Package: DecomposeR (via r-universe)

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Type Package

Title Empirical Mode Decomposition for Cyclostratigraphy

Version 1.0.6

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Description Tools to apply Ensemble Empirical Mode Decomposition (EEMD) for cyclostratigraphy purposes. Mainly: a new algorithm, extricate, that performs EEMD in seconds, a linear interpolation algorithm using the greatest rational common divisor of depth or time, different algorithms to compute instantaneous amplitude, frequency and ratios of frequencies, and functions to verify and visualise the outputs. The functions were developed during the CRASH project (Checking the Reproducibility of Astrochronology in the Hauterivian). When using for publication please cite Wouters, S., Crucifix, M., Sinnesael, M., Da Silva, A.C., Zeeden, C., Zivanovic, M., Boulvain, F., Devleeschouwer, X., 2022, ``A decomposition approach to cyclostratigraphic signal processing". Earth-Science Reviews 225 (103894).

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License GPL-3

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Suggests EMD, Rssa, astrochron, tidyverse

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| approx.cor | Correlation of time-series with different sampling rate | |
|------------|---|--|
| | | |

Description

Allows to correlate time-series having different sampling rate, if they have a comparable depth or time scale

Usage

```
approx.cor(xy1, dt1, xy2, dt2, plot = T, output = T, type = "p", ...)
```

Arguments

| xy1 | intensity values for the first data set |
|--------|---|
| dt1 | depth or time scale for the first data set |
| xy2 | intensity values for the second data set |
| dt2 | depth or time scale for the second data set |
| plot | whether to plot |
| output | whether to output |
| type | type of points in the plot (see help page of lines() for details) |
| | additional parameters to feed to the lines() function |

Value

a list of correlation (\$cor), slope (\$slope), intercept (\$intercept) (two values for each: interpolation to fit dt1 and dt2 respectively), and of the xy1 and xy2 values, interpolated for dt1 (\$df1) and df2 (\$df2)

```
set.seed(42)
n <- 600
t <- seq_len(n)

p1 <- 30
p2 <- 240

xy.pure <- (1 + 0.6 * sin(t*2*pi/p2)) * sin(t*2*pi/p1) + 2 * sin(t*2*pi/p2)

xy <- xy.pure + rnorm(n, sd = 0.5)

inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
```

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```
dt.pure <- cumsum(inter_dt)
keep <- runif(length(dt.pure)) < 0.5

xy <- xy[keep]
dt <- dt.pure[keep] + rnorm(sum(keep), -0.2, 0.2)

par(mfrow = c(1,2))

plot(xy, dt, type = "o", pch = 19)

plot(xy.pure, dt.pure, type = "o", pch = 19)

par(mfrow = c(1,1))

out <- approx.cor(xy, dt, xy.pure, dt.pure)

out$cor
out$cor
out$slope
out$intercept</pre>
```

as.emd

Create / Check emd objects

Description

Allows to convert the result of a decomposition into a standard list. The warnings of the is.emd checking function allow to identify the problems.

Usage

```
as.emd(
   xy,
   dt,
   imf,
   residue = NULL,
   ini = NULL,
   mode = NULL,
   repl = 1,
   order = NA
)
is.emd(emd)
```

Arguments

xy a vector of length n for the original signal at each dt dt a vector of length n for the depth or time reference

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| imf | a data.frame or matrix of n rows of the IMFs |
|---------|--|
| residue | a vector of length n for the residue of the decomposition |
| ini | an optional vector of length n of the eventual initial Intrinsic Mode Function xy would be a demodulation of, if it is a demodulation. |
| mode | the mode sequence index to give to each replicated IMFs |
| repl | the id of each replicates. The length of unique(repl) defines the amount of replicates. |
| order | the order of the imf, typically from higher frequency to lower frequency |
| emd | an emd object to check |

Value

a list made of \$xy (original signal), \$dt (depth/time), \$m (a matrix of the decomposition), \$repl (the replicate id of each point) and \$mode (the mode id of each point).

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
s30 \leftarrow (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1)
s240 \leftarrow 2 * sin(t*2*pi/p2)
sn <- rnorm(n, sd = 0.5)
xy < -s30 + s240 + sn
inter_dt \leftarrow round(runif(length(xy), min = 0.5, max = 1.5),1)
dt <- cumsum(inter_dt)</pre>
dec \leftarrow as.emd(xy = xy, dt = dt, imf = matrix(c(sn, s30, s240), ncol = 3))
plot_emd(dec, pdf = FALSE)
is.emd(dec)
## Not run:
dec$xy <- 1
is.emd(dec)
## End(Not run)
```

