### Version 1.11.0

Description 'rifi' analyses data from rifampicin time series created by microarray or RNAseq.

'rifi' is a transcriptome data analysis tool for the holistic identification of transcription and decay associated processes.

The decay constants and the delay of the onset of decay is fitted for each probe/bin. Subsequently, probes/bins of equal properties

are combined into segments by dynamic programming, independent of a existing genome annotation. This allows to detect transcript segments

of different stability or transcriptional events within one annotated gene.

In addition to the classic decay constant/half-

life analysis, 'rifi' detects processing sites, transcription pausing sites, internal transcription start sites in operons, sites of partial transcription termination in operons, identifies ar-

eas of likely transcriptional interference by the collision mechanism

and gives an estimate of the transcription velocity.

All data are integrated to give an estimate of continous transcriptional units, i.e. operons. Comprehensive output tables and visualizations of the full genome result and the individual fits for all probes/bins are produced.

# **Depends** R (>= 4.2)

**Imports** car, cowplot, doMC, parallel, dplyr, egg, foreach, ggplot2, graphics, grDevices, grid, methods, nls2, nnet, rlang, S4Vectors, scales, stats, stringr, SummarizedExperiment, tibble, rtracklayer, reshape2, utils

Suggests DescTools, devtools, knitr, rmarkdown, BiocStyle

# VignetteBuilder knitr

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

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

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apply\_ancova

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# 

apply\_ancova checks the variances between 2 segments showing either pausing site (ps) or internal starting site (ITSS) independently.

# Description

apply\_ancova is a statistical test to check if fragments showing ps and ITSS events have significant slope using Ancova test. The function uses ancova test. Ancova is applied when the data contains independent variables, dependent variables and covariant variables. In this case, segments are independent variables, position is the dependent variable and the delay is the covariant.

# Usage

apply\_ancova(inp)

# Arguments

inp

SummarizedExperiment: the input data frame with correct format.

# Value

the SummarizedExperiment with the columns regarding statistics:

ID:	The bin/probe specific ID.	
position:	The bin/probe specific position.	
strand:	The bin/probe specific strand.	
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_segmer	nt:	
	The position based segment.	
delay:	The delay value of the bin/probe.	
half_life:	The half-life of the bin/probe.	
TI_termination	_factor:	
	String, the factor of TI fragment.	
delay_fragment:		
	The delay fragment the bin belongs to.	
velocity_fragme	ent:	
	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.	
<pre>intensity_mean_fragment:</pre>		
	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\_duration: Integer, the duration between two delay fragments. event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I. p\_value\_slope: Integer, the p\_value added to the inp delay\_frg\_slope: Integer, the slope value of the fit through the respective delay fragment velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments

# Examples

```
data(stats_minimal)
apply_ancova(inp = stats_minimal)
```

apply\_event\_position %

# apply\_event\_position

apply\_event\_position extracts event time duration for pausing site or iTSS

# Description

apply\_event\_position is a short version of apply\_Ttest\_delay function to extract event time duration for pausing site or iTSS. Its adds a new column with the duration.

### Usage

```
apply_event_position(inp)
```

#### Arguments

inp

SummarizedExperiment: the input data frame with correct format.

The SummarizedExperiment with the columns regarding statistics:

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

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

strand: The bin/probe specific strand.

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: String, the factor of TI fragment.

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.

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\_duration: Integer, the duration between two delay fragments.

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

**p\_value\_slope:** Integer, the p\_value added to the inp.

delay\_frg\_slope: Integer, the slope value of the fit through the respective delay fragment.

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments.

event\_position: Integer, position of the event added to the input.

#### Examples

data(stats\_minimal)
apply\_event\_position(inp = stats\_minimal)

apply_manova	%
	======================================
	apply_manova checks if the ratio of hl ratio and intensity ratio is statistically significant.

# Description

apply\_manova compares the variance between two fold-changes HL and intensity within the same TU (half-life frgA/half-life frgB/intensity frgA/intensity frgB). HL fragment could cover two intensity fragments therefore this function sets first fragments borders and uses manova\_function. Manova checks the variance between 2 segments (independent variables) and two dependents variables (HL and intensity).

## Usage

apply\_manova(inp)

### Arguments

inp

SummarizedExperiment: the input data frame with correct format.

#### Value

The probe data frame with the columns regarding statistics:

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

position: The bin/probe specific position.

strand: The bin/probe specific strand.

**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: String, the factor of TI fragment.

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.

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\_duration: Integer, the duration between two delay fragments.

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

**p\_value\_slope:** Integer, the p\_value added to the inp.

delay\_frg\_slope: Integer, the slope value of the fit through the respective delay fragment.

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments.

event\_position: Integer, position of the event added to the input.

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

FC\_fragment\_HL: String, the fragments corresponding to HL fold change.

**p\_value\_HL:** Integer, the p\_value added to the input of 2 HL fragments.

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

FC\_fragment\_intensity: String, the fragments corresponding to intensity fold change.

p\_value\_intensity: Integer, the p\_value added to the input of 2 intensity fragments.

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

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

FC\_HL\_intensity: Integer, the value corresponding to HL and intensity fold change.

FC\_HL\_intensity\_fragment: String, the fragments corresponding to intensity and HL fold change.

**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.

p\_value\_Manova: Integer, the p\_value added to the input.

# Examples

```
data(stats_minimal)
apply_manova(inp = stats_minimal)
```

apply\_Ttest\_delay %

#### apply\_Ttest\_delay

apply\_Ttest\_delay checks the significance of the point between 2 segments showing pausing site (ps) and internal starting site (ITSS) independently

# Description

apply\_Ttest\_delay: is a statistical test to check the significance of the point between 2 segments showing pausing site (ps) and internal starting site (ITSS) independently. The function uses t-test. The last point from the first segment and the first point from the second segment are selected and added to the residuals of each model. The sum is subjected to t-test.

# Usage

apply\_Ttest\_delay(inp)

### Arguments

inp

SummarizedExperiment: the input data frame with correct format.

### Value

the SummarizedExperiment with the columns regarding statistics:

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

position: The bin/probe specific position.

strand: The bin/probe specific strand.

**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: String, the factor of TI fragment.

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.

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\_duration: Integer, the duration between two delay fragments.

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

### Examples

```
data(stats_minimal)
apply_Ttest_delay(inp = stats_minimal)
```

apply\_t\_test

#### \_\_\_\_\_

%

# apply\_t\_test

apply\_t\_test uses the statistical t\_test to check if the fold-change of half -life (HL) fragments and the fold-change intensity fragments respectively are significant.

# Description

apply\_t\_test compares the mean of two neighboring fragments within the same TU to check if the fold-change is significant.Fragments with distance above threshold are not subjected to t-test.Dataframes with less than 3 rows are excluded.

# Usage

```
apply_t_test(inp, threshold = 300)
```

# Arguments

inp	SummarizedExperiment: the input data frame with correct format.
threshold	integer: threshold.

### Details

The functions used are:

- 1. fragment\_function: checks number of fragments inside TU, less than 2 are excluded otherwise they are gathered for analysis.
- 2. t\_test\_function: excludes dataframes with less than 3 rows, makes fold-change and apply t-test, assign fragments names and ratio, add columns with the corresponding p\_values.

# Value

the SummarizedExperiment with the columns regarding statistics:

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

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

strand: The bin/probe specific strand.

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: String, the factor of TI fragment.

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.

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\_duration: Integer, the duration between two delay fragments.

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

**p\_value\_slope:** Integer, the p\_value added to the inp.

**delay\_frg\_slope:** Integer, the slope value of the fit through the respective delay fragment.

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments.

event\_position: Integer, position of the event added to the input.

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

FC\_fragment\_HL: String, the fragments corresponding to HL fold change.

**p\_value\_HL:** Integer, the p\_value added to the input of 2 HL fragments.

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

FC\_fragment\_intensity: String, the fragments corresponding to intensity fold change.

**p\_value\_intensity:** Integer, the p\_value added to the input of 2 intensity fragments.

#### Examples

```
data(stats_minimal)
apply_t_test(inp = stats_minimal, threshold = 300)
```

%

apply\_t\_test\_ti

## apply\_t\_test\_ti

apply\_t\_test\_ti compares the mean of two neighboring TI fragments within the same TU.

### Description

apply\_t\_test\_ti uses the statistical t\_test to check if two neighboring TI fragments are significant.

#### Usage

apply\_t\_test\_ti(inp)

#### Arguments

inp SummarizedExperiment: the input data frame with correct format.

#### Value

the SummarizedExperiment with the columns regarding statistics:

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

position: The bin/probe specific position.

strand: The bin/probe specific strand.

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: String, the factor of TI fragment. **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. 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\_duration: Integer, the duration between two delay fragments. event\_ps\_itss\_p\_value\_Ttest: p\_value of pausing site or iTSS\_I. **p** value slope: Integer, the p value added to the inp. **delay\_frg\_slope:** Integer, the slope value of the fit through the respective delay fragment. velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments. event position: Integer, position of the event added to the input. FC\_HL: Integer, the fold change value of 2 HL fragments. FC fragment HL: String, the fragments corresponding to HL fold change. **p\_value\_HL:** Integer, the p\_value added to the input of 2 HL fragments. FC intensity: Integer, the fold change value of 2 intensity fragments. FC\_fragment\_intensity: String, the fragments corresponding to intensity fold change. **p** value intensity: Integer, the p value added to the input of 2 intensity fragments. synthesis\_ratio: Integer, the value corresponding to synthesis rate. synthesis ratio event: String, the event assigned by synthesis rate either Termination or iTSS. FC HL intensity: Integer, the value corresponding to HL and intensity fold change. FC\_HL\_intensity\_fragment: String, the fragments corresponding to intensity and HL fold change. 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. p\_value\_Manova: Integer, the p\_value added to the input. **p\_value\_TI:** Integer, the p\_value added to the input.

TI\_fragments\_p\_value: String, the fragments subjected to statistical test.

## Examples

```
data(stats_minimal)
apply_t_test_ti(inp = stats_minimal)
```

check\_input

#### Check\_input

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%

Check\_input reviews the input given by the user

# Description

check\_input stops the operation if the input data frame has severe faults. Less severe faults lead to the removal of wrong IDs and a warnings describing the problem. The Summarized Experiment colData must have the columns "timepoint" with the timepoints convertible to numeric and containing the timepoint 0. If replicates are used the column in colData must be called "replicate". The replicate must be convertible to numeric. In the RowRanges, optionally, IDs can be given as character (except ",","|","\_"), but need to refer to a unique position/strand combination. Strand information needs to be given. The relative intensity in the assay must be numeric. The relative intensity for the first time point cannot be 0 or NA.

# Usage

check\_input(inp, thrsh = 0)

### Arguments

inp	SummarizedExperiment: the input data frame with correct format
thrsh	numeric: the minimal allowed intensity for time point "0".

### Value

the SummarizedExperiment object: checked, and with position, ID and filtration added to the rowRanges.

# Examples

```
data(example_input_minimal)
check_input(inp = example_input_minimal, thrsh = 0)
```

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dataframe\_summary

#### dataframe\_summary

%

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dataframe\_summary creates two tables relating gene annotation to fragments

# Description

dataframe\_summary creates two tables summary of segments and their half-lives. The first output is bin/probe features and the second one is intensity fragment based. The dataframe\_summary creates one table with feature\_type, gene, locus\_tag, position, strand, TU, delay\_fragment, HL\_fragment, half\_life, intensity\_fragment, intensity and velocity. The second table is similar to the first one but in compact form. It contains the same columns, the only difference is on position where a start and end position are indicated separately.

# Usage

dataframe\_summary(data, input)

#### Arguments

data	SummarizedExperiment: the input data frame with correct format.
input	dataframe: dataframe from event_dataframe function.

### Value

bin_df:	all information regarding bins:
	position: Integer, position of the bin/probe on the genome
	feature_type: 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 (+/-)
	segment: String, the bin/probe segment on the genome
	TU: String, The overarching transcription unit
	delay_fragment: The delay fragment the bin belongs to
	delay: Integer, the delay value of the bin/probe
	HL_fragment: The half-life fragment the bin belongs to
	half_life: The half-life of the bin/probe
	intensity_fragment: The intensity fragment the bin belongs to
	intensity: The relative intensity at time point 0
	flag: String, the flag of the bin/probe, contains information or the distribution
	for the #'different fitting models

	<b>TI_termination_factor:</b> String, the TI termination factor determined by TI
frag_df:	all information regarding fragments:
	feature_type: String, region annotation covering the fragments
	gene: String, gene annotation covering the fragments
	locus_tag: String, locus_tag annotation covering the fragments
	first_position_frg: Integer, the bin/probe specific first position
	last_position_frg: Integer, the bin/probe specific last position
	strand: Boolean. The bin/probe specific strand (+/-)
	TU: String, The overarching transcription unit
	segment: String, the bin/probe segment on the genome
	delay_fragment: String, the delay fragment the bin belongs to
	<b>HL_fragment:</b> Integer, the half_life fragment of the bin/probe belongs to
	half_life: Integer, the half-life of the bin/probe
	HL_SD: Integer, the half-life standard deviation of the HL fragment, bin/probe based
	HL_SE: Integer, the half-life standard error of the HL fragment, bin/probe based
	intensity_fragment: Integer, the intensity fragment the bin belongs to
	intensity: Integer, the relative intensity of bin/probe at time point 0
	<b>intensity_SD:</b> Integer, the intensity standard deviation of the intensity frag- ment, bin/probe based
	intensity_SE: Integer, the intensity standard error of the intensity fragment, bin/probe based
	velocity: The velocity value of the respective delay fragment

# Examples

```
data(stats_minimal)
data(res_minimal)
dataframe_summary(data = stats_minimal, input = res_minimal)
```

dataframe\_summary\_events %

dataframe\_summary\_events

dataframe\_summary\_events creates one table with all events between the segments

#### Description

dataframe\_summary\_events creates one table with the following columns: event, features, p\_value, event\_position, event\_duration, position, region, gene, locus\_tag, strand, TU, segment\_1, segment\_2, length, velocity\_ratio, FC\_HL, FC\_intensity, FC\_HL/FC\_intensity.

