Package 'clintools'

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Type Package

Title Tools for Clinical Research

Version 0.8.5

Description Every research team have their own script for data management, statistics and most importantly hemodynamic indices. The purpose is to standardize scripts utilized in clinical research. The hemodynamic indices can be used in a long-format dataframe, and add both periods of interest (trigger-periods), and delete artifacts with deleter-files. Transfer function analysis (Claassen et al. (2016) <doi:10.1177/0271678X15626425>) and Mx (Czosnyka et al. (1996) <doi:10.1161/01.str.27.10.1829>) can be calculated using this package.

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URL https://github.com/lilleoel/clintools

BugReports https://github.com/lilleoel/clintools/issues

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Description

clinmon() uses a *continuous* recording and returns a dataframe with hemodynamic indices for every period, epoch or block depending on the input. Calculates COest, CPPopt, CVRi, Dx, Mx, PI, PRx, PWA, RI, and Sx (see *Hemodynamic indices*).

Usage

```
clinmon(df, variables,
trigger = NULL, deleter = NULL,
blocksize = 3, epochsize = 20,
overlapping = FALSE, freq = 1000,
blockmin = 0.5, epochmin = 0.5,
output = "period", fast = FALSE)
```

Arguments

df	Raw <i>continuous</i> recording with all numeric data and first column has to be time in seconds. (dataframe)
variables	Defining the type and order of the recorded variables as a list. Middle cerebral artery blood velocity ('mcav'), Arterial blood pressure ('abp'), cerebral perfusion pressure ('cpp'), intracranial pressure ('icp'), and heart rate ('hr') is currently supported. It is necessary that time is the first row. (list)
trigger	Trigger with two columns: first is start, and second is end of periods to be analyzed. Every row corresponds to a period. Default is NULL, which results in analysis of the full dataframe. (dataframe)
deleter	Deleter with two columns: first is start and second is end of period with artefacts, which need to be deleted. Every row is a period with artefacts. Default is NULL. (dataframe)

blocksize	Length of a block, in seconds. Default is 3. (numeric)
epochsize	Size of epochs in number of blocks. Default is 20. (numeric)
overlapping	The number of block which should overlap when calculating correlation based indices, and remain blank if overlapping calculations should not be utilized. Default is FALSE. (numeric)
freq	Frequency of recorded data, in Hz. Default is 1000. (numeric)
blockmin	Minimum measurements required to create a block in ratio. Default is 0.5 corresponding to 50% . If the block holds less than the defined ratio the block will be omitted. (numeric)
epochmin	Minimum number of blocks required to create an epoch in ratio. Default is 0.5 corresponding to 50% . If the epoch holds less than the defined ration the epoch will be omitted. (numeric)
output	Select what each row should represent in the output. Correlation based indices are not presented when selecting blocks for every row. Currently 'block', 'epoch', 'period' or 'cppopt' is supported. Default is 'period'. (string)
fast	Select if you want the data to aggregated before analysis resulting in a faster, but perhaps more imprecise run, in Hz. Default is FALSE. (numeric)

Details

Using a *continuous* raw recording, clinmon() calculates hemodynamic indices for every period, epoch or block depending on the chosen output.

View(data)

time	abp	mcav
7.00	78	45
7.01	78	46
301.82	82	70
301.83	81	69

To calculate the indices insert the data and select the relevant variables.

```
clinmon(df=data, variables=c("abp","mcav"))
```

See Value for output description.

Value

Returns a dataframe with the results, with either every blocks, epochs or periods as rows, depending on the chosen output.

The columns of the output are:

• period - The period number corresponding to the row-number in the trigger file.

- epoch The epoch number, or if period is chosen as output it reflects the number of epochs in the period.
- block The block number, or if period or epoch is chosen as output it reflects the number of blocks in the period or epoch.
- time_min The minimum time value or the period, epoch or block.
- time_max The maximum time value or the period, epoch or block.
- missing_percent The percentage of missing data in the period, epoch or block.
- XX_mean The mean value of each variable for the period, epoch or block.
- XX_min The minimum value of each variable for the period, epoch or block.
- XX_max The maximum value of each variable for the period, epoch or block.
- YY The indices in each column.

