# Package 'workloopR'

October 21, 2019

Type	Package	
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**Title** Analysis of Work Loops and Other Data from Muscle Physiology Experiments

Version 1.0.1.9000

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Description Functions for the import, transformation, and analysis of data from muscle physiology experiments. The work loop technique is used to evaluate the mechanical work and power output of muscle. Josephson (1985) <a href="https://jeb.biologists.org/content/114/1/493">https://jeb.biologists.org/content/114/1/493</a> modernized the technique for application in comparative biomechanics. Although our initial motivation was to provide functions to analyze work loop experiment data, as we developed the package we incorporated the ability to analyze data from experiments that are often complementary to work loops. There are currently three supported experiment types: work loops, simple twitches, and tetanus trials. Data can be imported directly from .ddf files or via an object constructor function. Through either method, data can then be cleaned or transformed via methods typically used in studies of muscle physiology. Data can then be analyzed to determine the timing and magnitude of force development and relaxation (for isometric trials) or the magnitude of work, net power, and instantaneous power among other things (for work loops). Although we do not provide plotting functions, all resultant objects are designed to be friendly to visualization via either base-R plotting or 'tidyverse' functions.

URL https://github.com/vbaliga/workloopR

BugReports https://github.com/vbaliga/workloopR/issues
Imports pracma (>= 2.0.7),
 signal (>= 0.7-6)

Suggests testthat (>= 2.1.1),
 knitr,
 rmarkdown

License GPL (>= 3) + file LICENSE

Encoding UTF-8

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# R topics documented:

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analyze\_workloop

Analyze work loop object to compute work and power output

#### **Description**

Compute work and power output from a work loop experiment on a per-cycle basis.

# Usage

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```
analyze_workloop(x, simplify = FALSE, GR = 1, M = -1, vel_bf = 0.05, ...)
```

# **Arguments**

X	A workloop object of class muscle_stim that has been passed through select_cycles. See Details.
simplify	Logical. If FALSE, the full analyzed workloop object is returned. If TRUE a simpler table of net work and power (by cycle) is returned.
GR	Gear ratio, set to 1 by default
М	Velocity multiplier, set adjust the sign of velocity. This parameter should generally be either -1 (the default) or 1.
vel_bf	Critical frequency (scalar) for low-pass filtering of velocity via signal::butter()
•••	Additional arguments potentially passed down from read_analyze_wl() or read_analyze_wl_dir()

## **Details**

Please note that select\_cycles() must be run on data prior to using this function. This function relies on the input muscle\_stim object being organized by cycle number.

The muscle\_stim object (x) must be a workloop, preferably read in by one of our data import functions. Please see documentation for as\_muscle\_stim() if you need to manually construct a muscle\_stim object from a non .ddf source.

The gear ratio (GR) and velocity multiplier (M) parameters can help correct for issues related to the magnitude and sign of data collection. By default, they are set to apply no gear ratio adjustment and to positivize velocity. Instantaneous velocity is often noisy and the vel\_bf parameter allows for

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low-pass filtering of velocity data. See signal::butter() and signal::filtfilt() for details of how filtering is achieved.

Please also be careful with units! Se Warning section below.

#### Value

The function returns a list of class analyzed\_workloop that provides instantaneous velocity, a smoothed velocity, and computes work, instantaneous power, and net power from a work loop experiment. All data are organized by the cycle number and important metadata are stored as Attributes.

Within the list, each entry is labeled by cycle and includes:

Time, in sec

Position Length change of the muscle, corrected for gear ratio, in mm

Force, corrected for gear ratio, in mN

Stim When stimulation occurs, on a binary scale

Cycle ID, as a letter

Inst\_velocity Instantaneous velocity, computed from Position change, reported in meters/sec

Filt\_velocity Instantaneous velocity, after low-pass filtering, again in meter/sec

Inst\_Power Instantaneous power, a product of Force and Filt\_velocity, reported in J

Percent\_of\_Cycle

The percent of that particular cycle which has elapsed

In addition, the following information is stored in the analyzed\_workloop object's attributes:

stimulus\_frequency

Frequency at which stimulus pulses occurred

cycle\_frequency

Frequency of oscillations (assuming sine wave trajectory)

total\_cycles Total number of oscillatory cycles (assuming sine wave trajectory) that the mus-

cle experienced.

cycle\_def Specifies what part of the cycle is understood as the beginning and end. There

are currently three options: 'lo' for L0-to-L0; 'p2p' for peak-to-peak; and 't2t'

for trough-to-trough

amplitude Amplitude of length change (assuming sine wave trajectory)

phase Phase of the oscillatory cycle (in percent) at which stimulation occurred. Some-

what experimental, please use with caution

position\_inverted

Logical; whether position inversion has been applied)

units The units of measurement for each column in the object after running this func-

tion. See Warning

sample\_frequency

Frequency at which samples were collected

header Additional information from the header

units\_table Units from each Channel of the original ddf file

protocol\_table Protocol in tabular format; taken from the original ddf file

stim\_table Specific info on stimulus protocol; taken from the original ddf file

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stimulus\_pulses

Number of sequential pulses within a stimulation train

stimulus\_offset

Timing offset at which stimulus began

gear\_ratio Gear ratio applied by this function

file\_id File name

mtime Time at which file was last modified

retained\_cycles

Which cycles were retained, as numerics

summary Simple table showing work (in J) and net power (in W) for each cycle

## Warning

Most systems we have encountered record Position data in millimeters and Force in millinewtons, and therefore this function assumes data are recorded in those units. Through a series of internal conversions, this function computes velocity in meters/sec, work in Joules, and power in Watts. If your raw data do not originate in millimeters and millinewtons, please transform your data accordingly and ignore what you see in the attribute units.

## Author(s)

Vikram B. Baliga and Shreeram Senthivasan

#### References

Josephson RK. 1985. Mechanical Power output from Striated Muscle during Cyclic Contraction. Journal of Experimental Biology 114: 493-512.

#### See Also

