Package: GENEA classify (via r-universe)

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Type Package

Title Segmentation and Classification of Accelerometer Data

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Author Chris Campbell [aut], Aimee Gott [aut], Joss Langford [aut], Charles Sweetland [aut], Penny Sweetland [aut], Jia Ying Chua [aut, cre], Activinsights Ltd [cph]

Description Segmentation and classification procedures for data from the 'Activinsights GENEActiv'

https://activinsights.com/technology/geneactiv/ accelerometer that provides the user with a model to guess behaviour from test data where behaviour is missing. Includes a step counting algorithm, a function to create segmented data with custom features and a function to use recursive partitioning provided in the function rpart() of the 'rpart' package to create classification models.

License GPL

LazyData true

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Maintainer Jia Ying Chua <jiayingc@activinsights.com>

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GENEAclassify-package Classification of accelorometer data

Description

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This package provides tools to perform the segmentation and classification of GENEActiv accelerometer data. The high frequency time series data is split into segments based on activity changepoints. An **rpart** fit is trained against known activities at each segment. This fit can then then be used to guess behaviours from test data when activity at each time point has not been reported. This allows detailed behaviour profiles to be created for the wearer.

classifyGENEA

Classify Data into Categories defined in an rpart GENEA fit

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Description

Perform classification on segmented GENEActiv bin data using an rpart GENEA training fit.

Usage

```
classifyGENEA(
  testfile,
  start = NULL,
  end = NULL,
  Use.Timestamps = FALSE,
  radians = FALSE,
  mmap.load = (.Machine$sizeof.pointer >= 8),
  trainingfit = trainingFit,
  newdata,
  allprobs = FALSE,
  setinf = 100,
```

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```
outputname = "_classified",
  outputdir = "GENEAclassification",
  datacols = "default",
 changepoint = c("UpDownDegrees", "TempFreq", "UpDownFreq", "UpDownMean", "UpDownVar",
    "UpDownMeanVar", "DegreesMean", "DegreesVar", "DegreesMeanVar",
    "UpDownMeanVarDegreesMeanVar", "UpDownMeanVarMagMeanVar"),
  penalty = "Manual",
  pen.value1 = 40,
  pen.value2 = 400,
  intervalseconds = 30,
 mininterval = 5,
  samplefreq = 100,
  filterorder = 2,
  boundaries = c(0.5, 5),
  Rp = 3,
  plot.it = FALSE,
  hysteresis = 0.1,
  stft_win = 10,
  plot.seg = FALSE,
  plot.seg.outputfile = "Changepoint",
  verbose = TRUE,
)
```

Arguments

testfile character string stating path to a GENEActiv bin file, or a folder containing

GENEActiv bin files.

start Where to start reading observations.
end Where to end reading observations.

Use.Timestamps To use timestamps as the start and end time values this has to be set to TRUE.

(Default FALSE)

radians calculate degrees rotation in radians.

mmap.load Default is (.Machine\sizeof.pointer >= 8). see mmap for more details

trainingfit a GENEA rpart object created by createGENEAmodel that gives the decision

tree that was fitted from the training data. These are the parameters used to

predict the new data.

newdata a new data frame that is to be classified (provide instead of testfile). The data

must contain the features named in the trainingfit.

allprobs single logical should all estimated probabilities be reported rather than proba-

bility of selected class (default FALSE).

setinf single numeric an arbitrary value to replace Inf in calculated columns or NA to

ignore Inf values. (default 100). -setinf is used to replace -Inf. Alternatively, use

setinf NULL to leave Inf as is.

outputname file name root (excluding extension) for saving the classification output (default

"classified").

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absolute or relative path to directory in which artifacts (plot and changes files) outputdir should be created or NULL (default "GENEAclassification"). datacols a vector constructed 'column.summary' or 'default'. See segmentation for details. changepoint defines the change point analysis to use. UpDownDegrees performs the change point analysis on the variance of arm elevation and wrist rotation. TempFreq performs a change point on the variance in the temeprature and Frequency (Typically better for sleep behaviours) penalty single characgter, the penalty to use for changepoint detection. default ("SIC") pen.value1 Value of the type 1 error required when penalty is "Asymptotic". Default set as NULL and so equals pen.value1 if no input. pen.value2 intervalseconds An integer number of seconds between 5 and 30 during which at most one changepoint may occur. mininterval single numeric that defines the smallest changepoint initially found. Passed to cpt.var as the variable minseglen samplefreq The sampling frequency of the data, in hertz, when calculating the step number. (default 100) filterorder The order of the filter applied with respect to the cheby options. boundaries to passed to the filter in the step counting algorithm. Rр the decibel level that the cheby filter takes. see cheby1 plot.it (logical) Creates a plot showing the zero crossings counted by the step counting algorithm#' @param Centre Centres the xz signal about 0 when set to True. hysteresis The hysteresis applied after zero crossing. (default 100mg) numeric for the window to calculate the frequency of an event using the stft stft_win function. plot.seg (logical) Creates a plot displaying the changepoint locations plot.seg.outputfile The name of the png file created that shows the change points on a positionals single logical should additional progress reporting be printed at the console (deverbose fault TRUE). other arguments to be passed to dataImport, segmentation and other functions

Details

This function will apply the rules determined by the rpart GENEA decision tree passed to argument trainingfit to the columns of newdata to classify into classes (view using "levels").

Value

The function will return the data frame that was provided as newdata with additional columns.

1. Class, a factor indicating that the predicted category of the segment

createGENEAmodel 5

2. p.Class, estimated probability that the prediction is correct

Alternatively, by setting argument allprobs to TRUE, a column constructed 'p.level' containing the estimated probability of each possible class will be returned instead.

Examples