Hemodynamic indices

COest | Estimated cardiac output:

Required variables: abp, hr; Required output: -.

Estimated cardiac output (COest) is calculated by utilizing the method described by Koenig et al. [1]:

$$COest = PP/(SBP + DBP) * HR$$

PP: Pulse pressure; SBP: systolic blood pressure; DBP: diastolic blood pressure; HR: heart rate.

CPPopt | Optimal cerebral perfusion pressure:

Required variables: abp, icp; Required output: period.

Optimal cerebral perfusion pressure (CPPopt) is calculated utilizing the method described by Steiner et al. [2]. The CPPopt return NA if CPPopt is the maximum or minimum CPP investigated. CPPopt is recommended to only be calculated after 'several hours' of recording:

$$CPPopt = The 5mmHqCPPIntervalWithLowestMeanPRx$$

CPP: cerebral perfusion pressure; PRx: Pressure reactivity index.

CVRi | Cardiovascular resistance index:

Required variables: abp, mcav; Required output: -.

Cardiovascular resistance index (CVRi) is calculated utilizing the method described by Fan et al. [3]:

$$CVRi = meanABP/meanMCAv$$

ABP: arterial blood pressure; MCAv: middle cerebral artery blood velocity.

Dx | Diastolic flow index:

Required variables: cpp/abp, mcav; Required output: epoch, period.

Diastolic flow index (Dx) is calculated utilizing the method described by Reinhard et al. [4]:

$$Dx = cor(meanCPP/minMCAv)$$

$$Dxa = cor(meanABP/minMCAv)$$

cor: correlation coefficient; CPP: cerebral perfusion pressure; ABP: arterial blood pressure; MCAv: middle cerebral artery blood velocity.

Mx | Mean flow index:

Required variables: cpp/abp, mcav; Required output: epoch, period.

Mean flow index (Mx) is calculated utilizing the method described by Czosnyka et al. [5]:

$$Mx = cor(meanCPP/meanMCAv)$$

$$Mxa = cor(meanABP/meanMCAv)$$

cor: correlation coefficient; CPP: cerebral perfusion pressure; ABP: arterial blood pressure; MCAv: middle cerebral artery blood velocity.

PI | Gosling index of pulsatility:

Required variables: mcav; Required output: -.

Gosling index of pulsatility (PI) is calculated utilizing the method described by Michel et al. [6]:

$$PI = (systolicMCAv - diastolicMCAv)/meanMCAv$$

MCAv: middle cerebral artery blood velocity.

PRx | Pressure reactivity index:

Required variables: abp, icp; Required output: epoch, period.

Pressure reactivity index (PRx) is calculated utilizing the method described by Czosnyka et al. [7]:

$$PRx = cor(meanABP/meanICP)$$

cor: correlation coefficient; CPP: cerebral perfusion pressure; ICP: intracranial pressure.

PWA | Pulse wave amplitude:

Required variables: cpp/icp/abp/mcav; Required output: -.

Pulse wave amplitude (PWA) is calculated utilizing the method described by Norager et al. [8]:

$$PWA = systolic - diastolic$$

RI | Pourcelots resistive (resistance) index:

Required variables: mcav; Required output: -.

Pourcelots resistive (resistance) index (RI) is calculated utilizing the method described by Forster et al. [9]:

$$RI = (systolicMCAv - diastolicMCAv)/systolicMCAv$$

MCAv: middle cerebral artery blood velocity.

Sx | Systolic flow index:

Required variables: cpp/abp, mcav; Required output: epoch, period.

Systolic flow index (Sx) is calculated utilizing the method described by Czosnyka et al. [5]:

$$Sx = cor(meanCPP/systolicMCAv)$$

$$Sxa = cor(meanABP/systolicMCAv)$$

cor: correlation coefficient; CPP: cerebral perfusion pressure; ABP: arterial blood pressure; MCAv: middle cerebral artery blood velocity.