```
read_ddf, read_analyze_wl, select_cycles
Other data analyses: isometric_timing, read_analyze_wl_dir, read_analyze_wl
Other workloop functions: fix_GR, get_wl_metadata, invert_position, read_analyze_wl_dir, read_analyze_wl, select_cycles, summarize_wl_trials, time_correct
```

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```
# summary provides a bit more detail
summary(wl_analyzed)

# run the analysis but get the simplified version
wl_analyzed_simple <- analyze_workloop(wl_selected, simplify = TRUE, GR = 2)</pre>
```

as\_muscle\_stim

Create your own muscle\_stim object

# **Description**

For use when data are not stored in .ddf format and you would like to create a muscle\_stim object that can be used by other workloopR functions.

#### Usage

```
as_muscle_stim(x, type, sample_frequency, ...)
```

# **Arguments**

x A data. frame. See Details for how it should be organized.

type Experiment type; must be one of: "workloop", "tetanus", or "twitch."

sample\_frequency

Numeric value of the frequency at which samples were recorded; must be in Hz.

Please format as numeric, e.g. 10000 works but 10000 Hz does not

. Additional arguments that can be passed in as attributes. See Details.

# Details

muscle\_stim objects, which are required by (nearly) all workloopR functions, are automatically created via read\_ddf(). Should you have data that are stored in a format other than .ddf, use this function to create your own object of class muscle\_stim.

The input x must be a data.frame that contains time series of numeric data collected from an experiment. Each row must correspond to a sample, and these columns (exact title matches) must be included:

"Time" - time, recorded in seconds

"Position" - instantaneous position of the muscle, preferably in millimeters

"Force" - force, preferably in millinewtons

"Stim" - whether stimulation has occurred. All entries must be either 0 (no stimulus) or 1 (stimulus occurrence).

Additional arguments can be provided via .... For all experiment types, the following attributes are appropriate:

"units", "header", "units\_table", "protocol\_table", "stim\_table", "stimulus\_pulses", "stimulus\_offset", "stimulus\_width", "gear\_ratio", "file\_id", or "mtime".

Please ensure that further attributes are appropriate to your experiment type.

For workloops, these include: "stimulus\_frequency", "cycle\_frequency", "total\_cycles", "cycle\_def", "amplitude", "phase", and "position\_inverted"

For twitches or tetanic trials: "stimulus\_frequency", and "stimulus\_length"

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#### Value

An object of class workloop, twitch, or tetanus, all of which inherit class muscle\_stim. These objects behave like data.frames in most situations but also store metadata from the ddf as attributes.

The muscle\_stim object's columns contain:

Time Time

Position Length change of the muscle, uncorrected for gear ratio

Force, uncorrected for gear ratio

Stim When stimulation occurs, on a binary scale

In addition, the following information is stored in the data. frame's attributes:

sample\_frequency

Frequency at which samples were collected

pulses Number of sequential pulses within a stimulation train

total\_cycles\_lo

Total number of oscillatory cycles (assuming sine wave trajectory) that the muscle experienced. Cycles are defined with respect to initial muscle length (L0-to-

L0 as opposed to peak-to-peak).

amplitude amplitude of length change (again, assuming sine wave trajectory)

cycle\_frequency

Frequency of oscillations (again, assuming sine wave trajectory)

units The units of measurement for each column in the data.frame. This might be

the most important attribute so please check that it makes sense!