### Usage

dataframe\_summary\_events(data, data\_annotation)

### Arguments

data SummarizedExperiment: the input data frame with correct format.

data\_annotation

dataframe: dataframe from processed gff3 file.

### Value

event: String, event type either pausing site, iTSS\_I, iTSS\_II, Termination, HL\_event, Int\_event, HL\_Int\_event and velocity\_change

**p\_value:** Integer, **p\_value** of the event

p\_adjusted: Integer, p\_value adjusted

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

FC\_intensity: Integer, the fold change value of 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: Fold change of half-life/ fold change of intensity

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

velocity\_ratio: Integer, ratio of velocity between 2 delay fragments

feature\_type: 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 (+/-)

TU: String, The overarching transcription unit

segment\_1: String, the first segment of the event, includes the segment, TU, delay fragment in case
 of ps or iTSS\_I. The rest of the events include HL fragment and intensity fragment

segment\_2: String, same description as segment\_1 but is the second fragment of the event

event\_duration: Integer, the difference (min) between 2 delay fragment when ps or iTSS\_I happen

**gap\_fragments:** Integer, length in position (nt), calculated by the difference between the last position of the first fragment and the first position of the second fragment.

features: Integer, number of fragements involved on the event

# Examples

```
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
data(stats_minimal)
dataframe_summary_events(data = stats_minimal,
data_annotation = metadata(stats_minimal)$annot[[1]])
```

dataframe\_summary\_events\_HL\_int

%

# dataframe\_summary\_events\_HL\_int

dataframe\_summary\_events\_HL\_int creates one table with all events between the segments

# Description

The dataframe\_summary\_events\_HL\_int creates one table with the following columns: event, features, p\_value, event\_position, position, region, gene, locus\_tag, strand, TU, segment\_1, segment\_2, length, FC\_HL, FC\_intensity, FC\_HL/FC\_intensity.

# Usage

dataframe\_summary\_events\_HL\_int(data, data\_annotation)

### Arguments

data SummarizedExperiment: the input data frame with correct format.

data\_annotation

dataframe: dataframe from processed gff3 file.

### Value

event: String, event type.

p\_value: Integer, p\_value of the event.

p\_adjusted: Integer, p\_value adjusted.

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

FC\_intensity: Integer, the fold change value of 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.

FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity.

## 18

feature\_type: 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 (+/-).

TU: String, The overarching transcription unit.

- segment\_1: String, the first segment of the event, includes the segment, TU, delay fragment in case of ps or iTSS\_I. The rest of the events include HL fragment and could be extended intensity fragment.
- segment\_2: String, the second fragment of the two of fragments subjected to analysis.
- event\_duration: Integer, the difference (min) between 2 delay fragment when ps or iTSS\_I happen.
- **gap\_fragments:** Integer, length in position (nt), calculated by the difference between the last position of the first fragment and the first position of the second fragment.

features: Integer, number of fragements involved on the event

# Examples

```
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
data(stats_minimal)
dataframe_summary_events_HL_int(data = stats_minimal,
data_annotation = metadata(stats_minimal)$annot[[1]])
```

dataframe\_summary\_events\_ps\_itss

dataframe\_summary\_events\_ps\_itss creates one table with all events between the segments.

# Description

The dataframe\_summary\_events\_ps\_itss creates one table with the following columns: event, features, p\_value, event\_position, event\_duration, position, region, gene, locus\_tag, strand, TU, segment\_1, segment\_2, length, velocity\_ratio.

# Usage

```
dataframe_summary_events_ps_itss(data, data_annotation)
```

# Arguments

data SummarizedExperiment: the input data frame with correct format.

data\_annotation

dataframe: dataframe from processed gff3 file.

### Value

event: String, event type.

**p\_value:** Integer, p\_value of the event.

**p\_adjusted:** Integer, p\_value adjusted.

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

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments.

feature\_type: 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 (+/-).

TU: String, The overarching transcription unit.

- segment\_1: String, the first segment of the event, includes the segment, TU, delay fragment in case of ps or iTSS\_I. The rest of the events include HL fragment and could be extended intensity fragment.
- segment\_2: String, the second fragment of the two of fragments subjected to analysis.
- event\_duration: Integer, the difference (min) between 2 delay fragment when ps or iTSS\_I happen.
- **gap\_fragments:** Integer, length in position (nt), calculated by the difference between the last position of the first fragment and the first position of the second fragment.
- features: Integer, number of fragements involved on the event

### Examples

```
data(stats_minimal)
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
dataframe_summary_events_ps_itss(data = stats_minimal,
data_annotation = metadata(stats_minimal)$annot[[1]])
```

dataframe\_summary\_events\_velocity

dataframe\_summary\_events\_velocity creates one table with all events between the segments.

# Description

The dataframe\_summary\_events\_velocity creates one table with the following columns: event, features, p\_value, event\_position, event\_duration, position, region, gene, locus\_tag, strand, TU, segment\_1, segment\_2, length, velocity\_ratio.

### Usage

dataframe\_summary\_events\_velocity(data, data\_annotation)

#### Arguments

data SummarizedExperiment: the input data frame with correct format.

data\_annotation

dataframe: dataframe from processed gff3 file.

# Value

event: String, event type.

**p\_value:** Integer, p\_value of the event.

p\_adjusted: Integer, p\_value adjusted.

**event\_position:** Integer, the position of event, calculated dividing the last position of the first fragment and the first position of the next fragment on 2.

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments

feature\_type: 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 (+/-).

TU: String, The overarching transcription unit.

segment\_1: String, the first segment of the event, includes the segment, TU, delay fragment in case
of ps or iTSS\_I. The rest of the events include HL fragment and could be extended intensity
fragment.

segment\_2: String, the second fragment of the two of fragments subjected to analysis

- event\_duration: Integer, the difference (min) between 2 delay fragment when ps or iTSS\_I happen.
- **gap\_fragments:** Integer, length in position (nt), calculated by the difference between the last position of the first fragment and the first position of the second fragment.

features: Integer, number of fragements involved on the event.

# Examples

```
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
data(stats_minimal)
dataframe_summary_events_velocity(data = stats_minimal,
data_annotation = metadata(stats_minimal)$annot[[1]])
```

dataframe\_summary\_TI %

#### dataframe\_summary\_TI

dataframe\_summary\_TI creates one table with all TI fragments,p\_value and the coordinates

# Description

The dataframe\_summary creates one table with the following columns: event, TI\_fragment, TI\_factor, TI\_fragments\_TU, p\_value, feature\_type, gene, locus\_tag, strand, TU, features, event\_position, position\_1 and position\_2.

#### Usage

```
dataframe_summary_TI(data, input)
```

# Arguments

data	SummarizedExperiment: the input data frame with correct format.
input	dataframe: dataframe from event_dataframe function.

# Value

event: String, event type, transcription interference.

TI\_fragment: String, the fragment with TI.

TI\_termination\_factor: String, the factor of TI fragment.

**p\_value:** Integer, p\_value of the event.

**p\_adjusted:** Integer, p\_value adjusted.

feature\_type: 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 (+/-).

TU: String, The overarching transcription unit.

Features: Integer, The number of segments within the TI.

event\_position: Integer, the position middle between 2 TI fragments.

- **position\_1:** String, the first position of TI fragment, if 2 fragments, first position is from the first fragment.
- **position\_2:** String, the last position of TI fragment, if 2 fragments, last position is from the second fragment.

WIP

# Examples

```
data(stats_minimal)
data(res_minimal)
dataframe_summary_TI(data = stats_minimal, input = res_minimal)
```

event\_dataframe

# event\_dataframe

%

event\_dataframe creates a dataframe only with events

# Description

event\_dataframe creates a dataframe connecting segments, events and the annotation.

### Usage

```
event_dataframe(data, data_annotation)
```

### Arguments

data dataframe: the probe based data frame.

data\_annotation

dataframe: the coordinates are extracted from gff3

# Details

The functions used are:

position\_function: adds the specific position of ps or iTSS event.

annotation\_function\_event: adds the events to the annotated genes.

annotation file needs to be supplied. Strand is indicated in case of stranded data The event\_dataframe selects columns with statistical features. ID, position, strand and TU columns are required.

#### Value

A dataframe with unique intensity fragments

feature\_type: 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 (+/-)

TU: String, The overarching transcription unit

position: Integer, position of the bin/probe on the genome

segment: String, the bin/probe segment on the genome

FC\_fragment\_intensity: String, the fragments subjected to fold change

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

**p\_value\_intensity:** Integer, **p\_value** of the FC\_intensity

FC\_fragment\_HL: String, the fragments subjected to fold change

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

p\_value\_HL: Integer, p\_value of the FC\_HL

FC\_HL\_FC\_intensity\_fragment: String, fragments subjected to FC\_HL/FC\_intensity

- FC\_HL\_FC\_intensity: Integer, the value of FC\_HL/FC\_intensity
- **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
- p\_value\_manova: Integer, p\_value of the event FC\_HL/FC\_intensity

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

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

pausing\_site: Boolean, presence or absence of pausing\_site event (ps)

iTSS\_I: Boolean, presence or absence of internal starting site event (iTSS\_I)

ps\_ts\_fragment: String, fragments involved on the event ps or iTSS\_I

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

event\_duration: Integer, the duration between two delay fragments

delay: Integer, the delay value of the bin/probe

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

intensity: The relative intensity at time point 0

delay\_frg\_slope: Integer, the slope value of the fit through the respective delay fragment

p\_value\_slope: Integer, the p\_value added to the inp

### example\_input\_e\_coli

# Examples

```
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
data(stats_minimal)
event_dataframe(data = stats_minimal,
data_annotation = metadata(stats_minimal)$annot[[1]])
```

example\_input\_e\_coli An example SummarizedExperiment from E. coli An example SummarizedExperiment from RNA-seq containing information about the intensities at all time points (assay). Seqnames, IRanges and strand columns (rowRanges)and colData with time point series and replicates.

# Description

An example SummarizedExperiment from E. coli An example SummarizedExperiment from RNAseq containing information about the intensities at all time points (assay). Seqnames, IRanges and strand columns (rowRanges)and colData with time point series and replicates.

# Usage

```
data(example_input_e_coli)
```

### Format

A assay:

- 0: relative intensities at 0 min
- 1: relative intensities at 1 min
- 10: relative intensities at 10 min
- 15: relative intensities at 15 min
- 2: relative intensities at 2 min
- 20: relative intensities at 20 min
- 3: relative intensities at 3 min
- 4: relative intensities at 4 min
- 5: relative intensities at 5 min
- 6: relative intensities at 6 min
- 8: relative intensities at 8 min

# Source

https://github.com/CyanolabFreiburg/rifi

example\_input\_minimal An artificial example SummarizedExperiment An example SummarizedExperiment containing information about the intensities at all time points (assay). Seqnames, IRanges and strand columns (rowRanges) and colData with time point series and replicates.

# Description

An artificial example SummarizedExperiment An example SummarizedExperiment containing information about the intensities at all time points (assay). Seqnames, IRanges and strand columns (rowRanges) and colData with time point series and replicates.

# Usage

```
data(example_input_minimal)
```

# Format

An object of class RangedSummarizedExperiment with 4 rows and 33 columns.

#### Source

https://github.com/CyanolabFreiburg/rifi

example\_input\_synechocystis\_6803

An example input data frame from Synechocystis PCC 6803 A SummarizedExperiment from microarrays data containing information about the intensities at all time points (assay), Seqnames, IRanges and strand columns (rowRanges) and colData with time point series and averaged replicates.

# Description

An example input data frame from Synechocystis PCC 6803 A SummarizedExperiment from microarrays data containing information about the intensities at all time points (assay), Seqnames, IRanges and strand columns (rowRanges) and colData with time point series and averaged replicates.