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References

- 1. Koenig et al. (2015) Biomed Sci Instrum. 2015;51:85-90. (PubMed)
- 2. Steiner et al. (2002) Crit Care Med. 2002 Apr;30(4):733-8. (PubMed)
- 3. Fan et al. (2018) Front Physiol. 2018 Jul 16;9:869. (PubMed)
- 4. Reinhard et al. (2003) Stroke. 2003 Sep;34(9):2138-44. (PubMed)
- 5. Czosnyka et al. (1996) Stroke. 1996 Oct;27(10):1829-34. (PubMed)
- 6. Michel et al. (1998) Ultrasound Med Biol. 1998 May;24(4):597-9. (PubMed)
- 7. Czosnyka et al. (1997) Neurosurgery. 1997 Jul;41(1):11-7; discussion 17-9. (PubMed)
- 8. Norager et al. (2020) Acta Neurochir (Wien). 2020 Dec;162(12):2983-2989. (PubMed)
- 9. Forster et al. (2017) J Paediatr Child Health. 2018 Jan;54(1):61-68. (PubMed)

Examples

```
data(testdata)
clinmon(df.data10, variables=c('abp','mcav','hr'), freq=10)
```

df.data1000

Test-data - 1000 Hz

Description

Recording with four columns: time (t), non-invasive arterial blood pressure (abp), middle cerebral artery velocity measured using transcranial Doppler (mcav), and heart rate (hr).

Usage

```
data(testdata)
```

Format

An object of class "dataframe"; an example of the usage in clinmon-function.

References

```
Olsen MH et al. (Unpublished data, 2020) (GitHub)
```

```
data(testdata)
variables <- c("abp","mcav","hr")
clinmon(df.data1000,variables,fast=50)</pre>
```

df.deleter 7

df.deleter

Test-deleter

Description

Deleter dataframe with two columns: start (start) and end (end) of the deleter-period.

Usage

```
data(testdata)
```

Format

An object of class "dataframe"; an example of the usage in clinmon-function.

References

```
Olsen MH et al. (Unpublished data, 2020) (GitHub)
```

Examples

```
data(testdata)
variables <- c("abp","mcav","hr")
clinmon(df.data1000,variables,deleter=df.deleter,fast=50)</pre>
```

iscus

ISCUSFlex-values to dataframe (iscus)

Description

iscus() is a function which converts XML files extracted from the Microdialyis-apparatur of IS-CUSFlex apparatus to a dataframe.

Usage

```
iscus(filename)
```

Arguments

filename

path to the XML-file with the measurements

Value

Returns a dataframe with the measurements.

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Examples

```
## Not run:
    iscus("C:/ISCUSfiles/7888e844-1c7a-40af-a3f2-3bb27a8dd9e5.xml")
## End(Not run)
```

ortable

Logistic regression table with Odds ratio (ortable)

Description

ortable() is a small function which utilises the output from the glm-function to print a dataframe with odds ratio, confidence limits, and p-values.

Usage

```
ortable(x, d, d_p, intercept, simple)
```

Arguments

X	Utlises the output from a glm-function. (glm-output)
d	Refers to the number of digits for odds ratio and confidence intervals. Default is 2. (numeric)
d_p	Refers to the number of digits for odds ratio and confidence intervals. Default is 3. (numeric)
intercept	The intercept is presented in the table if TRUE. Default is FALSE. (boolian)
simple	Odds ratio and confidence intervals are merged into one column if TRUE. Default is TRUE. (boolian)

Value

Returns a dataframe with with odds ratio, confidence limits, and p-values.

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testdata10

Test-data - 10 Hz

Description

Recording with four columns: time (t), non-invasive arterial blood pressure (abp), middle cerebral artery velocity measured using transcranial Doppler (mcav), and heart rate (hr).

Usage

```
data(testdata)
```

Format

An object of class "dataframe"; an example of the usage in clinmon-function.