```
## segData <- read.csv(system.file(package = "GENEAclassify",
## "testdata", "trainingData9.csv"))
## The training fit can be created by provided the file path to the training data
## in the function getTrainingData - see the help file for more details
## Uses the fitted decision tree to predict the segmented data
## class9 <- classifyGENEA(testfile = "trainingData9.csv",
## newdata = segData,
## trainingfit = trainingFit)
## head(class9)
## table(class9$Class)</pre>
```

createGENEAmodel

Create training data decision tree model

Description

From data frame create a decision tree that can be used for classifying data into specified categories. The data frame may optionally contain a reserved column Source, specifying the provenance of the record. The data frame must contain a column, by default named Activity, specifing the classes into which the model fit should be classified.

Usage

```
createGENEAmodel(
  data,
  outputtree = NULL,
  features = c("Segment.Duration", "Principal.Frequency.mad", "UpDown.sd", "Degrees.sd"),
  category = "Activity",
  plot = TRUE,
  verbose = TRUE,
  ...
)
```

Arguments

data data frame containing segmented GENEActiv bin data.

outputtree name of the png file that shows the classification tree plot.

features character vector naming independent variables to use in classification. Alternatively, a numeric vector specifying the variables to pass to the classification or

tively, a numeric vector specifying the variables to pass to the classification or NULL, in which case all variables are used in the order of the supplied training dataset. Note that including large numbers of variables (>7) may result in long

run times.

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category	single character naming the dependent variable to use (default 'Activity').
plot	a logical value indicating whether a plot of the classification tree should be plotted. The default is TRUE.
verbose	single logical should additional progress reporting be printed at the console? (default $TRUE$)
	other arguments for rpart

Details

The function will create an rpart classification tree for the training data based upon the parameters passed to features. The model created, an GENEA rpart object can be used within the function "classifyGENEA" to classify GENEA bin files.

Value

A GENEA rpart fit.

See Also

The returned object can be interrogated with features, the variables used in defining the model, and "levels", the response categories predicted by the model.

Examples

```
## dataPath <- file.path(system.file(package = "GENEAclassify"),</pre>
##
                                       "testdata",
##
                                       "trainingData9.csv")
##
## t1 <- read.csv(file = dataPath)
## f1 <- createGENEAmodel(data = t1,
##
                           features = c("Degrees.var",
##
                                         "UpDown.mad",
                                         "Magnitude.mean"),
##
##
                           category = "Activity")
##
## class(f1)
## levels(f1)
## features(f1)
## plot(f1)
## text(f1)
```

dataImport

Data import function

Description

Loads the data into R and creates format required for segmentation.

dataImport 7

Usage