## Author(s)

Shreeram Senthivasan

# See Also

```
read_ddf
```

```
Other data import functions: get_wl_metadata, read_analyze_wl_dir, read_analyze_wl, read_ddf_dir, read_ddf
```

fix\_GR

```
str(attributes(wl_dat))
## End(Not run)
```

fix\_GR

Adjust for the gear ratio of a motor arm

#### **Description**

Fix a discrepancy between the gear ratio of the motor arm used and the gear ratio recorded by software.

## Usage

```
fix_GR(x, GR = 1)
```

# Arguments

x A muscle\_stim object
GR Gear ratio, set to 1 by default

#### **Details**

The muscle\_stim object can be of any type, including workloop, twitch, or tetanus.

If you have manually constructed the object via as\_muscle\_stim(), the muscle\_stim object should have columns as follows:

Position: length change of the muscle;

Force: force

#### Value

An object of the same class(es) as the input (x). The function will multiply Position by (1/GR) and multiply Force by GR, returning an object with new values in \$Position and \$Force. Other columns and attributes are welcome and will simply be passed on unchanged into the resulting object.

# Author(s)

Vikram B. Baliga

### See Also

```
analyze_workloop, read_analyze_wl, read_analyze_wl_dir

Other data transformations: invert_position, select_cycles

Other workloop functions: analyze_workloop, get_wl_metadata, invert_position, read_analyze_wl_dir, read_analyze_wl, select_cycles, summarize_wl_trials, time_correct

Other twitch functions: invert_position, isometric_timing

Other tetanus functions: invert_position
```

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#### **Examples**

get\_wl\_metadata

Get file info for a sequence of experiment files

## **Description**

Grab metadata from files stored in the same folder (e.g. a sequence of trials in an experiment).

#### Usage

```
get_wl_metadata(file_path, pattern = "*.ddf")
```

# Arguments

file\_path Path where files are stored. Should be in the same folder.

Regex pattern for identifying relevant files in the file\_path.

# **Details**

If several files (e.g. successive trials from one experiment) are stored in one folder, use this function to obtain metadata in a list format. Runs file.info from base R to extract info from files.

This function is not truly considered to be part of the batch analysis pipeline; see read\_analyze\_wl\_dir() for a similar function that not only grabs metadata but also imports & analyzes files. Instead, get\_wl\_metadata() is meant to be a handy function to investigate metadata issues that arise if running read\_analyze\_wl\_dir() goes awry.

Unlike read\_analyze\_wl\_dir(), this function does not necessarily need files to all be work loops. Any file type is welcome (as long as the Regex pattern argument makes sense).

# Author(s)

Vikram B. Baliga

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#### See Also

```
summarize_wl_trials
Other data import functions: as_muscle_stim, read_analyze_wl_dir, read_analyze_wl, read_ddf_dir,
read_ddf
Other workloop functions: analyze_workloop, fix_GR, invert_position, read_analyze_wl_dir,
read_analyze_wl, select_cycles, summarize_wl_trials, time_correct
Other batch analyses: read_analyze_wl_dir, summarize_wl_trials, time_correct
```

# **Examples**

 ${\tt invert\_position}$ 

Invert the position data

#### **Description**

Multiply instantaneous position by -1.

# Usage

```
invert_position(x)
```

# **Arguments**

Х

A muscle\_stim object

# **Details**

The muscle\_stim object can be of any type, including workloop, twitch, or tetanus.

If you have manually constructed the object via as\_muscle\_stim(), the muscle\_stim object should have a column entitled Position. Other columns and attributes are welcome and will be passed along unchanged.

# Value

A workloop object with inverted position. The position\_inverted attribute is set to TRUE and all others are retained.

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#### Author(s)

Vikram B. Baliga

#### See Also

```
Other data transformations: fix_GR, select_cycles
```

Other workloop functions: analyze\_workloop, fix\_GR, get\_wl\_metadata, read\_analyze\_wl\_dir,

read\_analyze\_wl, select\_cycles, summarize\_wl\_trials, time\_correct

Other twitch functions: fix\_GR, isometric\_timing

Other tetanus functions: fix\_GR

# **Examples**

isometric\_timing

Compute timing and magnitude of force in isometric trials

# Description

Calculate timing and magnitude of force at stimulation, peak force, and various parts of the rising (force development) and relaxation (falling) phases of the twitch.

# Usage

```
isometric\_timing(x, rising = c(10, 90), relaxing = c(90, 50))
```

# Arguments

relaxing

X	A muscle_stim object that contains data from an isometric twitch trial, ideally created via read_ddf.
rising	Set points of the rising phase to be described. By default: 10% and 90%.

Set points of the relaxation phase to be described. By default: 90% and 50%.