References

```
Olsen MH et al. (Unpublished data, 2020) (GitHub)
```

Examples

```
data(testdata)
variables <- c("abp","mcav","hr")
clinmon(df.data10,variables,freq=10)</pre>
```

TFA

Transfer function analysis of dynamic cerebral autoregulation (TFA)

Description

TFA() calculates dynamic cerebral autoregulation trough a transfer function analysis from a *continuous* recording. This function follows the recommendations from Claassen et al. [1] and mimicks the matlab script created by David Simpsons in 2015 (Matlab TFA function). TFA() also includes the possibility to analyse raw recordings with application of cyclic (beat-to-beat) average with the possibility of utilizing interpolation. (see **details**).

Usage

```
TFA(df, variables,
trigger = NULL, deleter = NULL,
freq = 1000, fast = 50, raw_data = FALSE,
interpolation = 3, output = "table",
vlf = c(0.02,0.07),lf = c(0.07,0.2),
hf = c(0.2,0.5), detrend = FALSE,
spectral_smoothing = 3,
```

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```
coherence2_thresholds = cbind(c(3:15),
c(0.51,0.40,0.34,0.29,0.25,0.22,0.20,0.18,
0.17,0.15,0.14,0.13,0.12)),
apply_coherence2_threshold = TRUE,
remove_negative_phase = TRUE,
remove_negative_phase_f_cutoff = 0.1,
normalize_ABP = FALSE,
normalize_CBFV = FALSE,
window_type = 'hanning',
window_length = 102.4,
overlap = 59.99,
overlap_adjust = TRUE,
na_as_mean = TRUE)
```

Arguments

df Raw continuous recording with numeric data and first column has to be time in

seconds. (dataframe)

variables Definition of the type and order of recorded variables as a list. Middle cerebral

artery blood velocity ('mcav') and arterial blood pressure ('abp') is currently

supported. (list)

trigger Trigger with two columns: first is start, and second is end of period to be an-

alyzed. Every row is a period for analysis. Default is NULL, which results in

analysis of the full dataframe. (dataframe)

deleter Deleter with two columns: first is start and second is end of period with artefacts,

which need to be deleted. Every row is a period with artefacts. Default is NULL.

(dataframe)

freq Frequency of recorded data, in Hz. Default is 1000. (numeric)

fast Select if you want the data to aggregated resulting in a faster, but perhaps more

imprecise run, in Hz. Default is 50 (numeric)

raw_data Select TRUE if the data is raw and cyclic mean should be calculated. **NB:** this

function have not been validated, why validated methods for calculating cyclic mean are preferred. Only 1 period can be analysed using raw_data. Default is

FALSE (boolian)

interpolation Select the number of beats which should be interpolated. Default is up to 3 beats

and 0 results in no interpolation. (numeric)

output Select what the output should be. 'table' results in a dataframe with values

for the three frequencies defined by Claassen et al. [1]; 'long' results in a dataframe with the results in a long format; 'plot' results in a daframe which can help plot gain, phase and coherence; 'plot-peak' results in a dataframe, which can be used to validate the cyclic average, and 'raw' results in a nested

list with results primarily for debugging. Default is 'table'. (string)

vlf, lf, hf, detrend, spectral_smoothing, coherence2_thresholds

See TFA-parameters

apply_coherence2_threshold, remove_negative_phase

See TFA-parameters

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```
remove_negative_phase_f_cutoff, normalize_ABP

See TFA-parameters

normalize_CBFV, window_type, window_length, overlap

See TFA-parameters

overlap_adjust, na_as_mean

See TFA-parameters
```

Details

Using a *continuous* raw recording, TFA() calculates dynamic cerebral autoregulation trough a transfer function analysis. This function utilizes the recommendations from Claassen et al [1] and mimicks the matlab script created by David Simpsons in 2015.

View(data)

time	abp	mcav
7.00	78	45
7.01	78	46
301.82	82	70
301.83	81	69

To calculate the variables insert the data and select the relevant variables.

```
TFA(df=data, variables=c("abp","mcav"))
```

See Value for output description.