```
dataImport(
  binfile,
  downsample = 100,
  start = NULL,
  end = NULL,
  Use.Timestamps = FALSE,
  radians = FALSE,
  keep_raw_data = TRUE,
  mmap.load = (.Machine$sizeof.pointer >= 8),
  ...
)
```

Arguments

binfile File path to binary data to be segmented.

downsample Rate to downsample the data, defaults to every 100th observation. For no down-

sampling set NULL.

start Where to start reading observations.
end Where to end reading observations.

Use. Timestamps To use timestamps as the startand end time values this has to be set to TRUE.

(Default FALSE)

radians calculate degrees rotation in radians.

keep_raw_data Keep the raw data for calculating steps using stepcounter.

mmap.load Default is (.Machine\sizeof.pointer >= 8). see mmap for more details

... additional arguments passed through.

Details

Reads in the binary data file and extracts the information required for the segmentation procedure.

Value

Returns a list containing a matrix of the data including the x, y and z axis data, vectors of the up down (elevation) and degrees (rotation), a vector of time stamps, a vector of vector magnitudes and the serial number of the device.

```
## segData = dataImport(bindata = file.path(system.file(package = "GENEAread"),
## "binfile",
## "TESTfile.bin"))
##
segData1 = dataImport(AccData)
## names(segData)
```

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find_peaks

Find_Peaks

Description

Peak detect function

Usage

```
find_peaks(x, m = 3)
```

Arguments

x timeseries signal

The number of points either side of the peak to required to be a peak.

Details

Finds the peaks and valleys within the signal passed to the function.

getGENEAsegments

import and segment one or more bin files.

Description

Import and summarize GENEActiv bin data for manual classification.

Usage

```
getGENEAsegments(
  testfile,
  start = NULL,
  end = NULL,
  Use.Timestamps = FALSE,
  radians = FALSE,
  keep_raw_data = TRUE,
  mmap.load = (.Machine$sizeof.pointer >= 8),
  outputtoken = "_segmented",
  outputtdir = "GENEAclassification",
  datacols = "default",
  decimalplaces = "default",
  filterWave = FALSE,
  filtername = "haar",
  j = 8,
  changepoint = c("UpDownDegrees", "TempFreq", "UpDownFreq", "UpDownMean", "UpDownVar",
```

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```
"UpDownMeanVar", "DegreesMean", "DegreesVar", "DegreesMeanVar",
    "UpDownMeanVarDegreesMeanVar", "UpDownMeanVarMagMeanVar"),
  penalty = "Manual",
  pen.value1 = 40,
  pen.value2 = 400,
  intervalseconds = 30,
 mininterval = 5,
  samplefreq = 100,
  filterorder = 2,
  boundaries = c(0.5, 5),
  Rp = 3,
  plot.it = FALSE,
  hysteresis = 0.1,
  stft_win = 10,
  plot_changepoints = FALSE,
  plot_changepoints_outputfile = "Changepoint",
  verbose = FALSE,
)
```

Arguments

testfile character vector stating path to a GENEActiv bin file, or a folder containing

GENEActiv bin files.

start Where to start reading observations.
end Where to end reading observations.

Use. Timestamps To use timestamps as the start and end time values this has to be set to TRUE.

(Default FALSE)

radians calculate degrees rotation in radians.

keep_raw_data Keep the raw data for calculating steps using stepcounter.

mmap.load Default is (.Machine\sizeof.pointer >= 8). see mmap for more details

outputtoken single character string to be appended to the file name for saving the segmena-

tion output (default '_segmentated').

outputdir The absolute or relative path to directory in which artifacts (plot and changes

files) should be created, or NULL (default "GENEAclassification").

datacols a vector constructed 'column.summary' or 'default'. See segmentation for

details.

decimalplaces named numeric vector of decimal places with which to round summary columns.

NULL returns unrounded values. The length 1 character vector 'default' applies default roundings:

raan rounamgs.