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#### **Details**

The data. frame (x) must have time series data organized in columns. Generally, it is preferred that you use a muscle\_stim object imported by read\_ddf().

The rising and relaxing arguments allow for the user to supply numeric vectors of any length. By default, these arguments are rising = c(10,90) and relaxing = c(90,50). Numbers in each of these correspond to percent values and capture time and force at that percent of the corresponding curve. These values can be replaced by those that the user specifies and do not necessarily need to have length = 2. But please note that 0 and 100 should not be used, e.g. rising = seq(10,90,5) works, but rising = seq(0,100,5) does not.

#### Value

A data.frame with the following metrics as columns:

File ID file\_ID time\_stim Time between beginning of data collection and when stimulation occurs force\_stim Magnitude of force at the onset of stimulation Absolute time of peak force, i.e. time between beginning of data collection and time\_peak when peak force occurs Magnitude of peak force force\_peak time\_rising\_X Time between beginning of data collection and X% of force development force\_rising\_X Magnitude of force at X% of force development time\_relaxing\_X Time between beginning of data collection and X% of force relaxation force\_relaxing\_X

# Author(s)

Vikram B. Baliga

## References

Ahn AN, and Full RJ. 2002. A motor and a brake: two leg extensor muscles acting at the same joint manage energy differently in a running insect. Journal of Experimental Biology 205, 379-389.

#### See Also

```
Other data analyses: analyze_workloop, read_analyze_wl_dir, read_analyze_wl Other twitch functions: fix_GR, invert_position
```

Magnitude of force at X% of relaxation

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```
# we'll use different set points than the defaults
analyze_twitch <- isometric_timing(twitch_dat,
    rising = c(25, 50, 75),
    relaxing = c(75, 50, 25)
)

# see the results
analyze_twitch</pre>
```

read\_analyze\_wl

All-in-one import function for work loop files

#### **Description**

read\_analyze\_wl() is an all-in-one function to read in a work loop file, select cycles, and compute work and power output.

#### Usage

```
read_analyze_wl(file_name, ...)
```

#### **Arguments**

file\_name A .ddf file that contains data from a single workloop experiment
... Additional arguments to be passed to read\_ddf(), select\_cycles(), or analyze\_workloop().

#### Details

Please be careful with units! See Warnings below. This function combines read\_ddf() with select\_cycles() and then ultimately analyze\_workloop() into one handy function.

As detailed in these three functions, possible arguments include:

cycle\_def - used to specify which part of the cycle is understood as the beginning and end. There are currently three options: 'lo' for L0-to-L0; 'p2p' for peak-to-peak; and 't2t' for trough-to-trough bworth\_order - Filter order for low-pass filtering of Position via signal::butter prior to finding peak lengths. Default: 2.

bworth\_freq - Critical frequency (scalar) for low-pass filtering of Position via signal::butter prior to finding peak lengths. Default: 0.05.

keep\_cycles - Which cycles should be retained. Default: 4:6.

GR - Gear ratio. Default: 1.

M - Velocity multiplier used to positivize velocity; should be either -1 or 1. Default: -1.

vel\_bf - Critical frequency (scalar) for low-pass filtering of velocity via signal::butter. Default: 0.05.

The gear ratio (GR) and velocity multiplier (M) parameters can help correct for issues related to the magnitude and sign of data collection. By default, they are set to apply no gear ratio adjustment and to positivize velocity. Instantaneous velocity is often noisy and the vel\_bf parameter allows for low-pass filtering of velocity data. See signal::butter() and signal::filtfilt() for details of how filtering is achieved.

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#### Value

The function returns a list of class analyzed\_workloop that provides instantaneous velocity, a smoothed velocity, and computes work, instantaneous power, and net power from a work loop experiment. All data are organized by the cycle number and important metadata are stored as Attributes.

Within the list, each entry is labeled by cycle and includes:

Time, in sec

Position Length change of the muscle, corrected for gear ratio, in mm

Force, corrected for gear ratio, in mN

Stim When stimulation occurs, on a binary scale

Cycle ID, as a letter

Inst\_velocity Instantaneous velocity, computed from Position change, reported in meters/sec

Filt\_velocity Instantaneous velocity, after low-pass filtering, again in meter/sec

Inst\_Power Instantaneous power, a product of Force and Filt\_velocity, reported in J

Percent\_of\_Cycle

The percent of that particular cycle which has elapsed

In addition, the following information is stored in the analyzed\_workloop object's attributes:

stimulus\_frequency

Frequency at which stimulus pulses occurred

cycle\_frequency

Frequency of oscillations (assuming sine wave trajectory)

total\_cycles Total number of oscillatory cycles (assuming sine wave trajectory) that the mus-

cle experienced.

cycle\_def Specifies what part of the cycle is understood as the beginning and end. There

are currently three options: 'lo' for L0-to-L0; 'p2p' for peak-to-peak; and 't2t'

for trough-to-trough

amplitude Amplitude of length change (assuming sine wave trajectory)

phase Phase of the oscillatory cycle (in percent) at which stimulation occurred. Some-

what experimental, please use with caution

position\_inverted

Logical; whether position inversion has been applied)

units The units of measurement for each column in the object after running this func-

tion. See Warning

sample\_frequency

Frequency at which samples were collected

header Additional information from the header

units\_table Units from each Channel of the original ddf file

protocol\_table Protocol in tabular format; taken from the original ddf file

stim\_table Specific info on stimulus protocol; taken from the original ddf file

stimulus\_pulses

Number of sequential pulses within a stimulation train

stimulus\_offset

Timing offset at which stimulus began

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gear\_ratio Gear ratio applied by this function

file\_id File name

mtime Time at which file was last modified

retained\_cycles

Which cycles were retained, as numerics

summary Simple table showing work (in J) and net power (in W) for each cycle

#### Warning

Most systems we have encountered record Position data in millimeters and Force in millinewtons, and therefore this function assumes data are recorded in those units. Through a series of internal conversions, this function computes velocity in meters/sec, work in Joules, and power in Watts. If your raw data do not originate in millimeters and millinewtons, please transform your data accordingly and ignore what you see in the attribute units.

#### Author(s)

Vikram B. Baliga

#### References

Josephson RK. 1985. Mechanical Power output from Striated Muscle during Cyclic Contraction. Journal of Experimental Biology 114: 493-512.

#### See Also