### Usage

data(example\_input\_synechocystis\_6803)

# finding\_PDD

# Format

Assay with 3000 rows and 10 variables:

- 0: relative intensities at 0 min
- 2: relative intensities at 2 min
- 4: relative intensities at 4 min
- 8: relative intensities at 8 min
- **16:** relative intensities at 16 min
- 32: relative intensities at 32 min
- 64: relative intensities at 64 min

### Source

https://github.com/CyanolabFreiburg/rifi

%

finding\_PDD

#### ======

#### finding\_PDD

finding\_PDD Flags potential candidates for post transcription decay

# Description

'finding\_PDD' uses 'score\_fun\_linear\_PDD' to make groups by the difference to the slope. The slope is further checked for steepness to decide for PDD. '*PDD*' is added to the 'flag' column. Post transcription decay is characterized by a strong decrease of intensity by position. The rowRanges need to contain at least 'ID', 'intensity', 'position' and 'position\_segment'!

### Usage

```
finding_PDD(inp, cores = 1, pen = 2, pen_out = 1, thrsh = 0.001)
```

# Arguments

inp	SummarizedExperiment: the input.
cores	integer: the number of assigned cores for the task
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Advised to be kept at 2. Default is 2.
pen_out	numeric: an internal parameter for the dynamic programming. Higher values result in fewer possible outliers. Advised to be kept at 1. Default is 1.
thrsh	numeric: an internal parameter that allows fragments with slopes steeper than the thrsh to be flagged with ' <i>PDD</i> '. Higher values result in fewer candidates. Advised to be kept at 0.001. Default is 0.001.

# Value

The SummarizedExperiment object: with "PDD" added to the flag column.

# Examples

```
data(preprocess_minimal)
finding_PDD(inp = preprocess_minimal, cores = 2, pen = 2,
pen_out = 1, thrsh = 0.001)
```

finding\_TI

# finding\_TI

%

==

finding\_TI flags potential candidates for transcription interference

# Description

finding\_TI uses 'score\_fun\_ave' to make groups by the mean of "probe\_TI". "TI" is added to the "flag" column. TI is characterized by relative intensities at time points later than "0". The rowRanges need to contain at least "ID", "probe\_TI" and "position\_segment"!

### Usage

finding\_TI(inp, cores, pen = 10, thrsh = 0.5, add = 1000)

# Arguments

inp	SummarizedExperiment: the input.
cores	integer: the number of assigned cores for the task
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Advised to be kept at 10. Default is 10.
thrsh	numeric: an internal parameter that allows fragments with a certain amount of IDs with higher relative intensities at time points later than "0" to be flagged as " <i>TI</i> ". Higher values result in fewer candidates0.5 is 25 %, 0 is 50%, 0.5 is 75%. Advised to be kept at 0.5. Default is 0.5.
add	integer: range of nucleotides before and after a potential TI event wherein IDs are fitted with the TI fit.

### Value

The SummarizedExperiment object: with "TI" added to the flag column.

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### fit\_e\_coli

# Examples

```
data(preprocess_minimal)
finding_TI(inp = preprocess_minimal, cores = 2, pen = 10, thrsh = 0.5,
add = 1000)
```

fit_e_coli	The result of rifi_fit for E.coli example data A SummarizedExperiment
	containing the output from rifi_fit as an extension of rowRanges and
	metadata.

#### Description

The result of rifi\_fit for E.coli example data A SummarizedExperiment containing the output from rifi\_fit as an extension of rowRanges and metadata.

#### Usage

data(fit\_e\_coli)

# Format

Three data frames with 290 rows and 10 variables, 155 rows and 5 variables, and 135 rows and 9 variables are generated. The columns of the first data frame are added to the rowRanges and the rest are added as metadata.

inp: The SummarizedExperiment:

ID: The bin/probe specific ID
position: The bin/probe specific position
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
postion\_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: String, the factor of TI fragment
fit\_obj\_STD: the fit object for the standard fit:

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

**delay:** The delay value of the bin/probe

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

**inty\_S0:** The relative intensity at time point 0

intyf: The background value of the fit

fit\_obj\_TI: the fit object for the TI fit:

delay: The delay value of the bin/probe

ti\_delay: The ti-delay value of the bin/probe
half\_life: The half-life of the bin/probe
ti\_value: The ti-value of the bin/probe
TI\_termination\_factor: String, the factor of TI fragment
synthesis\_rate: The synthesis rate of the bin/probe
TI\_background: The background value of the fit
position: The bin/probe specific position
ID: The bin/probe specific ID

# Source

https://github.com/CyanolabFreiburg/rifi

fit_minimal	The artificial result of rifi_fit for artificial example data A Summa-
	rizedExperiment containing the output from rifi_fit.

# Description

The artificial result of rifi\_fit for artificial example data A SummarizedExperiment containing the output from rifi\_fit.

# Usage

```
data(fit_minimal)
```

### Format

An object of class RangedSummarizedExperiment with 4 rows and 33 columns.

### Source

https://github.com/CyanolabFreiburg/rifi

fit\_synechocystis\_6803

The result of rifi\_fit for Synechocystis 6803 example data A SummarizedExperiment containing the output from rifi\_fit as an extension of rowRanges and metadata.

# Description

The result of rifi\_fit for Synechocystis 6803 example data A SummarizedExperiment containing the output from rifi\_fit as an extension of rowRanges and metadata.

#### Usage

```
data(fit_synechocystis_6803)
```

#### Format

Three data frames with 3000 rows and 10 variables, 2811 rows and 5 variables, and 189 rows and 9 variable are generated. The columns of the first data frame are added to the rowRanges and the rest are added as metadata.

inp: the SummarizedExperiment:

ID: The bin/probe specific ID
position: The bin/probe specific position
strand: The bin/probe specific strand
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
postion\_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: String, the factor of TI fragment

fit\_obj\_STD: the fit object for the standard fit:

ID: The bin/probe specific IDdelay: The delay value of the bin/probehalf\_life: The half-life of the bin/probeinty\_S0: The relative intensity at time point 0intyf: The background value of the fit

fit\_obj\_TI: the fit object for the TI fit:

**delay:** The delay value of the bin/probe

ti\_delay: The ti-delay value of the bin/probe

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

ti\_value: The ti-value of the bin/probe

### fold\_change

\_\_\_\_\_

TI\_termination\_factor: String, the factor of TI fragment
synthesis\_rate: The synthesis rate of the bin/probe
TI\_background: The background value of the fit
position: The bin/probe specific position
ID: The bin/probe specific ID

### Source

https://github.com/CyanolabFreiburg/rifi

fold_change	%
	fold_change sets a fold-change ratio between the neighboring fragments of Half-life (HL) and intensity

# Description

fold\_change sets fold change on intensity and fold change HL fragments of two successive fragments. Two intensity fragments could belong to one HL fragment. This function sets first the borders using the position and applies the fold change ratio between the neighboring fragments of HL and those from intensity log2(intensity frgA/intensity frgB/half-life frgA/half-life frgB). All grepped fragments are from the same TU excluding outliers.

# Usage

fold\_change(inp)

### Arguments

inp

SummarizedExperiment: the input data frame with correct format.

# Details

The function used is: synthesis\_r\_Function: assigns events depending on the ratio between HL and intensity of two consecutive fragments. intensity(int) = synthesis rate(k)/decay(deg) (steady state), int1/int2 = k1/deg1\*deg2/k2 int1 \* (deg1/int2) \* deg2 = k1/k2 => synthesis ratio. In case of synthesis ratio is: synthesis ratio > 0 -> New start synthesis ratio < 0 -> Termination

# fold\_change

#### Value

the SummarizedExperiment with the columns regarding statistics:

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

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

strand: The bin/probe specific strand.

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: String, the factor of TI fragment.

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.

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\_duration: Integer, the duration between two delay fragments.

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

**p\_value\_slope:** Integer, the p\_value added to the inp.

delay\_frg\_slope: Integer, the slope value of the fit through the respective delay fragment.

velocity\_ratio: Integer, the ratio value of velocity from 2 delay fragments.

event\_position: Integer, position of the event added to the input.

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

FC\_fragment\_HL: String, the fragments corresponding to HL fold change.

**p\_value\_HL:** Integer, the p\_value added to the input of 2 HL fragments.

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

FC\_fragment\_intensity: String, the fragments corresponding to intensity fold change.

**p\_value\_intensity:** Integer, the p\_value added to the input of 2 intensity fragments.

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

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

FC\_HL\_intensity: Integer, the value corresponding to HL and intensity fold change.

- FC\_HL\_intensity\_fragment: String, the fragments corresponding to intensity and HL fold change.
- **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.

#### Examples

```
data(stats_minimal)
fold_change(inp = stats_minimal)
```

fragmentation\_e\_coli The result of rifi\_fragmentation for E.coli example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension of rowRanges

### Description

The result of rifi\_fragmentation for E.coli example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension of rowRanges

#### Usage

data(fragmentation\_e\_coli)

## Format

rowRanges of the SummarizedExperiment with 290 rows and 22 variables:

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

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

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: String, the factor of TI fragment

**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

#### Source

https://github.com/CyanolabFreiburg/rifi

fragmentation\_minimal The result of rifi\_fragmentation for artificial example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension of rowRanges and metadata.

# Description

The result of rifi\_fragmentation for artificial example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension of rowRanges and metadata.

# Usage

```
data(fragmentation_minimal)
```

# Format

An object of class RangedSummarizedExperiment with 24 rows and 33 columns.

#### Source

https://github.com/CyanolabFreiburg/rifi

fragmentation\_synechocystis\_6803

The result of rifi\_fragmentation for Synechocystis 6803 example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension fo rowRanges

# Description

The result of rifi\_fragmentation for Synechocystis 6803 example data A SummarizedExperiment containing the output from rifi\_fragmentation as an extension fo rowRanges

### Usage

data(fragmentation\_synechocystis\_6803)

# Format

rowRanges of the SummarizedExperiment:

ID: The bin/probe specific ID position: The bin/probe specific position 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: String, the factor of TI fragment 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

# Source

https://github.com/CyanolabFreiburg/rifi

fragment\_delay

fragment\_delay

%

fragment\_delay performs the delay fragmentation

## Description

fragment\_delay makes delay\_fragments based on position\_segments and assigns all gathered information to the SummarizedExperiment object. The columns "delay\_fragment", "velocity\_fragment", "intercept" and "slope" are added. fragment\_delay makes delay\_fragments, assigns slopes, which are 1/velocity at the same time, and intercepts for the TU calculation. The function used is: score\_fun\_linear the input is the SummarizedExperiment object. pen is the penalty for new fragments in the dynamic programming, pen\_out is the outlier penalty.

## Usage

fragment\_delay(inp, cores = 1, pen, pen\_out)

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
cores	cores: integer: the number of assigned cores for the task.
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
pen_out	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.

## Value

the SummarizedExperiment object:

ID:	The bin/probe specific ID.	
position:	The bin/probe specific position.	
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:		
	String, the factor of TI fragment.	
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.	

## Examples

```
data(fragmentation_minimal)
fragment_delay(inp = fragmentation_minimal, cores = 2, pen = 2, pen_out = 1)
```

fragment\_HL

%

==

# fragment\_HL

fragment\_HL performs the half\_life fragmentation

## Description

fragment\_HL makes HL\_fragments based on delay\_fragments and assigns all gathered information to the SummarizedExperiment object.

## Usage

```
fragment_HL(inp, cores = 1, pen, pen_out)
```

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
cores	integer: the number of assigned cores for the task.
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
pen_out	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.

## fragment\_HL

#### Details

The columns "HL\_fragment" and "HL\_mean\_fragment" are added.

fragment\_HL makes half-life\_fragments and assigns the mean of each fragment.

The function used is:

.score\_fun\_ave.

The input the SummarizedExperiment object.

pen is the penalty for new fragments in the dynamic programming, pen\_out is the outlier penalty.

#### Value

The SummarizedExperiment object:

ID: The bin/probe specific ID
position: The bin/probe specific position
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: String, the factor of TI fragment
delay\_fragment: The delay fragment the bin belongs to
velocity\_fragment: The velocity value of the respective delay fragment
slope: The slope of the fit through the respective delay fragment
HL\_fragment: The mean half-life value of the respective half-life fragment

#### Examples

data(fragmentation\_minimal)
fragment\_HL(inp = fragmentation\_minimal, cores = 2, pen = 2, pen\_out = 1)

fragment\_inty

%

#### fragment\_inty

fragment\_inty performs the intensity fragmentation

## Description

fragment\_inty makes intensity\_fragments based on HL\_fragments and assigns all gathered information to the SummarizedExperiment object.

## Usage

fragment\_inty(inp, cores = 1, pen, pen\_out)

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
cores	cores: integer: the number of assigned cores for the task.
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
pen_out	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.

## Details

The columns "intensity\_fragment" and "intensity\_mean\_fragment" are added.

fragment\_inty makes intensity\_fragments and assigns the mean of each fragment.

The function used is:

.score\_fun\_ave.

The input is the the SummarizedExperiment object.

pen is the penalty for new fragments in the dynamic programming, pen\_out is the outlier penalty.