- Start.Time = 0,
- Degrees.mean = 3,
- Degrees.median = 3,
- Degrees.var = 3,
- Degrees.sd = 3,

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- Degrees.mad = 3,
- Magnitude.mean = 3,
- UpDown.mean = 3,
- UpDown.median = 3,
- UpDown.var = 3
- UpDown.sd = 3,
- UpDown.mad = 3,
- Principal.Frequency.median = 3,
- Principal.Frequency.mad = 3,
- Principal.Frequency.ratio = 3,
- Principal.Frequency.sumdiff = 3,
- Principal.Frequency.meandiff = 3,
- Principal.Frequency.abssumdiff = 3,
- Principal.Frequency.sddiff = 3,
- Light.mean = 0,
- Light.max = 0,
- Temp.mean = 1,
- Temp.sumdiff = 3
- Temp.meandiff = 3
- Temp.abssumdiff = 3
- Temp.sddiff = 3
- Step.count = 0
- Step.sd = 1
- Step.mean = 0
- Step.GENEAamplitude = 3
- Step.GENEAwavelength = 3
- Step.GENEAdistance = 3

This can be changed by using a named list. e.g decimalplaces = c(Start.Time = 2, Degrees.mean = 4).

filterWave

single logical, should a smoothing filter from wave.filter be applied? (default FALSE).

filtername

single character, the name of the wavelet to use for smoothing when filter is TRUE. (default "haar") Passed to link[waveslim]{wave.filter}.

j

single numeric, the level to which to smooth. Passed to link[waveslim]{wave.filter} (default 8).

changepoint

defines the change point analysis to use. UpDownDegrees performs the change point analysis on the variance of arm elevation and wrist rotation. TempFreq performs a change point on the variance in the temperature and frequency (Typically better for sleep behaviours).

penalty

single character, the penalty to use for changepoint detection. default ("SIC").

pen.value1

Value of the type 1 error required when penalty is "Asymptotic".

pen.value2

Default set as NULL and so equals pen.value1 if no input.

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An integer number of seconds between 5 and 30 during which at most one changepoint may occur.

mininterval single numeric that defines the smallest changepoint initially found. Passed to

cpt.var as the variable minseglen

samplefreq The sampling frequency of the data, in hertz, when calculating the step number.

(default 100).

filterorder The order of the filter applied with respect to the butter or cheby options. See

cheby1 or butter.

boundaries to pass to the filter in the step counting algorithm.

Rp the decibel level that the cheby filter takes. See cheby1.

plot.it single logical, Creates a plot showing the zero crossings counted by the step

counting algorithm#' @param Centre Centres the xz signal about 0 when set to

True.

hysteresis The hysteresis applied after zero crossing. (default 100mg)

stft_win numeric for the window to calculate the frequency of an event using the stft

function.

plot_changepoints

single logical, Creates a plot displaying the changepoint locations.

 $\verb|plot_changepoints_outputfile||$

The name of the png file created that shows the change points on a positionals

plots.

verbose single logical should additional progress reporting be printed at the console?

(default TRUE).

.. other arguments to be passed to dataImport, segmentation and other functions

with these functions.

Value

segmented data are returned

See Also

The returned object can be interrogated with head.

```
## testfile = file.path(system.file(package = "GENEAread"),
##
                                     "binfile",
                                     "TESTfile.bin")
##
##
## segData <- getGENEAsegments(testfile = testfile,</pre>
##
                                outputdir = file.path(tempdir(), "GENEAclassification"),
##
                                changepoint = "UpDownDegrees",
##
                                pen.value1 = 1,
                                pen.value2 = 1)
##
## head(segData)
## list.files(file.path(tempdir(), "GENEAclassification"))
```

segmentation

Perform Segmentation on GENEActiv Accelerometer Data

Description

Perform segmentation of Activinsights accelerometer data. Data are smoothed by the second, or by 10 data points, whichever number of records is greater.

Filtering is performed by tools from waveslim. Options are passed to wavelet.filter.

Usage

```
segmentation(
        data,
        outputfile = "detectedChanges",
        outputdir = "GENEAclassification",
        datacols = "default",
        decimalplaces = "default",
        filterWave = FALSE,
        filtername = "haar",
        j = 8,
     changepoint = c("UpDownDegrees", "TempFreq", "UpDownFreq", "UpDownMean", "UpDownVar",
                \label{thm:continuous} \begin{tabular}{ll} "UpDownMeanVar", "DegreesMeanVar", "DegreesMeanVar", "UpDownMeanVarMagMeanVar", "RadiansMean", "
                "RadiansVar", "RadiansMeanVar", "UpDownMeanDegreesVar"),
        penalty = "Manual",
        pen.value1 = 40,
        pen.value2 = 400,
        intervalseconds = 30,
       mininterval = 5,
        samplefreq = 100,
        filterorder = 2,
        boundaries = c(0.5, 5),
        Rp = 3,
        plot.it = FALSE,
        hysteresis = 0.1,
        stft_win = 10,
        verbose = FALSE,
        verbose_timer = FALSE,
)
```