```
read_ddf, select_cycles analyze_workloop

Other data analyses: analyze_workloop, isometric_timing, read_analyze_wl_dir

Other data import functions: as_muscle_stim, get_wl_metadata, read_analyze_wl_dir, read_ddf_dir, read_ddf

Other workloop functions: analyze_workloop, fix_GR, get_wl_metadata, invert_position, read_analyze_wl_dir, select_cycles, summarize_wl_trials, time_correct
```

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read\_analyze\_wl\_dir Read and analyze work loop files from a directory

## **Description**

All-in-one function to import multiple workloop .ddf files from a directory, sort them by mtime, analyze them, and store the resulting objects in an ordered list.

# Usage

```
read_analyze_wl_dir(file_path, pattern = "*.ddf", sort_by = "mtime",
    ...)
```

# **Arguments**

file_path	Directory in which files are located
pattern	Regular expression used to specify files of interest. Defaults to all .ddf files within file_path
sort_by	Metadata by which files should be sorted to be in the correct run order. Defaults to mtime, which is time of last modification of files.
	Additional arguments to be passed to read_analyze_wl(), analyze_workloop(), select_cycles(), or read_ddf().

#### **Details**

Work loop data files will be imported and then arranged in the order in which they were run (assuming run order is reflected in mtime). Chiefly used in conjunction with summarize\_wl\_trials() and time\_correct() if time correction is desired.

### Value

A list containing analyzed\_workloop objects, one for each file that is imported and subsequently analyzed. The list is sorted according to the sort\_by parameter, which by default uses the time of last modification of each file's contents (mtime).

#### Warning

Most systems we have encountered record Position data in millimeters and Force in millinewtons, and therefore this function assumes data are recorded in those units. Through a series of internal conversions, this function computes velocity in meters/sec, work in Joules, and power in Watts. If your raw data do not originate in millimeters and millinewtons, please transform your data accordingly and ignore what you see in the attribute units.

# Author(s)

Shreeram Senthivasan

## References

Josephson RK. 1985. Mechanical Power output from Striated Muscle during Cyclic Contraction. Journal of Experimental Biology 114: 493-512.

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#### See Also

```
read_analyze_wl, get_wl_metadata, summarize_wl_trials, time_correct

Other data analyses: analyze_workloop, isometric_timing, read_analyze_wl

Other data import functions: as_muscle_stim, get_wl_metadata, read_analyze_wl, read_ddf_dir, read_ddf

Other workloop functions: analyze_workloop, fix_GR, get_wl_metadata, invert_position, read_analyze_wl, select_cycles, summarize_wl_trials, time_correct

Other batch analyses: get_wl_metadata, summarize_wl_trials, time_correct
```

# **Examples**

read\_ddf

Import work loop or isometric data from .ddf files

#### **Description**

read\_ddf reads in workloop, twitch, or tetanus experiment data from .ddf files.

# Usage