## Value

The SummarizedExperiment object:

ID:	The bin/probe specific ID
position:	The bin/probe specific position
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

## fragment\_TI

<pre>position_segment:</pre>		
	The position based segment	
delay:	The delay value of the bin/probe	
half_life:	The half-life of the bin/probe	
TI_termination_	_factor:	
	String, the factor of TI fragment	
delay_fragment:	:	
	The delay fragment the bin belongs to	
velocity_fragme	ent:	
	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	

## Examples

```
data(fragmentation_minimal)
fragment_inty(inp = fragmentation_minimal, cores = 2, pen = 2, pen_out = 1)
```

fragment\_TI

fragment\_TI

%

\_

fragment\_TI performs the TI fragmentation

## Description

fragment\_TI makes TI\_fragments based on TUs and assigns all gathered information to the SummarizedExperiment object. The columns TI\_termination\_fragment" and the TI\_mean\_termination\_factor are added.

# Usage

fragment\_TI(inp, cores = 1, pen, pen\_out)

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
cores	cores: integer: the number of assigned cores for the task.
pen	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
pen_out	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.

## Details

The function used is:

.score\_fun\_ave.

The input is the SummarizedExperiment object.

pen is the penalty for new fragments in the dynamic programming, pen\_out is the outlier penalty.

## Value

The SummarizedExperiment object:

ID:	The bin/probe specific ID
position:	The bin/probe specific position
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_segmer	nt:
	The position based segment
delay:	The delay value of the bin/probe
half_life:	The half-life of the bin/probe
TI_termination_	_factor:
	String, the factor of TI fragment
delay_fragment:	
	The delay fragment the bin belongs to
velocity_fragme	ent:
	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_fragmer	nt:
	The mean half-life value of the respective half-life fragment
intensity_fragm	nent:
	The intensity fragment the bin belongs to
intensity_mean_	_fragment:
	The mean intensity value of the respective intensity fragment

#### gff3\_preprocess

### Examples

```
data(fragmentation_minimal)
fragment_TI(inp = fragmentation_minimal, cores = 2, pen = 2, pen_out = 1)
```

gff3\_preprocess

#### gff3\_preprocess

%

gff3\_preprocess process gff3 file from database for multiple usage

#### Description

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

#### Usage

```
gff3_preprocess(path)
```

### Arguments

path path: path to the directory containing the gff3 file.

## Details

The resulting dataframe contains region, positions, strand, gene and locus\_tag.

## 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_e_coli.gff3.gz", package = "rifi"))
)
```

make\_df

%

#### make\_df

make\_df adds important columns to the SummarizedExperiment object

## Description

'make\_df' adds to the SummarizedExperiment object with the columns: "intensity", "probe\_TI" and "flag".

## Usage

make\_df(inp, cores = 1, bg = 0, rm\_FLT = TRUE)

## Arguments

inp	SummarizedExperiment: the (checked) input.
cores	integer: the number of assigned cores for the task.
bg	numeric: threshold over which the last timepoint has to be fitted with the above background mode.
rm_FLT	logical: remove IDs where all replicates are marked as filtered. Default is FALSE.

## Details

The replicates are collapsed into their respective means.

"intensity" is the mean intensity from time point 0.

"probe\_TI" is a value needed for the distribution for the different fitting models.

"flag" contains information or the distribution for the different fitting models.

Probes that don't reach the background level expression are flagged as "ABG" ("above background"). This is only needed for microarray data and is controlled by the bg parameter. The default for bg = 0, resulting in all probes to be above background (0 is advised for RNAseq data).

Probes where all replicates were filtered in the optional filtration step can be fully removed by rm\_FLT = TRUE! If you wish to keep all information in the assay set to FALSE!

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## make\_pen

## Value

the SummarizedExperiment object: with intensity, probe\_TI and flag added to the rowRanges.

#### Examples

```
data(preprocess_minimal)
make_df(inp = preprocess_minimal, cores = 2, bg = 0, rm_FLT = TRUE)
```

make\_pen

%

====

### make\_pen

make\_pen assigns automatically a penalties

## Description

'make\_pen' calls one of four available penalty functions to automatically assign penalties for the dynamic programming.

#### Usage

```
make_pen(
    inp,
    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

inp	SummarizedExperiment: the input data frame with correct format
FUN	function: one of the four bottom level functions (see details)
cores	integer: the number of assigned cores for the task
logs	numeric vector: the logbook vector.

dpt	integer: the number of times a full iteration cycle is repeated with a more narrow range based on the previous cycle. Default is 2.
smpl_min	integer: the smaller end of the sampling size. Default is 10.
<pre>smpl_max</pre>	integer: the larger end of the sampling size. Default is 100.
sta_pen	numeric: the lower starting penalty. Default is 0.5.
end_pen	numeric: the higher starting penalty. Default is 4.5.
rez_pen	numeric: the number of penalties iterated within the penalty range. Default is 9.
sta_pen_out	numeric: the lower starting outlier penalty. Default is 0.5.
end_pen_out	numeric: the higher starting outlier penalty. Default is 3.5.
rez_pen_out	numeric: the number of outlier penalties iterated within the outlier penalty range. Default is 7.

## Details

The four functions to be called are:

- 1. fragment\_delay\_pen
- 2. fragment\_HL\_pen
- 3. fragment\_inty\_pen
- 4. fragment\_TI\_pen

These functions 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. Returns a penalty object (list of 4 objects) the first being the logbook.

#### Value

A list with 4 items:

logbook: Interger, the logbook vector containing all penalty information

penalties: Integer, a vetor with the respective penalty and outlier penalty

correct: Matrix, a matrix of the correct splits

wrong: Matrix, a matrix of the incorrect splits

## Examples

```
data(fit_minimal)
make_pen(
    inp = fit_minimal, FUN = rifi:::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,
end_pen_out = 3.5, rez_pen_out = 7
)
```

nls2\_fit

%

nls2\_fit estimates decay for each probe or bin

## Description

nls2\_fit uses nls2 function to fit a probe or bin using intensities of the time series data from different time point. nls2 uses different starting values through expand grid and selects the best fit. Different filters could be applied prior fitting to the model.

## Usage

```
nls2_fit(
    inp,
    cores = 1,
    decay = seq(0.01, 0.11, by = 0.02),
    delay = seq(0, 10, by = 0.1),
    k = seq(0.1, 1, 0.2),
    bg = 0.2
)
```

### Arguments

inp	SummarizedExperiment: the input with correct format.
cores	integer: the number of assigned cores for the task.
decay	numeric vector: A sequence of starting values for the decay. Default is seq(.08, 0.11, by=.02)
delay	numeric vector: A sequence of starting values for the delay. Default is seq(0,10, by=.1)
k	numeric vector: A sequence of starting values for the synthesis rate. Default is $seq(0.1,1,0.2)$
bg	numeric vector: A sequence of starting values. Default is 0.2.

#### Details

To apply nls2\_fit function, prior filtration could applied.

- generic\_filter\_BG: filter probes with intensities below background using threshold. Those
  probes are filtered.
- 2. filtration\_below\_backg: additional functions exclusive to microarrays could be applied. Its very strict to the background (not recommended in usual case).
- 3. filtration\_above\_backg: selects probes with a very high intensity and above the background (recommended for special transcripts). Probes are flagged with "*ABG*". Those transcripts are usually related to a specific function in bacteria. This filter selects all probes with the same ID, the mean is applied, the last time point is selected and compared to the threshold.

The model used estimates the delay, decay, intensity of the first time point (synthesis rate/decay) and the background. The coefficients are gathered in vectors with the corresponding IDs. Absence of the fit or a very bad fit are assigned with NA. In case of probes with very high intensities and above the background, the model used makes abstinence of background coefficient. The output of all coefficients is saved in the metadata. The fits are plotted using the function\_plot\_fit.r through rifi\_fit.

## Value

the SummarizedExperiment object: with delay and decay added to the rowRanges. The full fit data is saved in the metadata as "fit\_STD".

delay: Integer, the delay value of the bin/probe

half\_life: Integer, the half-life of the bin/probe

#### Examples

```
data(preprocess_minimal)
nls2_fit(inp = preprocess_minimal, cores = 2)
```

penalties_e_coli	The result of rifi_penalties for E.coli example data. A SummarizedEx-
	periment containing the output from rifi_penalties including the log-
	book and the four penalty objects as metadata.

## Description

The result of rifi\_penalties for E.coli example data. A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata.

#### Usage

```
data(penalties_e_coli)
```

## Format

A list with 5 items:

logbook: The logbook vector containing all penalty information

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

logbook: The logbook vector containing all penalty informationdelay\_penalties: a vetor with the delay penalty and delay outlier penaltycorrect: a matrix of the correct splitswrong: a matrix of the incorrect splits

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

logbook: The logbook vector containing all penalty informationHL\_penalties: a vetor with the half-life penalty and half-life outlier penaltycorrect: a matrix of the correct splitswrong: a matrix of the incorrect splits

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

logbook: The logbook vector containing all penalty informationinty\_penalties: a vetor with the intensity penalty and intensity outlier penaltycorrect: a matrix of the correct splitswrong: a matrix of the incorrect splits

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

**logbook:** The logbook vector containing all penalty information **TI\_penalties:** a vetor with the TI penalty and TI outlier penalty **correct:** a matrix of the correct splits **wrong:** a matrix of the incorrect splits

## Source

https://github.com/CyanolabFreiburg/rifi

penalties_minimal	The result of rifi_penalties for artificial example data A Summarized-
	Experiment containing the output from rifi_penalties including the log-
	book and the four penalty objects as metadata.

## Description

The result of rifi\_penalties for artificial example data A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata.

### Usage

```
data(penalties_minimal)
```

#### Format

An object of class RangedSummarizedExperiment with 24 rows and 33 columns.

#### Source

https://github.com/CyanolabFreiburg/rifi

penalties\_synechocystis\_6803

The result of rifi\_penalties for Synechocystis 6803 example data. A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata.

## Description

The result of rifi\_penalties for Synechocystis 6803 example data. A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata.

## Usage

data(penalties\_synechocystis\_6803)

#### Format

A list with 5 items:

logbook: The logbook vector containing all penalty information

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

**logbook:** The logbook vector containing all penalty information **delay\_penalties:** a vetor with the delay penalty and delay outlier penalty **correct:** a matrix of the correct splits **wrong:** a matrix of the incorrect splits

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

**logbook:** The logbook vector containing all 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

**pen\_obj\_inty:** A list with 4 items:

logbook: The logbook vector containing all penalty information
inty\_penalties: a vetor with the intensity penalty and intensity outlier penalty
correct: a matrix of the correct splits
wrong: a matrix of the incorrect splits

**pen\_obj\_TI:** A list with 4 items:

**logbook:** The logbook vector containing all penalty information **TI\_penalties:** a vetor with the TI penalty and TI outlier penalty **correct:** a matrix of the correct splits **wrong:** a matrix of the incorrect splits

#### Source

https://github.com/CyanolabFreiburg/rifi

%

predict\_ps\_itss

predict\_ps\_itss

predict\_ps\_itss predicts pausing sites (ps) and internal starting sites (ITSS) between delay fragments.

#### Description

predict\_ps\_itss predicts ps and ITSS within the same TU. Neighboring delay segments are compared to each other by positioning the intercept of the second segment into the first segment using slope and intercept coefficients.

### Usage

predict\_ps\_itss(inp, maxDis = 300)

#### Arguments

inp	SummarizedExperiment: the input data frame with correct format.
maxDis	integer: the maximal distance allowed between two successive fragments

## Details

predict\_ps\_itss uses 3 steps to identify ps and ITSS:

- 1. select unique TU.
- 2. select from the input dataframe the columns: ID, position, strand, delay. delay fragment, TU and slope coordinates, velocity\_fragment and intercept.
- 3. select delay segments in the TU.
- 4. loop into all delay segments and estimate the coordinates of the last point of the first segment using the coefficients of the second segment and vice versa. We get two predicted positions, the difference between them is compared to the threshold.

In case the strand is "-", additional steps are added:

The positions of both segments are ordered from the last position to the first one.

All positions are merged in one column and subtracted from the maximum position. the column is split in 2. The first and second correspond to the positions of the first and second segments respectively.

Both segments are subjected to lm fit and the positions predicted are used on the same way as the opposite strand.

If the difference between the positions predicted is lower than negative threshold, ps is assigned otherwise, and if the difference is higher than the positive threshold, ITSS is assigned.

#### Value

The SummarizedExperiment with the columns regarding statistics:

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

position: The bin/probe specific position.

strand: The bin/probe specific strand.

**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: String, the factor of TI fragment.

**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.

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\_duration: Integer, the duration between two delay fragments.

## preprocess\_e\_coli

## Examples

```
data(fragmentation_minimal)
predict_ps_itss(inp = fragmentation_minimal, maxDis = 300)
```

preprocess\_e\_coli The result of rifi\_preprocess for E.coli example data A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata. A list containing the output from rifi\_preprocess, including the inp and the modified input\_df.