Arguments

the GENEActiv bin object to be segmented which should be the output of the data dataImport function.

outputfile single character, file name for saving the segmentation output as CSV (and if plot.it is TRUE, corresponding plot PNG). If NULL, create no files.

outputdir

single character, the absolute or relative path to directory in which plot and changes files) should be created, or NULL (default "GENEAclassification"). Ignored if outputfile is NULL.

datacols

character vector constructed 'column.summary'. This object specifies the data and summary to output for the classification. The first part of each element must name column in the GENEAbin datasets specified by filePath. Derived columns may also be selected:

- Step (zero-crossing step counter method),
- Principal.Frequency.

The second should be the name of a function that evaluates to lenth one. The functions must contain only alphabetical characters (no numbers, underscores or punctuation). The default matrix, specified using the length 1 character vector 'default' is:

- UpDown.mean
- UpDown.sd
- UpDown.mad
- · Degrees.mean
- · Degrees.var
- · Degrees.sd
- Light.mean
- Light.max
- Temp.mean
- · Temp.sumdiff
- · Temp.meandiff
- · Temp.abssumdiff
- · Temp.sddiff
- Magnitude.mean
- Step.GENEAcount
- Step.sd
- Step.mean
- Step.GENEAamplitude
- Step.GENEAwavelength
- Principal.Frequency.median
- · Principal.Frequency.mad
- Principal.Frequency.ratio
- · Principal.Frequency.sumdiff
- · Principal.Frequency.meandiff
- Principal.Frequency.abssumdiff
- · Principal.Frequency.sddiff

decimalplaces

named numeric vector of decimal places with which to round summary columns. NULL returns unrounded values. The length 1 character vector 'default' applies default roundings:

• Start.Time = 0,

```
• Degrees.mean = 3,
                    • Degrees.median = 3,
                    • Degrees.var = 3,
                    • Degrees.sd = 3,
                    • Degrees.mad = 3,
                    • Magnitude.mean = 3,
                    • UpDown.mean = 3,
                    • UpDown.median = 3,
                    • UpDown.var = 3
                    • UpDown.sd = 3,
                    • UpDown.mad = 3,
                    • Principal.Frequency.median = 3,
                    • Principal.Frequency.mad = 3,
                    • Principal.Frequency.ratio = 3,
                    • Principal.Frequency.sumdiff = 3,
                    • Principal.Frequency.meandiff = 3,
                    • Principal.Frequency.abssumdiff = 3,
                    • Principal.Frequency.sddiff = 3,
                    • Light.mean = 0,
                    • Light.max = 0,
                    • Temp.mean = 1,
                    • Temp.sumdiff = 3
                    • Temp.meandiff = 3
                    • Temp.abssumdiff = 3
                    • Temp.sddiff = 3
                    • Step.GENEAcount = 0
                    • Step.sd = 1
                    • Step.mean = 0
filterWave
                  single logical, should a smoothing filter from wave.filter be applied? (default
                  FALSE).
filtername
                  single character, the name of the wavelet to use for smoothing when filter is
                  TRUE. (default "haar") Passed to link[waveslim] {wave.filter}.
                  single numeric, the level to which to smooth. Passed to link[waveslim]{wave.filter}
                  (default 8).
changepoint
                  defines the change point analysis to use. UpDownDegrees performs the change
                  point analysis on the variance of arm elevation and wrist rotation. TempFreq
                  performs a change point on the variance in the temeprature and frequency (Typ-
                  ically better for sleep behaviours).
```

single characgter, the penalty to use for changepoint detection. default ("SIC"). penalty

Value of the type 1 error required when penalty is "Asymptotic". pen.value1

pen.value2 Default set as NULL and so equals pen.value1 if no input.

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An integer number of seconds between 5 and 30 during which at most one

changepoint may occur.

mininterval single numeric that defines the smallest changepoint initially found. Passed to

cpt.var as the variable minseglen

samplefreq The sampling frequency of the data, in hertz, when calculating the step number.