```
read_ddf(file_name, file_id = NA, rename_cols = list(c(2, 3),
    c("Position", "Force")), skip_cols = 4:11, phase_from_peak = FALSE,
    ...)
```

# **Arguments**

file_name	A .ddf file that contains data from a single workloop, twitch, or tetanus experiment
file_id	A string identifying the experiment. The file name is used by default.
rename_cols	List consisting of a vector of indices of columns to rename and a vector of new column names. See Details.
skip_cols	Numeric vector of column indices to skip. See Details.
phase_from_peal	<
	Logical, indicating whether percent phase of stimulation should be recorded relative to peak length or relative to L0 (default)
	Additional arguments passed to/from other functions that work with read_ddf()

read\_ddf

#### **Details**

Read in a .ddf file that contains data from an experiment. If position and force do not correspond to columns 2 and 3 (respectively), replace "2" and "3" within rename\_cols accordingly. Similarly, skip\_cols = 4:11 should be adjusted if more than 11 columns are present and/or columns 4:11 contain important data.

Please note that there is no correction for gear ratio or further manipulation of data. See fix\_GR to adjust gear ratio. Gear ratio can also be adjusted prior to analyses within the analyze\_workloop() function, the data import all-in-one function read\_analyze\_wl(), or the batch analysis all-in-one read\_analyze\_wl\_dir().

Please also note that organization of data within the .ddf file is assumed to conform to that used by Aurora Scientific's Dynamic Muscle Control and Analysis Software. YMMV if using a .ddf file from another source. The as\_muscle\_stim() function can be used to generate muscle\_stim objects if data are imported via another function. Please feel free to contact us with any issues or requests.

#### Value

An object of class workloop, twitch, or tetanus, all of which inherit class muscle\_stim. These objects behave like data.frames in most situations but also store metadata from the ddf as attributes.

The muscle\_stim object's columns contain:

Time Time

Position Length change of the muscle, uncorrected for gear ratio

Force, uncorrected for gear ratio

Stim When stimulation occurs, on a binary scale

In addition, the following information is stored in the data. frame's attributes:

sample\_frequency

Frequency at which samples were collected

pulses Number of sequential pulses within a stimulation train

total\_cycles\_lo

Total number of oscillatory cycles (assuming sine wave trajectory) that the muscle experienced. Cycles are defined with respect to initial muscle length (L0-to-

L0 as opposed to peak-to-peak).

amplitude amplitude of length change (again, assuming sine wave trajectory)

cycle\_frequency

Frequency of oscillations (again, assuming sine wave trajectory)

units The units of measurement for each column in the data. frame. This might be

the most important attribute so please check that it makes sense!

#### Author(s)

Vikram B. Baliga and Shreeram Senthivasan

## See Also

Other data import functions: as\_muscle\_stim, get\_wl\_metadata, read\_analyze\_wl\_dir, read\_analyze\_wl, read\_ddf\_dir

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#### **Examples**

read\_ddf\_dir

Import a batch of work loop or isometric data files from a directory

#### **Description**

Uses read\_ddf() to read in workloop, twitch, or tetanus experiment data from multiple .ddf files.

#### Usage

```
read_ddf_dir(file_path, pattern = "*.ddf", sort_by = "mtime", ...)
```

# Arguments

file_path	Path where files are stored. Should be in the same folder.
pattern	Regex pattern for identifying relevant files in the file_path.
sort_by	Metadata by which files should be sorted to be in the correct run order. Defaults to mtime, which is time of last modification of files.
	Additional arguments to be passed to read_ddf().

## **Details**

Read in a .ddf file that contains data from an experiment. If position and force do not correspond to columns 2 and 3 (respectively), replace "2" and "3" within rename\_cols accordingly. Similarly, skip\_cols = 4:11 should be adjusted if more than 11 columns are present and/or columns 4:11 contain important data.

Please note that there is no correction for gear ratio or further manipulation of data. See fix\_GR to adjust gear ratio. Gear ratio can also be adjusted prior to analyses within the analyze\_workloop() function, the data import all-in-one function read\_analyze\_wl(), or the batch analysis all-in-one read\_analyze\_wl\_dir().

Please also note that organization of data within the .ddf file is assumed to conform to that used by Aurora Scientific's Dynamic Muscle Control and Analysis Software. YMMV if using a .ddf file from another source. The as\_muscle\_stim() function can be used to generate muscle\_stim objects if data are imported via another function. Please feel free to contact us with any issues or requests.

read\_ddf\_dir

#### Value

A list of objects of class workloop, twitch, or tetanus, all of which inherit class muscle\_stim. These objects behave like data.frames in most situations but also store metadata from the ddf as attributes.

Each muscle\_stim object's columns contain:

Time Time

Position Length change of the muscle, uncorrected for gear ratio

Force, uncorrected for gear ratio

Stim When stimulation occurs, on a binary scale

In addition, the following information is stored in each data. frame's attributes:

sample\_frequency

Frequency at which samples were collected

pulses Number of sequential pulses within a stimulation train

total\_cycles\_lo

Total number of oscillatory cycles (assuming sine wave trajectory) that the muscle experienced. Cycles are defined with respect to initial muscle length (L0-to-

L0 as opposed to peak-to-peak).

amplitude amplitude of length change (again, assuming sine wave trajectory)

cycle\_frequency

Frequency of oscillations (again, assuming sine wave trajectory)

units The units of measurement for each column in the data.frame. This might be

the most important attribute so please check that it makes sense!

#### Author(s)

Vikram B. Baliga and Shreeram Senthivasan

#### See Also

Other data import functions: as\_muscle\_stim, get\_wl\_metadata, read\_analyze\_wl\_dir, read\_analyze\_wl, read\_ddf

20 select\_cycles

select_cycles Select cycles from a work loop object
---

# **Description**

Retain data from a work loop experiment based on position cycle

## Usage