## Description

The result of rifi\_preprocess for E.coli example data A SummarizedExperiment containing the output from rifi\_penalties including the logbook and the four penalty objects as metadata. A list containing the output from rifi\_preprocess, including the inp and the modified input\_df.

#### Usage

```
data(preprocess_e_coli)
```

#### Format

A SummarizedExperiment:

inp: the SummarizedExperiment:

ID: The bin/probe specific IDposition: The bin/probe specific positionintensity: The relative intensity at time point 0probe\_TI: An internal value to determine which fitting model is appliedflag: Information on which fitting model is appliedpostion\_segment: The position based segment

fit\_obj\_TI: the fit object for the TI fit:

- **0:** relative intensities at 0 min
- 1: relative intensities at 1 min
- 10: relative intensities at 10 min

15: relative intensities at 15 min

2: relative intensities at 2 min

- 20: relative intensities at 20 min
- 3: relative intensities at 3 min
- 4: relative intensities at 4 min
- **5:** relative intensities at 5 min
- 6: relative intensities at 6 min
- 8: relative intensities at 8 min

**ID:** The bin/probe specific ID**position:** The bin/probe specific position**filtration:** indicator wether the replicate is filtered or not

#### Source

https://github.com/CyanolabFreiburg/rifi

preprocess_minimal	The result of rifi_preprocess for artificial example data A Summarized-
	Experiment containing the output from rifi_preprocess

### Description

The result of rifi\_preprocess for artificial example data A SummarizedExperiment containing the output from rifi\_preprocess

#### Usage

data(preprocess\_minimal)

#### Format

An object of class RangedSummarizedExperiment with 4 rows and 33 columns.

### Source

https://github.com/CyanolabFreiburg/rifi

preprocess\_synechocystis\_6803

The result of rifi\_preprocess for Synechocystis 6803 example data is a A SummarizedExperiment containing the output of rifi\_preprocess as an extention to rowRanges

## Description

The result of rifi\_preprocess for Synechocystis 6803 example data is a A SummarizedExperiment containing the output of rifi\_preprocess as an extention to rowRanges

### Usage

data(preprocess\_synechocystis\_6803)

res\_minimal

## Format

A SummarizedExperiment:

inp: the SummarizedExperiment:

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

position: The bin/probe specific position

strand: The bin/probe specific strand

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

postion\_segment: The position based segment

fit\_obj\_TI: the fit object for the TI fit:

- 0: relative intensities at 0 min
- 2: relative intensities at 2 min
- 4: relative intensities at 4 min
- 8: relative intensities at 8 min
- 16: relative intensities at 16 min
- 32: relative intensities at 32 min
- 64: relative intensities at 64 min
- **ID:** The bin/probe specific ID

position: The bin/probe specific position

filtration: indicator wether the replicate is filtered or not

## Source

https://github.com/CyanolabFreiburg/rifi

res\_minimal

The result of event\_dataframe for E.coli artificial example. A data frame combining the processed genome annotation and a SummarizedExperiment data from rifi\_stats. The dataframe is

## Description

The result of event\_dataframe for E.coli artificial example. A data frame combining the processed genome annotation and a SummarizedExperiment data from rifi\_stats. The dataframe is

#### Usage

data(res\_minimal)

### Format

A list with 2 items:

region: the region from the gff file

gene: String, gene annotation covering the fragments

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

strand: the strand of the annotation

TU: The overarching transcription unit

position: The bin/probe specific position

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

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

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

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\_HL\_intensity\_fragment: fragments involved on ratio of fold change between 2 half-life fragments and fold change between 2 intensity fragments
- **FC\_HL\_intensity:** 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
- **p\_value\_Manova:** p\_value of the variance between two fold-changes, HL and intensity

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

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

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

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

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

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

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

event\_duration: Integer, the duration between two delay fragments

delay\_frg\_slope: the slope value of the respective delay fragment

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

delay: The delay value of the bin/probe

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

intensity: The relative intensity at time point 0

#### Source

https://github.com/CyanolabFreiburg/rifi

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## Description

rifi\_fit wraps the functions:

- 1. nls2\_fit
- 2. TI\_fit
- 3. plot\_nls2\_function
- 4. plot\_singleProbe\_function

## Usage

```
rifi_fit(
  inp,
  cores = 1,
 viz = FALSE,
 restr = 0.2,
 decay = seq(0.08, 0.11, by = 0.02),
 delay = seq(0, 10, by = 0.1),
 k = seq(0.1, 1, 0.2),
 bg = 0.2,
 TI_k = seq(0, 1, by = 0.5),
 TI_decay = c(0.05, 0.1, 0.2, 0.5, 0.6),
 TI = seq(0, 1, by = 0.5),
 TI_delay = seq(0, 2, by = 0.5),
 TI\_rest\_delay = seq(0, 2, by = 0.5),
 TI_bg = 0
)
```

## Arguments

inp	SummarizedExperiment: the input with correct format.
cores	integer: the number of assigned cores for the task.
viz	logical: whether to visualize the output.
restr	numeric: a parameter that restricts the freedom of the fit to avoid wrong TI-term_factors, ranges from 0 to $0.2$
decay	numeric vector: A sequence of starting values for the decay. Default is seq(.08, 0.11, by=.02)

numeric vector: A sequence of starting values for the delay. Default is seq(0,10, by=.1)
numeric vector: A sequence of starting values for the synthesis rate. Default is $seq(0.1,\!1,\!0.2)$
numeric vector: A sequence of starting values. Default is 0.2.
numeric vector: A sequence of starting values for the synthesis rate. Default is $seq(0, 1, by = 0.5)$ .
numeric vector: A sequence of starting values for the decay. Default is $c(0.05, 0.1, 0.2, 0.5, 0.6)$ .
numeric vector: A sequence of starting values for the TI. Default is seq $(0, 1, by = 0.5)$ .
numeric vector: A sequence of starting values for the delay. Default is seq $(0, 2, by = 0.5)$ .
numeric vector: A sequence of starting values. Default is $seq(0, 2, by = 0.5)$ .
numeric vector: A sequence of starting values. Default is 0.

## Value

the SummarizedExperiment object: with delay, decay and TI\_termination\_factor added to the rowRanges. The full fit data is saved in the metadata as "fit\_STD" and "fit\_TI". A plot is given if viz = TRUE.

## See Also

nls2\_fit TI\_fit plot\_nls2 plot\_singleProbe

## Examples

```
data(preprocess_minimal)
rifi_fit(
    inp = preprocess_minimal,
    cores = 1, viz = FALSE, restr = 0.1,
    decay = seq(.08, 0.11, by = .02),
    delay = seq(0, 10, by = .1), k = seq(0.1, 1, 0.2), bg = 0.2,
    TI_k = seq(0, 1, by = 0.5), TI_decay = c(0.05, 0.1, 0.2, 0.5, 0.6),
    TI = seq(0, 1, by = 0.5), TI_delay = seq(0, 2, by = 0.5),
    TI_rest_delay = seq(0, 2, by = 0.5), TI_bg = 0
)
```

rifi\_fragmentation %

## rifi\_fragmentation

rifi\_fragmentation wraps conveniently all fragmentation steps

## Description

rifi\_fragmentation is wrapper of the following functions:

==

- 1. fragment\_delay
- 2. fragment\_HL
- 3. fragment\_inty
- 4. TUgether
- 5. fragment\_TI

## Usage

```
rifi_fragmentation(
    inp,
    cores = 1,
    pen_delay = NULL,
    pen_out_delay = NULL,
    pen_HL = NULL,
    pen_out_HL = NULL,
    pen_inty = NULL,
    pen_out_inty = NULL,
    pen_TU = NULL,
    pen_TI = NULL,
    pen_out_TI = NULL
)
```

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.	
cores	integer: the number of assigned cores for the task.	
pen_delay	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.	
pen_out_delay	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.	
pen_HL	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.	

pen_out_HL	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.
pen_inty	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
<pre>pen_out_inty</pre>	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.
pen_TU	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default -0.75.
pen_TI	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Default is the auto generated value.
pen_out_TI	numeric: an internal parameter for the dynamic programming. Higher values result in fewer allowed outliers. Default is the auto generated value.

### Value

the SummarizedExperiment object: with delay\_fragment, HL\_fragment, intensity\_fragment, TI\_termination\_fragment and TU, and the respective values added to the rowRanges.

## See Also

fragment\_delay
fragment\_HL
fragment\_inty
TUgether
fragment\_TI

## Examples

```
data(penalties_minimal)
rifi_fragmentation(inp = penalties_minimal, cores = 2)
```

%

=

rifi\_penalties

rifi\_penalties

rifi\_penalties wraps conveniently all penalty steps

## Description

rifi\_penalties wraps the functions:

- 1. make\_pen,
- 2. viz\_pen\_obj

rifi\_penalties

## Usage

```
rifi_penalties(
  inp,
 details = FALSE,
 viz = FALSE,
 top_i = 25,
  cores = 1,
 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 = 4.5,
 rez_pen_out = 9
)
```

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
details	logical: whether to return the penalty objects or just the logbook.
viz	logical: whether to visualize the output or not. Default is FALSE
top_i	integer: the number of top results visualized. Default is all.
cores	integer: the number of assigned cores for the task.
dpt	integer: the number of times a full iteration cycle is repeated with a more narrow range based on the previous cycle. Default is 2.
smpl_min	integer: the smaller end of the sampling size. Default is 10.
<pre>smpl_max</pre>	integer: the larger end of the sampling size. Default is 100.
sta_pen	numeric: the lower starting penalty. Default is 0.5.
end_pen	numeric: the higher starting penalty. Default is 4.5.
rez_pen	numeric: the number of penalties iterated within the penalty range. Default is 9.
sta_pen_out	numeric: the lower starting outlier penalty. Default is 0.5.
end_pen_out	numeric: the higher starting outlier penalty. Default is 3.5.
rez_pen_out	numeric: the number of outlier penalties iterated within the outlier penalty range. Default is 7.

#### Value

The SummarizedExperiment object: with the penalties in the logbook added to the metadata. Also adds logbook\_details if details is TRUE, and plots the penalties if viz is TRUE.

## See Also

make\_pen
viz\_pen\_obj

## Examples

```
data(fit_minimal)
rifi_penalties(
    inp = fit_minimal, details = FALSE, viz = FALSE,
    top_i = 25, cores = 2, 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 = 4.5, rez_pen_out = 9
)
```

rifi\_preprocess

#### 

%

rifi\_preprocess wraps conveniently all pre-processing steps

## Description

rifi\_preprocess wraps the functions:

- 1. check\_input
- 2. make\_df
- 3. function\_seg
- 4. finding\_PDD
- 5. finding\_TI

## Usage

```
rifi_preprocess(
  inp,
  cores,
 FUN_filter = function(x) {
                                  FALSE },
 bg = 0,
  rm_FLT = FALSE,
  thrsh_check = 0,
  dista = 300,
  run_PDD = FALSE,
  pen_PDD = 2,
  pen_out_PDD = 1,
  thrsh_PDD = 0.001,
  pen_TI = 10,
  thrsh_{TI} = 0.5,
  add = 1000
)
```

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## rifi\_preprocess

## Arguments

inp	SummarizedExperiment: the input.
cores	integer: the number of assigned cores for the task.
FUN_filter	function: A function of x, returning a logical. x is the numeric vector of the intensity from all time points for a specific replicate.
bg	numeric: threshold over which the last time point has to be to be fitted with the above background mode.
rm_FLT	logical: remove IDs where all replicates are marked as filtered by the back- ground check. Default is FALSE.
thrsh_check	numeric: the minimal allowed intensity for time point "0". Advised to be kept at 0! Default is 0.
dista	integer: the amount of nucleotides defining the gap. Default is 300.
run_PDD	logical: running the PDD flag function
pen_PDD	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Advised to be kept at 2. Default is 2.
pen_out_PDD	numeric: an internal parameter for the dynamic programming. Higher values result in fewer possible outliers. Advised to be kept at 1. Default is 1.
thrsh_PDD	numeric: an internal parameter that allows fragments with slopes steeper than the threshold to be flagged with " <i>PDD</i> ". Higher values result in fewer candidates . Advised to be kept at 0.001. Default is 0.001.
pen_TI	numeric: an internal parameter for the dynamic programming. Higher values result in fewer fragments. Advised to be kept at 10. Default is 10.
thrsh_TI	numeric: an internal parameter that allows fragments with a certain amount of IDs with higher relative intensities at time points later than "0" to be flagged as " <i>TI</i> ". Higher values result in fewer candidates0.5 is 25 %, 0 is 50%, 0.5 is 75%. Advised to be kept at 0.5. Default is 0.5.
add	integer: range of nucleotides before a potential TI event where in IDs are fitted with the TI fit.

## Details

rifi\_preprocess allows for the optional integration of filter functions. Filter functions mark replicates with TRUE. Those are then not considered in the fit! FUN\_filter is a general filter usually to exclude probes with low expression or "bad" patterns.