(default 100).

filterorder The order of the filter applied with respect to the butter or cheby options if

stepCounter is used. The order of the moving average filter if step counter 2 is

used. See cheby1 or butter.

boundaries to pass to the filter in the step counting algorithm.

Rp the decibel level that the cheby filter takes. See cheby1.

plot.it single logical, Creates a plot showing the zero crossings counted by the step

counting algorithm#' @param Centre Centres the xz signal about 0 when set to

True.

hysteresis The hysteresis applied after zero crossing. (default 100mg)

stft_win numeric for the window to calculate the frequency of an event using the stft

function.

verbose single logical to print additional progress reporting (default FALSE).

verbose_timer single logical tp print additional progress reporting on time for each section of

the function (default FALSE).

.. other arguments to be passed to dataImport, segmentation and other functions

with these functions.

Details

Performs the segmentation procedure on the provided elevation data. Optionally a wavelet filter is first applied to smooth the data. The number of changes occuring in a given number of seconds may be controlled using the intervalseconds argument. Changes will be removed based on which segments are the closest match in terms of variance. A series of features for each of the segments will then be calculated and returned as a csy file.

Value

The segment data are returned invisibly. This data frame has columns:

- · Serial.Number
- Start.Time
- Segment.Start.Time
- · Segment.Duration
- UpDown.median
- UpDown.var
- · Degrees.median
- Degrees.mad

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In addition, the requested columns are included. Optionally, as a side effect a csv file is returned listing all of the segments found in the data along with a variety of features for that segment. Optionally a png file plotting the data and the detected changes can also be produced.

Examples

```
### Load the data to segment keeping only the first quarter of the data
## library(GENEAread)
## testfile = file.path(system.file(package = "GENEAread"),
                                      "binfile",
##
                                      "TESTfile.bin")
##
## segData <- dataImport(binfile = testfile,</pre>
       downsample = 100, start = 0, end = 0.25)
## head(segData)
### Run loaded data through segmentation function
## segment <- segmentation(data = segData, outputfile = NULL)</pre>
## head(segment)
## segment2 <- segmentation(data = segData, outputfile = NULL,</pre>
       datacols = "Degrees.skew")
## head(segment2)
```

stepCounter

Step Counter

Description

Function to calculate the number and variance of the steps in the data.

Usage

```
stepCounter(
  AccData,
  samplefreq = 100,
  filterorder = 2,
  boundaries = c(0.5, 5),
  Rp = 3,
  plot.it = FALSE,
  hysteresis = 0.05,
  verbose = verbose,
  fun = c("GENEAcount", "mean", "sd", "mad")
)
```

Arguments

or a vector.

samplefreq The sampling frequency of the data, in hertz, when calculating the step number

(default 100).

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filterorder single integer, order of the Chebyshev bandpass filter, passed to argument n of cheby1. length 2 numeric vector specifying lower and upper bounds of Chebychev filter boundaries (default c(0.5, 5) Hz), passed to argument W of butter or cheby1. Rр the decibel level that the cheby filter takes, see cheby1. single logical create plot of data and zero crossing points (default FALSE). plot.it hysteresis The hysteresis applied after zero crossing. (default 100mg) verbose single logical should additional progress reporting be printed at the console?

(default FALSE).

fun character vector naming functions by which to summarize steps. "count" is an

internally implemented summarizing function that returns step count.

Value

Returns a vector with length fun.

Examples

```
d1 <- \sin(\sec(0.1, 100, 0.1))/2 + rnorm(1000)/10 + 1
Steps4 = stepCounter(d1)
length(Steps4)
mean(Steps4)
sd(Steps4)
plot(Steps4)
```

TrainingData

Example Training Data set

Description

A manually classified training data set provided with the package.

Format

Manually classified segmented data that can be used to build a classification model.

Source

Manually classified by Actvinsights.

See Also

```
"levels" features
```

```
data(TrainingData)
class(TrainingData)
```

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 ${\it trainingFit}$

Example classification tree

Description

rpart object declaring the decision tree for the classification of GENEActiv bin data into typical groups.

Format

An rpart object with class GENEA containing the decision tree information required for prediction.

Source

Output of createGENEAmodel on experimental training data.

See Also

```
"levels" features
```

```
data(trainingFit)
class(trainingFit)
levels(trainingFit)
features(trainingFit)
plot(trainingFit)
text(trainingFit, cex = 0.5)
```

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