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| | 7 | |
|-----|-----|----|
| as. | puı | se |

Create / Check pulse objects

Description

Allows to convert instantaneous frequency determination results into a single 'pulse' object. This is the format generated by inst.pulse (and gzc if output = 2)

Usage

```
as.pulse(
  dt,
  f,
  a = NULL,
  m = NULL,
  idt = NULL,
  mode = NULL,
  repl = 1,
  order = NA
)
is.pulse(pulse)
```

Arguments

| dt | a vector of length n for the depth or time reference |
|-------|---|
| f | a data.frame or matrix of n rows of the instantaneous frequencies |
| а | a data.frame or matrix of n rows of the instantaneous amplitudes |
| m | a data.frame or matrix of n rows of the components from which the frequencies and amplitudes were computed from |
| idt | data.frame or matrix of n rows of identity tuning: new dt coordinates to remove the frequency modulation |
| mode | the mode sequence index to give to each replicated IMFs |
| repl | a vector for the number of replicates or a matrix, indicating in which replicate set each point is |
| order | the order of the imf, typically from higher frequency to lower frequency |
| pulse | a pulse object to check |

Value

a list made of \$dt (depth/time), \$f (instantaneous frequency), \$a (instantaneous amplitude) if a is provided, \$repl (the replicate id of each point) and \$mode (the mode id of each point).

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Examples

```
set.seed(42)
n <- 600
dt <- seq_len(n)</pre>
p1 <- 30
p2 <- 240
s30 < (1 + 0.6 * sin(dt*2*pi/p2)) * sin(dt*2*pi/p1)
s240 \leftarrow 2 * sin(dt*2*pi/p2)
xy < - s30 + s240
dec \leftarrow as.emd(xy = xy, dt = dt, imf = matrix(c(s30, s240), ncol = 2))
plot_emd(dec, pdf = FALSE, style = 1)
pulse <- inst.pulse(dec, last = TRUE, breaks = 200, bins = 40, cut = 10)</pre>
is.pulse(pulse)
simp.pulse <- as.pulse(pulse$dt, pulse$f)</pre>
str(simp.pulse)
```

check.emd

Check an EMD object

Description

Provides an ensemble of check on the quality of a decomposition presented as an emd object (see as.emd for more information)

Usage

```
check.emd(emd, xy = NULL, timelimit = 15)
```

Arguments

| emd | an amd object to test |
|-----------|---|
| ху | the original signal that was decomposed: this parameter is simply to insure that you are indeed comparing the decomposition to the original signal, and not cheating by providing the sum of your decomposition |
| timelimit | a time limit for the computation of the greatest common rational divisor. A too long time may be indicative of a problem, typically depth/time values that are |

not rounded adequately.

8 condense

Examples

condense

Condenses columns of matrix

Description

Condenses columns of a matrix by averaging or summing them. The condensing can be done partially: a multiple of the repetitions can be averaged or summed to keep some repetitions.

Usage

```
condense(m, n, fun = "mean")
```

Arguments

m matrix of repeated signal, each column being a repetition
 n the number of repetitions that will be averaged/summed
 fun the function to apply to each repetition: "mean" or "sum".

Value

a matrix with n times less columns

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Examples

```
m <- matrix(rep(seq(100, 800, 100), each = 10) + rep(1:10, 8), ncol = 8)
m
condense(m, 4)</pre>
```

DecomposeR

DecomposeR: Empirical Mode Decomposition for Cyclostratigraphy

Description

This package provides tools to apply Ensemble Empirical Mode Decomposition (EEMD) for cyclostratigraphy purposes. It proposes a new algorithm, that performs EEMD in seconds, a linear interpolation algorithm using the greatest rational common divisor of depth or time, different algorithms to compute instantaneous amplitude, frequency and ratios of frequencies, and functions to verify and visualise the outputs.