Value

TFA() returns a dataframe depending on the output selected. 'table' results in a dataframe with values for the three frequencies defined by Claassen et al. [1]; 'long' results in a dataframe with the results in a long format; 'plot' results in a daframe which can help plot gain, phase and coherence; 'plot-peak' results in a dataframe, which can be used to validate the cyclic average, and 'raw' results in a nested list with results primarily for debugging.

Some generic variables are listed below:

- abp_power The blood pressure power measured in mmHg^2.
- cbfv_power The cerebral blood flow velocity power measured in cm^2*s^-2
- coherence Coherence.
- gain_not_normal Not normalized gain measured in cm*s^-1*mmHg^-1.
- gain_normal Normalized gain measured in %*mmHg^-1.
- phase Phase measured in radians.

output = 'table':

Wide format output table with period, VLF, LF, and HF as columns, and the TFA-variables as rows.

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period	variable	vlf	1f	hf
1	abp_power	6.25	1.56	0.21
1	cbfv_power	3.22	2.25	0.30
3	gain_normal	1.04	1.48	1.85
3	phase	53.0	25.4	9.38

output = 'long':

Long format output table which can be manipulated depending on the intended use, with period, interval, variables and values as columns.

period	interval	variable	values
1	hf	abp_power	6.25
1	hf	cbfv_power	3.22
2	vlf	gain_norm	1.85
2	vlf	phase	9.38

output = 'plot':

Plot format output table which can be used to draw figures with gain, phase and coherence depending on frequency.

period	freq	gain	phase	coherence
1	0.00	0.16	0.00	0.04
1	0.01	0.29	4.22	0.29
2	1.55	1.15	-43.2	0.64
2	1.56	1.16	-41.1	0.42

TFA-paramters

A series of parameters that control TFA analysis (window-length, frequency bands ...). If this is not provided, default values, corresponding to those recommended in the white paper, will be used. These default values are given below for each parameter.

- vlf Limits of *very low frequency* band (in Hz). This corresponds to the matematical inclusion of [X:Y[. Default is c(0.02-0.07).
- If Limits of *low frequency* band (in Hz). This corresponds to the matematical inclusion of [X:Y]. Default is c(0.07-0.2).
- hf Limits of *high frequency* band (in Hz). This corresponds to the matematical inclusion of [X:Y[. Default is c(0.2-0.5).
- detrend Linear detrending of data prior to TFA-analysis (detrending is carried out as one continuous trend over the whole length of the recording, not segment-by-segment). Default is FALSE.

• spectral_smoothing The length, in samples, of the triangular spectral smoothing function. Note that this must be an odd number, to ensure that smoothing is symmetrical around the centre frequency. Default is 3.