## Value

The SummarizedExperiment object: checked, and with position, ID, intensity, probe\_TI, position\_segment, flag and filtration added to the rowRanges.

## See Also

check\_input make\_df

```
segment_pos
finding_PDD
finding_TI
```

## Examples

```
data(example_input_minimal)
rifi_preprocess(
    inp = example_input_minimal, cores = 2, bg = 100, rm_FLT = FALSE,
    thrsh_check = 0, dista = 300, run_PDD = FALSE
    )
```

rifi\_stats

## rifi\_stats wraps all statistical prediction steps conveniently

## Description

rifi\_stats wraps the functions:

- 1. predict\_ps\_itss
- 2. apply\_Ttest\_delay
- 3. apply\_ancova
- 4. apply\_event\_position
- 5. apply\_t\_test
- 6. fold\_change
- 7. apply\_manova
- 8. apply\_t\_test\_ti
- 9. gff3\_preprocess

## Usage

rifi\_stats(inp, dista = 300, path)

## Arguments

inp	SummarizedExperiment: the input data frame with correct format.
dista	integer: the maximal distance allowed between two successive fragments. De- fault is the auto generated value.
path	path: to the directory containing the gff3 file.

#### rifi\_summary

## Value

The SummarizedExperiment object: 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, TI\_fragments\_p\_value

### See Also

```
predict_ps_itss
apply_Ttest_delay
apply_ancova
apply_event_position
apply_t_test
fold_change
apply_manova
apply_t_test_ti
gff3_preprocess
```

#### Examples

```
data(fragmentation_minimal)
rifi_stats(inp = fragmentation_minimal, dista = 300,
path = gzfile(system.file("extdata", "gff_e_coli.gff3.gz",
package = "rifi")))
```

rifi\_summary

## %

## rifi\_summary

rifi\_summary wraps conveniently and summarize all rifi outputs

#### Description

rifi\_summary wraps the functions:

- 1. event\_dataframe
- 2. dataframe\_summary
- 3. dataframe\_summary\_events

- 4. dataframe\_summary\_events\_HL\_int
- 5. dataframe\_summary\_events\_ps\_itss
- 6. dataframe\_summary\_events\_velocity
- 7. dataframe\_summary\_TI

## Usage

```
rifi_summary(inp, data_annotation = metadata(inp)$annot[[1]])
```

## Arguments

```
inp SummarizedExperiment: the input data frame with correct format.
```

data\_annotation

dataframe: gff3 dataframe after processing.

## Value

WIP

## See Also

event\_dataframe
dataframe\_summary
dataframe\_summary\_events
dataframe\_summary\_events\_HL\_int
dataframe\_summary\_events\_ps\_itss
dataframe\_summary\_events\_velocity
dataframe\_summary\_TI

### Examples

```
data(stats_minimal)
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
rifi_summary(inp = stats_minimal, data_annotation =
metadata(stats_minimal)$annot[[1]])
```

rifi\_visualization %

#### rifi visualization

rifi\_visualization plots all the data with fragments and events from both strands

## Description

rifi\_visualization plots the whole genome with genes, transcription units (TUs), delay, half-life (HL), intensity fragments, features, events, velocity, annotation, coverage if available.

### Usage

```
rifi_visualization(
  data,
  genomeLength,
  annot,
  coverage = 0,
  chr_fwd = NA,
  chr_rev = NA,
  region = c("CDS", "asRNA", "5'UTR", "ncRNA", "3'UTR", "tRNA"),
 color_region = c("grey0", "red", "blue", "orange", "yellow", "green", "white",
    "darkseagreen1", "grey50", "black"),
  color_text.1 = "grey0",
  color_text.2 = "black",
  color_TU = "blue",
  Alpha = 0.5,
  size_tu = 1.6,
  size_locusTag = 1.6,
  size_gene = 1.6,
  Limit = 10,
  shape = 22,
  col_outiler = "grey50",
  col_coverage = "grey",
  shape_outlier = 13,
  limit_intensity = NA,
  face = "bold",
  tick_length = 0.3,
  arrow.color = "darkseagreen1",
 minVelocity = 3000,
 medianVelocity = 6000,
  col_above20 = "#00FFFF",
  fontface = "plain",
  shape_above20 = 14,
```

```
col_outlierabove10 = "darkorchid",
  shape_outlierabove10 = 5,
 axis_text_y_size = 3,
 axis_title_y_size = 6,
 TI_threshold = 1.1,
  termination_threshold = -0.5,
 iTSS_threshold = 0.5,
 p_value_int = 0.05,
 p_value_event = 0.05,
 p_value_hl = 0.05,
 p_value_TI = 0.05,
 p_value_manova = 0.05,
 event_duration_ps = 1,
 event_duration_itss = -1,
 HL_threshold_1 = log2(1.5),
 HL_threshold_2 = -log2(1.5),
 vel_threshold = 200,
 HL_threshold_color = "black",
 vel_threshold_color = "grey52",
 ps_color = "orange",
 iTSS_I_color = "blue"
)
```

### Arguments

data	SummarizedExperiment: the input data frame with correct format.
genomeLength	integer: genome length output of gff3_preprocess function and element of meta- data of SummarizedExperiment.
annot	dataframe: the annotation file, output of gff3_preprocess function and element of metadata of SummarizedExperiment.
coverage	integer: in case the coverage is available.
chr_fwd	string object: coverage of the forward strand.
chr_rev	string object: coverage of the reverse strand.
region	dataframe: gff3 features of the genome.
color_region	string vector: vector of colors.
color_text.1	string: TU color text
color_text.2	string: genes color text
color_TU	string. TU color
Alpha	integer: color transparency degree.
size_tu	integer: TU size
<pre>size_locusTag</pre>	integer: locus_tag size
size_gene	integer: font size for gene annotation.
Limit	integer: value for y-axis limit.
shape	integer: value for shape.

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col_outiler	string: outlier color.
col_coverage	integer: color for coverage plot.
shape_outlier	integer: value for outlier shape.
limit_intensity	y
	integer: intensity limit if applicable.
face	string: label font.
tick_length	integer: value for ticks.
arrow.color	string: arrows color.
minVelocity	integer: threshold to fix the minimum of velocity.
medianVelocity	integer: threshold to fix the maximum of velocity.
col_above20	string: color for probes/bin above value 20.
fontface	integer: font type
shape_above20	integer: shape for probes/bins above value 20.
col_outlierabov	
-h	string: color for probes/bin outliers between 10 and 20,
shape_outlieral	integer: shape for probes/bin outliers between 10 and 20,
axis_text_y_siz	
	integer: text size for y-axis.
axis_title_y_s:	
	integer: title size for y-axis.
TI_threshold	integer: threshold for TI between two fragments in case the TI termination factor drops from the first segment to the second, default 1.1. If threshold is reached a line is drawn to seperates the two TI segments.
termination_th	
	integer: threshold for termination to plot, default .8.
iTSS_threshold	integer: threshold for iTSS_II selected to plot, default 1.2.
p_value_int	integer: p_value of intensity fragments fold-change to plot, default 0.05.
p_value_event	integer: p_value of t-test from pausing site and iTSS_I events to plot, default 0.05.
p_value_hl	integer: p_value of half_life fragments fold-change to plot, default 0.05.
p_value_TI	integer: p_value of TI fragments selected to be plotted, default 0.05.
p_value_manova	integer: p_value of manova test fragments to plot, default 0.05.
event_duration	
	integer: threshold for pausing sites selected to plot, default -2.
event_duration_	integer: threshold for iTSS_I selected to plot, default 2.
HL_threshold_1	integer: threshold for log2FC(HL) selected to plot, default log2(1.5). log2FC(HL)
	>= log2(1.5) are indicated by black color. If p_value <= p_value_hl (default 0.05), log2FC(HL) is indicated by HL* otherwise HL.

HL_threshold_2	integer: threshold for $\log 2FC(HL)$ selected to plot, default $-\log 2(1.5)$ . $\log 2FC(HL)$	
	<= -log2(1.5) are indicated by green color. If p_value <= p_value_hl (default	
	0.05), log2FC(HL) is indicated by HL* otherwise HL. In case of p_value is sig-	
	nificant and the log2FC(HL) is between $-\log_2FC(1.5)$ and $\log_2FC(1.5)$ , FC is assigned by green color and HL*.	
vel_threshold	integer: threshold for velocity ratio selected to plot, default 200.	
HL_threshold_color		
	string: color for HL fold change plot.	
vel_threshold_c	olor	
	string: color for velocity ratio plot.	
ps_color	string: color for pausing site plot.	
iTSS_I_color	string: color for iTSS_I plot.	

#### Details

rifi\_visualization uses several functions to plot the genes including as-RNA and ncRNA and TUs as segments. The function plots delay, HL and intensity fragments with statistical t-test between the neighboring fragment, significant t-test is assigned with '. *t-test and Manova statistical test are also depicted as* ''.

The functions used are:

- 1. annotation\_plot: plots the corresponding annotation.
- 2. positive\_strand\_function: plots delay, HL, intensity and events of positive strand.
- 3. negative\_strand\_function: plots delay, HL, intensity and events of negative strand.
- 4. empty\_data\_positive: plots empty boxes in case no data is available for positive strand.
- 5. empty\_data\_negative: plots empty boxes in case no data is available for negative strand.
- 6. strand\_selection: check if data is stranded and arrange by position.
- 7. splitGenome\_function: splits the genome into fragments.
- 8. indice\_function: assign a new column to the data to distinguish between fragments, outliers from delay or HL or intensity.
- 9. TU\_annotation: designs the segments border for the genes and TUs annotation
- 10. 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.
- 11. label\_log2\_function: used to add log scale to intensity values.
- 12. label\_square\_function: used to add square scale to coverage values.
- 13. coverage\_function: this function is used only in case of coverage is available.
- 14. 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.
- 15. outlier\_plot: plot the outliers with half-life between 10 and 30 on the maximum of the yaxis.
- 16. add\_genomeBorders: when the annotated genes are on the borders, they can not be plotted, therefore the region was split in 2 adding the row corresponding to the split part to the next annotation (i + 1) except for the first page.

- 17. my\_arrow: creates an arrow for the annotation.
- 18. arrange\_byGroup: selects the last row for each segment and add 40 nucleotides in case of negative strand for a nice plot.
- 19. regr: plots the predicted delay from linear regression if the data is on negative strand.
- 20. meanPosition: assign a mean position for the plot.
- 21. delay\_mean: adds a column in case of velocity is NA or equal to 60. The mean of the delay is calculated outliers.
- 22. my\_segment\_T: plots terminals and pausing sites labels.
- 23. my\_segment\_NS: plots internal starting sites 'iTSS'.
- 24. min\_value: returns minimum value for event plots in intensity plot.
- 25. velocity\_fun: function for velocity plot.
- 26. limit\_function: for values above 10 or 20 in delay and hl. Limit of the axis is set differently. y-axis limit is applied only if we have more than 3 values above 10 and lower or equal to 20. An exception is added in case a dataframe has less than 3 rows and 1 or more values are above 10, the rest of the values above 20 are adjusted to 20 on "secondaryAxis" function.
- 27. empty\_boxes: used only in case the dataframe from the positive strand is not empty, the TU are annotated.
- 28. function\_TU\_arrow: used to avoid plotting arrows when a TU is split into two pages.
- 29. terminal\_plot\_lm: draws a linear regression line when terminal outliers have an intensity above a certain threshold and are consecutive. Usually are smallRNA (ncRNA, asRNA).
- 30. slope\_function: replaces slope lower than 0.0009 to 0.
- 31. velo\_function: replaces infinite velocity with NA.
- 32. plot the coverage of RNA\_seq in exponential phase growth

#### Value

The visualization plot.

#### Examples

```
data(stats_minimal)
if(!require(SummarizedExperiment)){
suppressPackageStartupMessages(library(SummarizedExperiment))
}
rifi_visualization(data = stats_minimal,
genomeLength = metadata(stats_minimal)$annot[[2]],
annot = metadata(stats_minimal)$annot[[1]])
```

rifi_wrapper	%
	======================================
	rifi_wrapper wraps conveniently all functions included on rifi workflow

# Description

rifi\_wrapper wraps the functions:

- 1. rifi\_preprocess
- 2. rifi\_fit
- 3. rifi\_penalties
- 4. rifi\_fragmentation
- 5. rifi\_stats
- 6. rifi\_summary
- 7. rifi\_visualization.

## Usage

rifi\_wrapper(inp, cores, path, bg, restr)

# Arguments

inp	data frame: the input data frame with correct format.
cores	integer: the number of assigned cores for the task.
path	path: path to an annotation file in gff format.
bg	numeric: threshold over which the last time point has to be to be fitted with the above background mode.
restr	numeric: a parameter that restricts the freedom of the fit to avoid wrong TI-term_factors, ranges from 0 to $0.2$

## Value

All intermediate objects

segment\_pos

## See Also

```
rifi_preprocess
rifi_fit
rifi_penalties
rifi_fragmentation
rifi_stats
rifi_summary
rifi_visualization
```

# Examples

```
data(example_input_minimal)
rifi_wrapper(inp = example_input_minimal, cores = 2, path =
gzfile(system.file("extdata", "gff_e_coli.gff3.gz", package = "rifi")),
bg = 0, restr = 0.01)
```

<pre>segment_pos</pre>	%
	======================================
	segment_pos divides all IDs by position into position_segments

# Description

segment\_pos adds the column "position\_segment" to the rowRanges. To reduce run time, the data is divided by regions of no expression larger than "dist" nucleotides.