Details

Package: DecomposeR

Type: R package

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License: GPL-3

Note

If you want to use this package for publication or research purposes, please cite Wouters, S., Crucifix, M., Sinnesael, M., Da Silva, A.C., Zeeden, C., Zivanovic, M., Boulvain, F., Devleeschouwer, X., 2022, "A decomposition approach to cyclostratigraphic signal processing". Earth-Science Reviews 225 (103894). <doi:10.1016/j.earscirev.2021.103894>.

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DecomposeR.Datasets

Datasets for Testing DecomposeR

Description

Datasets for testing DecomposeR: the ace dataset is from from Sinnesael et al. (2016), the cip2 and cip3 data sets are from the signals 2 and 3 of the CIP project (Sinnesael et al., 2019), respectively, and cip1 was derived from cip1_raw which is a rasterisation of the .tif image provided as signal 1 of the CIP project. A real case study is also provided, out of ODP 926 in Ceara Rise, limited between 5 & 9 Millions of years ago (Ma): the data sets z13 and z13amp are from Zeeden et al., 2013, and are respectively the greyscale, and its amplitude modulation for the eccentricity; w17 is from Wilkens et al., 2017, which proposes a revised splice for magnetic susceptibility; sc97amp is the amplitude modulation of eccentricity as it was calculated on the magnetic susceptibility by Shackleton & Crowhurst (1997). Excerpts from the Laskar et al., 2004 solution are further provided from http://vo.imcce.fr/insola/earth/online/earth/online/index.php: they are the insolation input for the CIP1 signal (cip1_imput), and various solutions for precession, eccentricity and obliquity for given time intervals (in millions of years ago): La04_pre_0_20, La04_ecc_6_8, La04_obl_6_8 & La04_pre_obl_5_9.

Details

xy Values of the signal

pre Values of the signal

dt Depth or time of the signal

age Tuned age of the signal

References

Laskar, J., Robutel, P., Joutel, F., Gastineau, M. Correia, A. C. M., & Levrard, B. (2004). A long-term numerical solution for the insolation of the Earth. Astronomy & Astrophysics. 428. 261-285. doi:10.1051/00046361:20041335

Shackleton, N. J., & Crowhurst, S. (1997). Sediment fluxes based on an orbitally tuned time scale 5 Ma to 14 Ma, site 926. Proceedings of the Ocean Drilling Program, Scientific Results. 154. doi:10.2973/odp.proc.sr.154.102.1997

Sinnesael, M., Zivanovic, M., De Vleeschouwer, D., Claeys, P. & Schoukens, J. (2016). Astronomical component estimation (ACE v.1) by time-variant sinusoidal modeling. Geoscientific Model Development. 9. 3517-3531. doi:10.5194/gmd935172016

Sinnesael, M., De Vleeschouwer, D., Zeeden, C., et al. (2019). The Cyclostratigraphy Intercomparison Project (CIP): consistency, merits and pitfalls. Earth-Science Reviews. 199. 102965. doi:10.1016/j.earscirev.2019.102965

Wilkens, R. H., Westerhold, T., Drury A. D., Lyle, M., Gorgas, T., Tian, J. (2017). Revisiting the Ceara Rise, equatorial Atlantic Ocean: isotope stratigraphy of ODP Leg 154 from 0 to 5Ma. Climate of the Past. 13. 779-793. doi:10.5194/cp137792017

Zeeden, C., Hilgen, F., Westerhold, T., Lourens, L., Röhl, U. & Bickert, T. (2013). Revised Miocene splice, astronomical tuning and calcareous plankton biochronology of ODP Site 926 between 5

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and 14.4 Ma. Palaeogeography, Palaeoclimatology, Palaeoecology. 369. 430–451. doi:10.1016/j.palaeo.2012.11.009

Description

Calculates instantaneous frequency of frequency carriers using the direct quadrature method from Huang et al., 2009.

