# Package 'TockyPrep'

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<b>Title</b> Data Preprocessing for Fluorescent Timer Reporters Using the Timer-Of-Cell-Kinetics-of-activitY (Tocky)
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<b>Description</b> This package provides data preprocessing methods for analyzing Fluorescent Timer data obtained by flow cytometry. Specifically, it provides the trigonometric transformation of Timer fluorescence to generate Timer Angle and Timer Intensity.
<b>Depends</b> R (>= 4.2.0), utils, stats, graphics, grDevices
Suggests knitr, rmarkdown, KernSmooth
VignetteBuilder knitr
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<pre>URL https://github.com/MonoTockyLab/TockyPrep, https:    //MonoTockyLab.github.io/TockyPrep</pre>
BugReports https://github.com/MonoTockyLab/TockyPrep/issues
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log\_data

Log-transform selected variables in a data frame

## **Description**

This function applies a custom log10 transformation to selected variables in a data frame. It allows optional adjustment of the data before logging and removal of rows with zeros in the transformed variables.

#### Usage

```
log_data(x, columns = NULL, add_offset = TRUE)
```

## **Arguments**

x A data frame containing the data to be log-transformed.

columns Optional character vector specifying the names of the columns to be log-transformed.

If NULL, the function will prompt the user to select columns interactively. De-

fault is NULL.

add\_offset Logical value indicating whether to adjust the data before logging by subtracting

the 1% quantile and adding 1. This helps in handling negative values and zeros.

Default is TRUE.

#### Value

A data frame with the selected variables log-transformed and combined with the remaining variables.

## **Examples**

```
## Not run:
# Create an example data frame
df <- data.frame(
    Var1 = rnorm(100, mean = 100, sd = 20),
    Var2 = rnorm(100, mean = 50, sd = 10),
    Var3 = rnorm(100, mean = 10, sd = 2)
)

# Log-transform Var1 and Var2 with adjustment
df_logged <- log_data(df, columns = c("Var1", "Var2"), add_offset = TRUE)
# Log-transform variables with interactive selection
df_logged <- log_data(df)
## End(Not run)</pre>
```

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plot_tocky	Generate basic QC plots for Tocky data

## **Description**

This function visualizes either Timer fluorescence (Blue vs Red) or Timer dynamics by the Tocky method (Angle vs Intensity) based on the specified mode.

## Usage

```
plot_tocky(
  file = "PlotTocky",
  pseudocolour = TRUE,
  pdf = FALSE,
  output = "QC",
  n = 4,
  plot_mode = "Timer fluorescence",
  lower_quantile_cutoff = 0.01,
  select = FALSE,
  group = TRUE,
  group_order = NULL,
  interactive = TRUE,
  save = FALSE,
  samplefile = NULL,
  verbose = TRUE
)
```

## **Arguments**

X	A TockyPrepData o	bject returned by '	'timer_transform',	which sample grouping
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has been defined by 'sample\_definition'.

file File name.

pseudocolour A logical argument for whether to use pseudocolour in plots.

A logical argument; if FALSE, a jpeg file is generated instead.

output The output directory name for output files.

n A number; n x n plots will be generated in the output Tocky plot file, max is 4

for 16 plots.

plot\_mode Either "Timer fluorescence" for Blue vs Red plots, "Normalized Timer fluo-

rescence" for normalized plots, or "Timer Angle and Intensity" for Angle vs

Intensity plots.

lower\_quantile\_cutoff

Lower quantile cutoff for setting the plot ranges in fluorescence mode.

select Logical indicating whether to manually select samples for plotting.

group Logical indicating whether to group plots based on the 'group' field in 'sam-

ple\_definition'.

group\_order Optional character vector for specifying the order of the panels when using the

group option.

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interactive Logical indicating whether to prompt the user to select plot\_mode. Defaults to

'TRUE'.

save A logical argument; if FALSE, plots are shown in an X window.

samplefile Character vector specifying the sample files. Defaults to 'NULL'.

verbose Logical indicating whether to print progress messages. Default is 'TRUE'.

## **Examples**

```
## Not run:
   plot_tocky(x, plot_mode = "Timer fluorescence")
   plot_tocky(x, plot_mode = "Timer Angle and Intensity")
## End(Not run)
```

prep\_tocky

Prepare Data for Timer Transformation Using Flow Cytometric Data

## **Description**

This function prepares the dataset for timer transformation analysis by identifying common variables across sample files, configuring necessary control files, and setting up variables for downstream analysis. The function supports both interactive and non-interactive file selection modes.

#### Usage

```
prep_tocky(path = ".", interactive = TRUE, negfile = NULL, samplefile = NULL)
```

#### **Arguments**

path	Character string specifying the directory where the data files are located. Defaults to the current directory ".".
interactive	Logical indicating whether to prompt the user to select sample files. Defaults to 'TRUE'.
negfile	Character string specifying the negative control file. If 'NULL', the user will be prompted to select a file. Defaults to 'NULL'.
samplefile	Character vector specifying the sample files. If 'NULL' and 'samplefilechoice'

is 'TRUE', the user will be prompted to select files. Defaults to 'NULL'.