## Usage

```
segment_pos(inp, dista = 300)
```

# Arguments

inp	SummarizedExperiment: the input.
dista	integer: the amount of nucleotides defining the gap. Default is 300.

# Value

The SummarizedExperiment object:

ID:	The bin/probe specific ID
position:	The bin/probe specific position
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
<pre>position_segment:</pre>	
	The position based segment

# Examples

```
data(preprocess_minimal)
segment_pos(inp = preprocess_minimal, dista = 300)
```

stats_e_coli	The result of rifi_stats for E.coli example data A SummarizedExperi-
	ment containing the output from rifi_stats

# Description

The result of rifi\_stats for E.coli example data A SummarizedExperiment containing the output from rifi\_stats

# Usage

data(stats\_e\_coli)

## Format

A SummarizedExperiment:

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

position: The bin/probe specific position

strand: The bin/probe specific strand

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: String, the factor of TI fragment

**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 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

#### Source

https://github.com/CyanolabFreiburg/rifi

stats\_minimal

The result of rifi\_stats for artificial example data A SummarizedExperiment containing the output of rifi\_stats as an extention to rowRanges and metadata (gff file processed, see gff file documentation)

#### Description

The result of rifi\_stats for artificial example data A SummarizedExperiment containing the output of rifi\_stats as an extention to rowRanges and metadata (gff file processed, see gff file documentation)

## Usage

data(stats\_minimal)

## Format

A rowRanges of SummarizedExperiment with 24 rows and 45 variables:

ID: The bin/probe specific ID position: The bin/probe specific position 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: String, the factor of TI fragment 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 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

#### Source

https://github.com/CyanolabFreiburg/rifi

stats\_synechocystis\_6803

The result of rifi\_stats for Synechocystis 6803 example data A SummarizedExperiment containing the output of rifi\_stats as an extention to rowRanges

#### Description

The result of rifi\_stats for Synechocystis 6803 example data A SummarizedExperiment containing the output of rifi\_stats as an extention to rowRanges

#### Usage

data(stats\_synechocystis\_6803)

#### Format

The rowRanges of SummarizedExperiment:

**ID:** The bin/probe specific ID position: The bin/probe specific position 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: String, the factor of TI fragment 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 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

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- 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/rifi

summary_e_coli	The result of rifi_summary for E.coli example data A Summarized-
	Experiment containing the output of rifi_stats as an extention to rowRanges

## Description

The result of rifi\_summary for E.coli example data A SummarizedExperiment containing the output of rifi\_stats as an extention to rowRanges

#### Usage

data(summary\_e\_coli)

#### Format

The rowRanges of SummarizedExperiment:

**bin\_df:** all information regarding bins:

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

**feature\_type:** String, region annotation covering the fragments **gene:** String, gene annotation covering the fragments

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

position: The bin/probe specific position strand: The bin/probe specific strand segment: The segment the bin/probe belongs to TU: The overarching transcription unit delay\_fragment: The delay fragment the bin/probe belongs to **delay:** The delay of the bin/probe HL fragment: The half-life fragment the bin/probe belongs to half\_life: The half-life of the bin/probe intensity fragment: The intensity fragment the bin/probe belongs to **intensity:** The relative intensity at time point 0 flag: The flag of the bin/probe(TI, PDD) TI\_termination\_factor: String, the factor of TI fragment frag\_df: all information regarding fragments: feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments first position frg: The first position of the fragment on the genome last\_position\_frg: The last position of the fragment on the genome strand: The bin/probe specific strand TU: The overarching transcription unit segment: The segment the fragment belongs to delay fragment: The delay fragment of the fragment HL\_fragment: The half-life fragment of the fragment half life: The half-life mean of the fragment HL\_SD: The half-life standard deviation of the fragment HL SE: The half-life standard error of the fragment intensity\_fragment: The intensity\_fragment of the fragment intensity: The relative intensity at time point 0 intensity\_SD: The intensity standard deviation of the fragment intensity\_SE: The intensity standard error of the fragment velocity: The velocity value of the respective delay fragment event\_df: all information regarding events: event: String, event type **p\_value:** Integer, p\_value of the event **p\_adjusted:** Integer, p\_value adjusted FC HL: Integer, the fold change value of 2 HL fragments FC\_intensity: Fold change of intensity 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 FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity event\_position: Integer, the position middle between 2 fragments with an event velocity\_ratio: Integer, ratio of velocity between 2 delay fragments

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feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment 1: String, the first fragment of the two of fragments subjected to analysis segment\_2: String, the second fragment of the two of fragments subjected to analysis event duration: Integer, the duration between two delay fragments gap fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event events\_HL\_int\_df: all information regarding events related to half-life and intensity: event: String, event type **p\_value:** Integer, p\_value of the event **p\_adjusted:** Integer, p\_value adjusted FC\_HL: Integer, the fold change value of 2 HL fragments FC intensity: Integer, the fold change value of 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 FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity event\_position: Integer, the position middle between 2 fragments with an event feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment 1: String, the first fragment of the two of fragments subjected to analysis segment\_2: String, the second fragment of the two of fragments subjected to analysis event duration: Integer, the duration between two delay fragments gap\_fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event events ps itss df: all information regarding events related to pausing sites and iTSS I: event: String, event type **p\_value:** Integer, **p\_value** of the event p\_adjusted: Integer, p\_value adjusted event\_position: Integer, the position middle between 2 fragments with an event velocity ratio: Integer, ratio of velocity between 2 delay 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 feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand

**TU:** The overarching transcription unit segment 1: String, the first fragment of the two of fragments subjected to analysis segment\_2: String, the second fragment of the two of fragments subjected to analysis event\_duration: Integer, the duration between two delay fragments gap\_fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event events velocity df: all information regarding events related to velocity: event: String, event type **p** value: Integer, p value of the event p\_adjusted: Integer, p\_value adjusted event\_position: Integer, the position middle between 2 fragments with an event velocity\_ratio: Integer, ratio of velocity between 2 delay fragments feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment\_1: String, the first fragment of the two of fragments subjected to analysis segment\_2: String, the second fragment of the two of fragments subjected to analysis event duration: Integer, the duration between two delay fragments gap\_fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event TI\_df: all information regarding TI: event: String, event type TI\_fragment: String, the fragment with TI TI\_termination\_factor: String, the factor of TI fragment **p\_value:** Integer, p\_value of the event **p\_adjusted:** Integer, p\_value adjusted feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit features: Integer, number of fragements involved on the event event\_position: Integer, the position middle between 2 fragments with an event **position 1:** the first position of TI fragment, if 2 fragments, first position is from the first fragment position\_2: the last position of TI fragment, if 2 fragments, last position is from the second

#### Source

https://github.com/CyanolabFreiburg/rifi

fragment.

summary\_minimal

The result of rifi\_summary for artificial example data A Summarized-Experiment with the output from rifi\_summary as metadata

#### Description

The result of rifi\_summary for artificial example data A SummarizedExperiment with the output from rifi\_summary as metadata

#### Usage

data(summary\_minimal)

# Format

A list of 7 data frames with 290 rows and 11 variables, 36 rows and 11 variables, 57 rows and 18 variables, and 8 rows and 14 variables:

**bin\_df:** all information regarding bins:

**ID:** The bin/probe specific ID feature type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments position: The bin/probe specific position strand: The bin/probe specific strand segment: The segment the bin/probe belongs to TU: The overarching transcription unit delay\_fragment: The delay fragment the bin/probe belongs to delay: The delay of the bin/probe HL\_fragment: The half-life fragment the bin/probe belongs to half\_life: The half-life of the bin/probe intensity\_fragment: The intensity fragment the bin/probe belongs to intensity: The relative intensity at time point 0 flag: The flag of the bin/probe(TI, PDD) TI\_termination\_factor: String, the factor of TI fragment frag\_df: all information regarding fragments: feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments

first\_position\_frg: The first position of the fragment on the genome

**last\_position\_frg:** The last position of the fragment on the genome

strand: The bin/probe specific strand

TU: The overarching transcription unit

segment: The segment the fragment belongs to

delay\_fragment: The delay fragment of the fragment

HL\_fragment: The half-life fragment of the fragment

half\_life: The half-life mean of the fragment

**HL\_SD:** The half-life standard deviation of the fragment

**HL\_SE:** The half-life standard error of the fragment

intensity\_fragment: The intensity\_fragment of the fragment

intensity: The relative intensity at time point 0

intensity\_SD: The intensity standard deviation of the fragment

intensity\_SE: The intensity standard error of the fragment

velocity: The velocity value of the respective delay fragment

event\_df: all information regarding events:

event: String, event type

**p\_value:** Integer, **p\_value** of the event

p\_adjusted: Integer, p\_value adjusted

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

FC\_intensity: Fold change of intensity

**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

FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity

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

velocity\_ratio: Integer, ratio of velocity between 2 delay fragments

feature\_type: String, region annotation covering the fragments

gene: String, gene annotation covering the fragments

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

strand: The bin/probe specific strand

TU: The overarching transcription unit

segment\_1: String, the first fragment of the two of fragments subjected to analysis

segment\_2: String, the second fragment of the two of fragments subjected to analysis

event\_duration: Integer, the duration between two delay fragments

gap\_fragments: Integer, the distance between two delay fragments

features: Integer, number of fragements involved on the event

events\_HL\_int\_df: all information regarding events related to half-life and intensity:

event: String, event type

p\_value: Integer, p\_value of the event

p\_adjusted: Integer, p\_value adjusted

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

FC\_intensity: Integer, the fold change value of 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

FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity

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

feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment\_1: String, the first fragment of the two of fragments subjected to analysis segment 2: String, the second fragment of the two of fragments subjected to analysis event\_duration: Integer, the duration between two delay fragments gap fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event events ps itss df: all information regarding events related to pausing sites and iTSS I: event: String, event type **p\_value:** Integer, p\_value of the event p\_adjusted: Integer, p\_value adjusted event\_position: Integer, the position middle between 2 fragments with an event velocity\_ratio: Integer, ratio of velocity between 2 delay 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 feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment\_1: String, the first fragment of the two of fragments subjected to analysis segment 2: String, the second fragment of the two of fragments subjected to analysis event\_duration: Integer, the duration between two delay fragments gap\_fragments: Integer, the distance between two delay fragments features: Integer, number of fragements involved on the event events\_velocity\_df: all information regarding events related to velocity: event: String, event type **p** value: Integer, p value of the event p\_adjusted: Integer, p\_value adjusted event\_position: Integer, the position middle between 2 fragments with an event velocity\_ratio: Integer, ratio of velocity between 2 delay fragments feature type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments strand: The bin/probe specific strand TU: The overarching transcription unit segment\_1: String, the first fragment of the two of fragments subjected to analysis segment\_2: String, the second fragment of the two of fragments subjected to analysis event\_duration: Integer, the duration between two delay fragments

**gap\_fragments:** Integer, the distance between two delay fragments **features:** Integer, number of fragements involved on the event

TI\_df: all information regarding TI:

event: String, event type

TI\_fragment: String, the fragment with TI

TI\_termination\_factor: String, the factor of TI fragment

p\_value: Integer, p\_value of the event

p\_adjusted: Integer, p\_value adjusted

feature\_type: String, region annotation covering the fragments

gene: String, gene annotation covering the fragments

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

strand: The bin/probe specific strand

TU: The overarching transcription unit

features: Integer, number of fragements involved on the event

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

- position\_1: the first position of TI fragment, if 2 fragments, first position is from the first
  fragment
- **position\_2:** the last position of TI fragment, if 2 fragments, last position is from the second fragment.

#### Source

https://github.com/CyanolabFreiburg/rifi

summary\_synechocystis\_6803

The result of rifi\_summary for Synechocystis 6803 example data A list containing the output from rifi\_summary, including the fragment based data frame, bin based data frame, event data frame and the TI dataframe.

## Description

The result of rifi\_summary for Synechocystis 6803 example data A list containing the output from rifi\_summary, including the fragment based data frame, bin based data frame, event data frame and the TI dataframe.