```
select_cycles(x, cycle_def, keep_cycles = 4:6, bworth_order = 2,
   bworth_freq = 0.05, ...)
```

#### **Arguments**

Х	A workloop object (see Details for how it should be organized)
cycle_def	A string specifying how cycles should be defined; one of: "lo", "p2p", or "t2t". See Details more info
keep_cycles	The indices of the cycles to keep. Include 0 to keep data identified as being outside complete cycles
bworth_order	Filter order for low-pass filtering of Position via signal::butter() prior to finding L0
bworth_freq	Critical frequency (scalar) for low-pass filtering of Position via signal::butter() prior to finding L0
	Additional arguments passed to/from other functions that make use of select_cycles()

# **Details**

select\_cycles() subsets data from a workloop trial by position cycle. The cycle\_def argument is used to specify which part of the cycle is understood as the beginning and end. There are currently three options:

```
'lo' for L0-to-L0;
```

'p2p' for peak-to-peak; and 't2t' for trough-to-trough

Peaks are identified using pracma::findpeaks(). L0 points on the rising edge are found by finding the midpoints between troughs and the following peak. However the first and last extrema and L0 points may be misidentified by this method. Please plot your Position cycles to ensure the edge cases are identified correctly.

The keep\_cycles argument is used to determine which cycles (as defined by cycle\_def should be retained in the final dataset. Zero is the index assigned to all data points that are determined to be outside a complete cycle.

The  $muscle\_stim$  object (x) must be a workloop, preferably read in by one of our data import functions. Please see documentation for  $as\_muscle\_stim()$  if you need to manually construct a  $muscle\_stim$  object from another source.

#### Value

A workloop object with rows subsetted by the chosen position cycles. A Cycle column is appended to denote which cycle each time point is associated with. Finally, all attributes from the input workloop object are retained and one new attribute is added to record which cycles from the original data were retained.

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#### Author(s)

Vikram B. Baliga and Shreeram Senthivasan

#### See Also

```
analyze_workloop, read_analyze_wl, read_analyze_wl_dir
Other data transformations: fix_GR, invert_position
Other workloop functions: analyze_workloop, fix_GR, get_wl_metadata, invert_position, read_analyze_wl_dir, read_analyze_wl, summarize_wl_trials, time_correct
```

#### **Examples**

```
summarize\_wl\_trials \qquad \textit{Summarize work loop files}
```

# **Description**

Summarize important info from work loop files stored in the same folder (e.g. a sequence of trials in an experiment) including experimental parameters, run order, and mtime.

# Usage

```
summarize_wl_trials(wl_list)
```

# **Arguments**

```
wl_list List of analyzed_workloop objects, preferably one created by read_analyze_wl_dir().
```

22 summarize\_wl\_trials

#### **Details**

If several files (e.g. successive trials from one experiment) are stored in one folder, use this function to obtain summary stats and metadata and other parameters. This function requires a list of analyze\_workloop objects, which can be readily obtained by first running read\_analyze\_wl\_dir() on a specified directory.

#### Value

A data.frame of information about the collection of workloop files. Columns include:

File\_ID Name of the file

Cycle\_Frequency

Frequency of Position change

Amplitude amplitude of Position change

Phase Phase of the oscillatory cycle (in percent) at which stimulation occurred. Some-

what experimental, please use with caution

Stimulus\_Pulses

Number of stimulation pulses

mtime Time at which file's contents were last changed (mtime)

Mean\_Work Mean work output from the selected cycles

Mean\_Power Net power output from the selected cycles

#### Author(s)

Vikram B. Baliga and Shreeram Senthivasan

#### References

Josephson RK. 1985. Mechanical Power output from Striated Muscle during Cyclic Contraction. Journal of Experimental Biology 114: 493-512.

## See Also