## Value

A list containing paths to the control file, selected sample files, and the standardized variables used in the analysis.

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#### **Examples**

```
## Not run:
    # Interactive file selection
    prep_data <- prep_tocky(path='data', output='output')

# Specifying files directly for non-interactive usage
prep_data <- prep_tocky(
    path='data',
    output='output',
    negfile='neg_control.csv',
    samplefile=c('sample1.csv', 'sample2.csv')
)

## End(Not run)</pre>
```

sample\_definition

Update sample definitions and group assignments

### **Description**

This function takes the output from 'timer\_transform', specifically the 'sample\_definition' data frame, exports it to a CSV file for the user to edit group assignments, and then reads the updated file back into R.

#### Usage

```
sample_definition(
    x,
    sample_definition = NULL,
    output_dir = NULL,
    filename = "sample_definition.csv",
    sep = ",",
    verbose = TRUE,
    interactive = FALSE
)
```

#### **Arguments**

x A TockyPrepData object returned by 'timer\_transform', containing 'sample\_definition'. sample\_definition

(Optional) to use a data frame object as an input, specify the input object by this

parameter. Defaul is 'NULL'.

output\_dir Character string specifying the directory to save the 'sample\_definition.csv' file.

If 'NULL', the file is saved in the current working directory. Default is 'NULL'.

filename Character string specifying the name of the sample definition file. Default is

"sample definition.csv".

sep Character string indicating the field separator used in the CSV file. Default is

",""

verbose Logical indicating whether to display messages. Default is 'TRUE'.

interactive Logical indicating whether to use an interactive session to export a file for sam-

ple grouping and enable user to edit it and import. Defaults to 'TRUE'.

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#### Value

An updated TockyPrepData with user-assigned groupings.

## **Examples**

```
## Not run:
    # Assuming `result` is the output from `timer_transform`
    sample_def <- sample_definition(result, output_dir = "output_directory")
# The function will pause, allowing you to edit the 'group' column in the CSV file.
# After editing and saving the file, press Enter in R to continue.
# The updated sample definitions will be returned as a data frame.
## End(Not run)</pre>
```

timer\_transform

Perform Timer Transformation on Flow Cytometry Data

## Description

This function processes flow cytometry data by applying FSC correction, normalization, and trigonometric transformation to the Blue and Red fluorescence channels.

#### Usage

```
timer_transform(
  prep,
  select = TRUE,
  blue_channel = NULL,
  red_channel = NULL,
  normalization_method = "MAD",
  red_threshold = NULL,
  blue_threshold = NULL,
  interactive_gating = FALSE,
  verbose = TRUE,
  q = 0.998,
  normalization = TRUE
)
```

## **Arguments**

prep A list containing file paths and variables, typically the output from prep\_tocky.

select Logical indicating whether to choose Timer fluorescence channels interactively.

Default is 'TRUE'.

blue\_channel Character string specifying the Blue fluorescence channel name. If 'NULL', the

function attempts to determine it automatically.

red\_channel Character string specifying the Red fluorescence channel name. If 'NULL', the

function attempts to determine it automatically.

normalization\_method

Charcter string specifying the normalization method to be applied to Timer fl-

fuorescence Default is 'MAD'. The alternative is 'SD'.

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red\_threshold Numeric specifying the Red channel gating threshold. If 'NULL', gating is performed automatically or interactively based on 'interactive\_gating'. blue\_threshold Numeric specifying the Blue channel gating threshold. If 'NULL', gating is performed automatically or interactively based on 'interactive\_gating'. interactive\_gating

Logical indicating whether to perform interactive gating when thresholds are not provided. Default is 'FALSE'.

verbose Logical indicating whether to print progress messages. Default is 'TRUE'.

q Quantile value used for automatic Timer thresholds. Default is 0.998.

normalization Logical indicating whether to apply Timer fluorescence normalization. Default

is 'TRUE'.

#### Value

The function returns a new TockyPrepData object containing:

- transformed\_data: Data frame with angle, intensity, and other variables.
- normalization\_parameters: List with normalization values and coefficients.
- cell\_counts: Data frame with cell counts for each sample.
- sample\_definition: Data frame with sample file names and placeholder group column.

#### **Examples**

```
## Not run:
    # Assuming `prep_data` is the output from `prep_tocky`
    result <- timer_transform(prep_data)
## End(Not run)</pre>
```

TockyPrepData-class A class representing a TockyPrepData object for output of  $timer\_transform$ 

### **Description**

This class is designed to encapsulate and structure the output of the timer\_transform function in the TockyPrep package. It organizes various outputs into a coherent structure, ensuring that all necessary data components are included and validated. This structure is particularly useful for downstream processing and analysis in related packages like TockyLocus and TockyBase.

#### **Slots**

transformed\_data A data.frame containing transformed data.

cell\_counts A data.frame containing counts of cells per sample.

sample\_definition A data.frame containing metadata for each sample.

timer\_fluorescence A list containing fluorescence timing data.

input A list of raw input data.

normalization\_parameters A list of parameters used for data normalization.

Tocky A list containing other Tocky-specific analysis features

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