#### Usage

data(summary\_synechocystis\_6803)

#### Format

A list of 4 data frames with 3000 rows and 11 variables, 297 rows and 11 variables, 486 rows and 18 variables, and 10 rows and 14 variables:

**bin\_df:** all information regarding bins:

**ID:** The bin/probe specific ID feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments position: The bin/probe specific position strand: The bin/probe specific strand segment: The segment the bin/probe belongs to **TU:** The overarching transcription unit delay fragment: The delay fragment the bin/probe belongs to delay: The delay of the bin/probe **HL fragment:** The half-life fragment the bin/probe belongs to half\_life: The half-life of the bin/probe intensity fragment: The intensity fragment the bin/probe belongs to intensity: The relative intensity at time point 0 flag: The flag of the bin/probe(TI, PDD) TI\_termination\_factor: String, the factor of TI fragment frag\_df: all information regarding fragments: feature\_type: String, region annotation covering the fragments gene: String, gene annotation covering the fragments locus\_tag: String, locus\_tag annotation covering the fragments first\_position\_frg: The first position of the fragment on the genome last\_position\_frg: The last position of the fragment on the genome strand: The bin/probe specific strand TU: The overarching transcription unit segment: The segment the fragment belongs to delay\_fragment: The delay fragment of the fragment **HL\_fragment:** The half-life fragment of the fragment half life: The half-life mean of the fragment

HL SD: The half-life standard deviation of the fragment

HL\_SE: The half-life standard error of the fragment

intensity\_fragment: The intensity\_fragment of the fragment

intensity: The relative intensity at time point 0

intensity\_SD: The intensity standard deviation of the fragment

intensity\_SE: The intensity standard error of the fragment

velocity: The velocity value of the respective delay fragment

event\_df: all information regarding events:

event: String, event type

- **p\_value:** Integer, p\_value of the event
- **p\_adjusted:** Integer, p\_value adjusted
- FC\_HL: Integer, the fold change value of 2 HL fragments
- FC\_intensity: Fold change of intensity
- **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
- FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity
- event\_position: Integer, the position middle between 2 fragments with an event
- velocity\_ratio: Integer, ratio of velocity between 2 delay fragments
- feature\_type: String, region annotation covering the fragments
- gene: String, gene annotation covering the fragments
- locus\_tag: String, locus\_tag annotation covering the fragments
- strand: The bin/probe specific strand
- TU: The overarching transcription unit
- segment\_1: String, the first fragment of the two of fragments subjected to analysis
- segment\_2: String, the second fragment of the two of fragments subjected to analysis
- event\_duration: Integer, the duration between two delay fragments
- gap\_fragments: Integer, the distance between two delay fragments
- features: Integer, number of fragements involved on the event
- events\_HL\_int\_df: all information regarding events related to half-life and intensity:
  - event: String, event type
  - **p\_value:** Integer, p\_value of the event
  - p\_adjusted: Integer, p\_value adjusted
  - FC\_HL: Integer, the fold change value of 2 HL fragments
  - FC\_intensity: Integer, the fold change value of 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
  - FC\_HL\_FC\_intensity: Fold change of half-life/ fold change of intensity
  - event\_position: Integer, the position middle between 2 fragments with an event
  - feature\_type: String, region annotation covering the fragments
  - gene: String, gene annotation covering the fragments
  - locus\_tag: String, locus\_tag annotation covering the fragments
  - strand: The bin/probe specific strand
  - TU: The overarching transcription unit
  - segment\_1: String, the first fragment of the two of fragments subjected to analysis
  - segment 2: String, the second fragment of the two of fragments subjected to analysis
  - event\_duration: Integer, the duration between two delay fragments
  - gap\_fragments: Integer, the distance between two delay fragments
  - features: Integer, number of fragements involved on the event
- events\_ps\_itss\_df: all information regarding events related to pausing sites and iTSS\_I:
  - event: String, event type
  - **p\_value:** Integer, **p\_value** of the event

**p\_adjusted:** Integer, p\_value adjusted

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

velocity\_ratio: Integer, ratio of velocity between 2 delay 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

feature\_type: String, region annotation covering the fragments

gene: String, gene annotation covering the fragments

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

strand: The bin/probe specific strand

TU: The overarching transcription unit

segment\_1: String, the first fragment of the two of fragments subjected to analysis

segment\_2: String, the second fragment of the two of fragments subjected to analysis

event\_duration: Integer, the duration between two delay fragments

gap\_fragments: Integer, the distance between two delay fragments

features: Integer, number of fragements involved on the event

events\_velocity\_df: all information regarding events related to velocity:

event: String, event type

**p\_value:** Integer, p\_value of the event

p\_adjusted: Integer, p\_value adjusted

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

velocity\_ratio: Integer, ratio of velocity between 2 delay fragments

**feature\_type:** String, region annotation covering the fragments

gene: String, gene annotation covering the fragments

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

strand: The bin/probe specific strand

TU: The overarching transcription unit

segment\_1: String, the first fragment of the two of fragments subjected to analysis

segment\_2: String, the second fragment of the two of fragments subjected to analysis

event\_duration: Integer, the duration between two delay fragments

gap\_fragments: Integer, the distance between two delay fragments

features: Integer, number of fragements involved on the event

**TI\_df:** all information regarding TI:

event: String, event type

TI\_fragment: String, the fragment with TI

TI\_termination\_factor: String, the factor of TI fragment

**p\_value:** Integer, **p\_value** of the event

p\_adjusted: Integer, p\_value adjusted

feature\_type: String, region annotation covering the fragments

gene: String, gene annotation covering the fragments

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

strand: The bin/probe specific strand

TU: The overarching transcription unit

features: Integer, number of fragements involved on the event

- event\_position: Integer, the position middle between 2 fragments with an event
- position\_1: the first position of TI fragment, if 2 fragments, first position is from the first
  fragment
- **position\_2:** the last position of TI fragment, if 2 fragments, last position is from the second fragment.

## Source

https://github.com/CyanolabFreiburg/rifi

TI_fit	%
	======================================
	TI_fit estimates transcription interference and termination factor using nls function for probe or bin flagged as "TI".

## Description

TI\_fit uses nls2 function to fit the flagged probes or bins with "TI" found using finding\_TI.r. It estimates the transcription interference level (referred later to TI) as well as the transcription factor fitting the probes/bins with nls function looping into several starting values.

## Usage

```
TI_fit(
    inp,
    cores = 1,
    restr = 0.2,
    k = seq(0, 1, by = 0.5),
    decay = c(0.05, 0.1, 0.2, 0.5, 0.6),
    ti = seq(0, 1, by = 0.5),
    ti_delay = seq(0, 2, by = 0.5),
    rest_delay = seq(0, 2, by = 0.5),
    bg = 0
)
```

#### Arguments

inp	SummarizedExperiment: the input with correct format.
cores	integer: the number of assigned cores for the task.
restr	numeric: a parameter that restricts the freedom of the fit to avoid wrong TI-term_factors, ranges from 0 to 0.2.

k	numeric vector: A sequence of starting values for the synthesis rate. Default is $seq(0, 1, by = 0.5)$ .
decay	numeric vector: A sequence of starting values for the decay Default is $c(0.05, 0.1, 0.2, 0.5, 0.6)$ .
ti	numeric vector: A sequence of starting values for the delay. Default is seq $(0, 1, by = 0.5)$ .
ti_delay	numeric vector: A sequence of starting values for the delay. Default is seq $(0, 2, by = 0.5)$ .
rest_delay	numeric vector: A sequence of starting values. Default is $seq(0, 2, by = 0.5)$ .
bg	numeric vector: A sequence of starting values. Default is 0.

# Details

To determine TI and termination factor, TI\_fit function is applied to the flagged probes and to the probes localized 1000 nucleotides upstream. Before applying TI\_fit function, some probes/bins are filtered out if they are below the background using generic\_filter\_BG. The model loops into a dataframe containing sequences of starting values and the coefficients are extracted from the fit with the lowest residuals. When many residuals are equal to 0, the lowest residual can not be determined and the coefficients extracted could be wrong. Therefore, a second filter was developed. First we loop into all starting values, we collect nls objects and the corresponding residuals. They are sorted and residuals non equal to 0 are collected in a vector. If the first residuals are not equal to 0, 20 % of the best residuals are collected in tmp\_r\_min vector and the minimum termination factor is selected. In case the first residuals are equal to 0 then values between 0 to 20% of the values collected in tmp\_r\_min vector are gathered. The minimum termination factor coefficient is determined and saved. The coefficients are gathered in res vector and saved as an object.

#### Value

the SummarizedExperiment object: with delay, decay and TI\_termination\_factor added to the rowRanges. The full fit data is saved in the metadata as "fit\_TI".

## Examples

```
data(preprocess_minimal)
TI_fit(inp = preprocess_minimal, cores=2, restr=0.01)
```

TUgether

%

## TUgether

TUgether combines delay fragments into TUs

## Description

TUgether combines delay fragments into TUs. The column "TU" is added. It uses score fun\_increasing on the start and end points of delay\_fragments.

#### Usage

```
TUgether(inp, cores = 1, pen = -0.75)
```

#### Arguments

inp	SummarizedExperiment: the input data frame with correct format.
cores	cores: integer: the number of assigned cores for the task.
pen	numeric: an internal parameter for the dynamic programming. Higher values
	result in fewer fragments. Default -0.75.

## Details

The function used is: .score\_fun\_increasing

The input is the SummarizedExperiment object. pen is the penalty for new fragments in the dynamic programming. Since high scores are aimed, pen is negative.

#### Value

The SummarizedExperiment with the columns regarding the TU:

**ID:** The bin/probe specific ID. position: The bin/probe specific position. 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: String, the factor of TI fragment. 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.

viz\_pen\_obj

# Examples

```
data(fragmentation_minimal)
TUgether(inp = fragmentation_minimal, cores = 2, pen = -0.75)
```

viz\_pen\_obj

#### viz\_pen\_obj

======

%

viz\_pen\_obj visualizes penalty objects

# Description

viz\_pen\_obj provides an optional visualization of any penalty object created by make\_pen. the function aan be customized to show only the  $n = top_i$  top results.

# Usage

viz\_pen\_obj(obj, top\_i = nrow(obj[[3]][[1]]) \* ncol(obj[[3]][[1]]))

## Arguments

obj	object: penalty object(make_pen output)
top_i	integer: the number of top results visualized. Default is all.

## Value

A visualization of the penalty object

# Examples

```
data(penalties_e_coli)
viz_pen_obj(penalties_e_coli$pen_obj_delay,25)
```

wrapper\_e\_coli The result of rifi\_wrapper for E.coli example data A list of SummarizedExperiment containing the output of rifi\_wrapper. The list contains 6 elements of SummarizedExperiment output of rifi\_preprocess, rifi\_fit, rifi\_penalties, rifi\_fragmentation, rifi\_stats and rifi\_summary. The plot is generated from rifi\_visualization. for more detail, please refer to each function separately.

## Description

The result of rifi\_wrapper for E.coli example data A list of SummarizedExperiment containing the output of rifi\_wrapper. The list contains 6 elements of SummarizedExperiment output of rifi\_preprocess, rifi\_fit, rifi\_penalties, rifi\_fragmentation, rifi\_stats and rifi\_summary. The plot is generated from rifi\_visualization. for more detail, please refer to each function separately.

#### Usage

data(wrapper\_e\_coli)

#### Format

An object of class list of length 6.

## Source

https://github.com/CyanolabFreiburg/rifi

wrapper_minimal	The result of rifi_wrapper for E.coli artificial example. A list
	of SummarizedExperiment containing the output of rifi_wrapper.
	The list contains 6 elements of SummarizedExperiment output of
	rifi_preprocess, rifi_fit, rifi_penalties, rifi_fragmentation, rifi_stats
	and rifi_summary. The plot is generated from rifi_visualization. for
	more detail, please refer to each function separately.

# Description

The result of rifi\_wrapper for E.coli artificial example. A list of SummarizedExperiment containing the output of rifi\_wrapper. The list contains 6 elements of SummarizedExperiment output of rifi\_preprocess, rifi\_fit, rifi\_penalties, rifi\_fragmentation, rifi\_stats and rifi\_summary. The plot is generated from rifi\_visualization. for more detail, please refer to each function separately.

## Usage

```
data(wrapper_minimal)
```

#### Format

An object of class list of length 6.

#### Source

https://github.com/CyanolabFreiburg/rifi

```
wrapper_summary_synechocystis_6803
```

The result of rifi\_wrapper for summary\_synechocystis\_6803 example data A list of SummarizedExperiment containing the output of rifi\_wrapper. The list contains 6 elements of SummarizedExperiment output of rifi\_preprocess, rifi\_fit, rifi\_penalties, rifi\_fragmentation, rifi\_stats and rifi\_summary. The plot is generated from rifi\_visualization. for more detail, please refer to each function separately.

# Description

The result of rifi\_wrapper for summary\_synechocystis\_6803 example data A list of SummarizedExperiment containing the output of rifi\_wrapper. The list contains 6 elements of SummarizedExperiment output of rifi\_preprocess, rifi\_fit, rifi\_penalties, rifi\_fragmentation, rifi\_stats and rifi\_summary. The plot is generated from rifi\_visualization. for more detail, please refer to each function separately.

#### Usage

data(wrapper\_summary\_synechocystis\_6803)

#### Format

An object of class list of length 6.

#### Source

https://github.com/CyanolabFreiburg/rifi

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