```
read_analyze_wl_dir, get_wl_metadata, time_correct

Other workloop functions: analyze_workloop, fix_GR, get_wl_metadata, invert_position, read_analyze_wl_dir, read_analyze_wl, select_cycles, time_correct

Other batch analyses: get_wl_metadata, read_analyze_wl_dir, time_correct
```

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```
summarized_wls <- summarize_wl_trials(analyzed_wls)
# or on your own directory
## Not run:
my_meta <- read_analyze_wl_dir("./my/file/path/")
my_summaries <- summarize_wl_trials(my_meta)
## End(Not run)</pre>
```

time\_correct

Time correction for work loop experiments

# Description

Correct for potential degradation of muscle over time.

# Usage

```
time_correct(x)
```

#### **Arguments**

Х

A data.frame with summary data, e.g. an object created by summarize\_wl\_trials().

#### Details

This function assumes that across a batch of successive trials, the stimulation parameters for the first and final trials are identical. If not, DO NOT USE. Decline in power output is therefore assumed to be a linear function of time. Accordingly, the difference between the final and first trial's (absolute) power output is used to 'correct' trials that occur in between, with explicit consideration of run order and time elapsed (via mtime). A similar correction procedure is applied to work.

#### Value

```
A data. frame that additionally contains:
```

Time\_Corrected\_Work

Time corrected work output, transformed from \$Mean\_Work

Time\_Corrected\_Power

Time corrected net power output, transformed from \$Mean\_Power

And new attributes:

power\_difference

Difference in mass-specific net power output between the final and first trials.

time\_difference

Difference in mtime between the final and first trials.

time\_correction\_rate

Overall rate; power\_difference divided by time\_difference.

#### Author(s)

Vikram B. Baliga and Shreeram Senthivasan

#### See Also

```
summarize_wl_trials
Other workloop functions: analyze_workloop, fix_GR, get_wl_metadata, invert_position,
read_analyze_wl_dir, read_analyze_wl, select_cycles, summarize_wl_trials
Other batch analyses: get_wl_metadata, read_analyze_wl_dir, summarize_wl_trials
```

# **Examples**

```
library(workloopR)
# batch read and analyze files included with workloopR
analyzed_wls <- read_analyze_wl_dir(system.file("extdata/wl_duration_trials",</pre>
                                                   package = 'workloopR'),
                                      phase_from_peak = TRUE,
                                      cycle_def = "p2p", keep_cycles = 2:4)
# now summarize
summarized_wls <- summarize_wl_trials(analyzed_wls)</pre>
# mtimes within the package are not accurate, so we'll supply
# our own vector of mtimes
summarized_wls$mtime <- read.csv(</pre>
                           system.file(
                              "extdata/wl_duration_trials/ddfmtimes.csv",
                              package="workloopR"))$mtime
# now time correct
timecor_wls <- time_correct(summarized_wls)</pre>
timecor_wls
# or on your own directory
## Not run:
my_meta <- read_analyze_wl_dir("./my/file/path/")</pre>
my_summaries <- summarize_wl_trials(my_meta)</pre>
my_timecors <- time_correct(my_summaries)</pre>
## End(Not run)
```

 $trapezoidal\_integration$ 

Approximate the definite integral via the trapezoidal rule

# Description

Mostly meant for internal use in our analysis functions, but made available for other use cases. Accordingly, it does not strictly rely on objects of class muscle\_stim.

#### Usage

```
trapezoidal_integration(x, f)
```

# **Arguments**

```
x a variable, e.g. vector of positions
f integrand, e.g. vector of forces
```

# **Details**

In the functions analyze\_workloop(), read\_analyze\_wl(), and read\_analyze\_wl\_dir(), work is calculated as the difference between the integral of the upper curve and the integral of the lower curve of a work loop.

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# Author(s)

Vikram B. Baliga

#### References

Atkinson, Kendall E. (1989), An Introduction to Numerical Analysis (2nd ed.), New York: John Wiley & Sons

#### See Also

```
analyze_workloop, read_analyze_wl, read_analyze_wl_dir
```

```
# create a circle centered at (x = 10, y = 20) with radius 2
t <- seq(0, 2 * pi, length = 1000)
coords <- t(rbind(10 + sin(t) * 2, 20 + cos(t) * 2))

# sanity check: does it look like a circle?
## Not run:
plot(coords, asp = 1)

## End(Not run)

# use the function to get the area
trapezoidal_integration(coords[, 1], coords[, 2])

# does it match (pi * r^2)?
3.14159265358 * (2^2) # very close</pre>
```

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