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- coherence2_thresholds The critical values (alpha=5%, second column) for coherence for a number of windows (first column, here from 3 to 15). These values were obtained by Monte Carlo simulation, using the default parameter settings for the TFA-analysis (Hanning window, overlap of 50% and 3-point spectral smoothing was assumed). These values should be recalculated for different settings. Note that if overlap_adjust=TRUE, the overlap will vary depending on the length of data. With an overlap of 60% (see below), the critical values increase by between 0.04 (for 3 windows) and 0.02 (for 15 windows). Default is cbind(c(3:15),c(0.51,0.40, 0.34,0.29,0.25,0.22,0.20,0.18,0.17, 0.15,0.14,0.13,0.12)).
- apply_coherence2_threshold Apply the thresholds given above to the TFA-estimates. All
 frequencies with magnitude-squared coherence below the threshold value are excluded from
 averaging when calculating the mean values of gain and phase across the bands. Note that low
 values of coherence are not excluded in the average of coherence across the bands. Default is
 TRUE.
- remove_negative_phase Remove (ignore) negative values of phase in averaging across bands. Negative phase values are removed only for frequencies below the frequency given below, when calculating the average phase in bands. Default is TRUE.
- remove_negative_phase_f_cutoff The cut-off frequency below-which negative phase values are neglected (only if remove_negative_phase is TRUE). Default is 0.1.
- normalize_ABP Normalize ABP by dividing by the mean and multiplying by 100, to express ABP change in %. Note that mean-values are always removed from ABP prior to analysis. Default is FALSE.
- normalize_CBFV Normalize CBFV by dividing by the mean and multiplying by 100, to express CBFV change in %. Note that the band-average values of gain are always calculated both with and without normalization of CBFV, in accordance with the recommendations. Note also that mean-values are always removed from CBFV prior to analysis. Default is FALSE.
- window_type Chose window 'hanning' or 'boxcar'. Default is 'hanning'.
- window_length Length of the data-window, in seconds. Default is 102.4.
- overlap Overlap of the windows, in %. If overlap_adjust is TRUE (see below), then this value may be automatically reduced, to ensure that windows cover the full length of data. Default is 59.99% rather than 60%, so that with data corresponding to 5 windows of 100 s at an overlap of 50%, 5 windows are indeed chosen.
- overlap_adjust Ensure that the full length of data is used (i.e. the last window finishes as near as possible to the end of the recording), by adjusting the overlap up to a maximum value given by params.overlap. Default is TRUE.
- na_as_mean Changes all missing non-interpolated values to the mean value of the corresponding variable. This have not been addressed in the paper by Claassen, and to ensure the dataframes are not 'gathered' this should generate the most stable results. Default is TRUE.

References

1. Claassen et al. (2016) J Cereb Blood Flow Metab. 2016 Apr;36(4):665-80. (PubMed)

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Examples

```
data(tfa_sample_data)
TFA(tfa_sample_data[,c(1:3)], variables=c("abp","mcav"), freq=10)
```

tfa_sample_data

TFA sample data

Description

Dataframe with data provided by Prof. Simpsons, with time (t), arterial blood pressure (abp), left MCAv (mcav_l), right MCAv (mcav_r), and end-tidal CO2 (etco2).

Usage

```
data(tfa_sample_data)
```

Format

An object of class "dataframe"; an example of the usage in TFA-function.

Source

GitHub

References

- Simpsons D (2015) (Cerebral Autoregulation Research Network)
- Claassen et al. (2016) J Cereb Blood Flow Metab. 2016 Apr;36(4):665-80. (PubMed)

```
data(tfa_sample_data)
TFA(tfa_sample_data[,c(1:3)], variables=c("abp","mcav"), freq=10)
```

tfa_sample_data_1

tfa_sample_data_1

TFA sample data - 1

Description

Dataframe with data provided by Prof. Simpsons, with time (t), arterial blood pressure (abp), left MCAv (mcav_l), right MCAv (mcav_r), and end-tidal CO2 (etco2).

Usage

```
data(tfa_sample_data)
```

Format

An object of class "dataframe"; an example of the usage in TFA-function.

Source

GitHub

References

- Simpsons D (2015) (Cerebral Autoregulation Research Network)
- Claassen et al. (2016) J Cereb Blood Flow Metab. 2016 Apr;36(4):665-80. (PubMed)

Examples

```
data(tfa_sample_data)
TFA(tfa_sample_data_1[,c(1:3)], variables=c("abp","mcav"), freq=10)
```

tfa_sample_data_2

TFA sample data - 2

Description

Dataframe with data provided by Prof. Simpsons, with time (t), arterial blood pressure (abp), left MCAv (mcav_l), right MCAv (mcav_r), and end-tidal CO2 (etco2).

Usage

```
data(tfa_sample_data)
```

Format

An object of class "dataframe"; an example of the usage in TFA-function.

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Source

GitHub

References

- Simpsons D (2015) (Cerebral Autoregulation Research Network)
- Claassen et al. (2016) J Cereb Blood Flow Metab. 2016 Apr;36(4):665-80. (PubMed)

```
data(tfa_sample_data)
TFA(tfa_sample_data_2[,c(1:3)], variables=c("abp","mcav"), freq=10)
```

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