Usage

```
dq.algorithm(fc, dt)
```

Arguments

fc a matrix of amplitude between -1 and 1, making up the frequency carrier dt a vector of depth or time values

Value

a list of the depth/time (dt), frequency (f), and identity tuning (idt), i.e. depths adapted to transform the frequency carrier into a cosine of period 1.

References

Huang, Norden E., Zhaohua Wu, Steven R. Long, Kenneth C. Arnold, Xianyao Chen, and Karin Blank. 2009. "On Instantaneous Frequency". Advances in Adaptive Data Analysis 01 (02): 177–229. https://doi.org/10.1142/S1793536909000096.

```
n <- 600
t <- seq_len(n)
p1 <- 30
xy <- sin(t*2*pi/p1 + 50)
int <- c(rep(1, 99 + 100), seq(1,3,2/100), seq(3,1,-2/100), rep(1,100 + 99))
dt <- cumsum(int)
cond <- dt < 75
xy <- xy[!cond]</pre>
```

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```
dt <- dt[!cond]/1.2 - 62.5

res <- dq.algorithm(xy, dt)

opar <- par("mfrow")

par(mfrow = c(3,1))

plot(dt, xy, type = "o", pch = 19, main = "Frequency carrier")

plot(dt, 1/res$f, pch = 19, type = "1", log = "y", lwd = 2, ylim = c(25,80), main = "Period (Direct Quadrature method)", ylab = "Period")

plot(res$idt[,1], xy, type = "o", pch = 19, main = "Identity tuning", axes = FALSE, ylab = "xy", xlab = "dt")

ap <- approx(x = dt, y = res$idt[,1], xout = seq(0,600, by = 20))

axis(1, at = ap$y, labels = ap$x)
axis(2)
box()

par(mfrow = opar)</pre>
```

extremist

Gives local extrema and zero crossings intervals

Description

Gives local minimas, maximas and zero crossings. Optimised for large data sets; the sky is the limit (and by the sky I mean the ability of R and your computer to memorise large data sets; but within this limit the algorithm can handle millions of points quickly).

Usage

```
extremist(xy, bound = FALSE, local = TRUE, zc = TRUE)
```

Arguments

| ху | the values where to find the local extremas |
|-------|---|
| bound | whether to consider the first and last points as both minima and maxima, for special purposes. Default is F, has it should be. |
| local | whether to consider the first and last points as local minima and maxima, if TRUE by default, otherwise these first and last points will be ignored |
| zc | whether to return the zero crossings |

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Value

a list of the indexes of the left (l) and right (r) boundaries for the minima (minindex), maxima (maxindex) and zero crossing (cross), along with the number of extrema and zero crossings

```
# Function script ----
xy \leftarrow c(1,0,0,0,4,5,5,0.5,-0.5,0.5,0,2,2,1,-1,-1,1,1,0,0,-4,-2,2,1,0,0.5,0,
        NA, 0.5,0,-0.5,3,2,3,0,0.5,4,4,0)
impressme <- 0 # Increase up to 5 or 6 to be impressed (bugs if your system
                # can't handle the size of the data).
               # If you increase it, do not run the plot script.
xy <- rep(xy, round(10^impressme))</pre>
print(paste("You are running ", length(xy), " points", sep = ""))
res <- extremist(xy)</pre>
# Plot script: do not run if you increase the impressme parameter ----
mini <- unique(c(res$minindex[[1]], res$minindex[[2]]))</pre>
maxi <- unique(c(res$maxindex[[1]], res$maxindex[[2]]))</pre>
zeri <- unique(c(res$cross[[1]], res$cross[[2]]))</pre>
1 <- length(xy)</pre>
opar <- par("mfrow")</pre>
par(mfrow = c(3,1))
plot(1:1, xy, type = "o", pch = 19)
points(mini, xy[mini], pch = 19, col = "blue")
plot(1:1, xy, type = "o", pch = 19)
points(maxi, xy[maxi], pch = 19, col = "red")
plot(1:1, xy, type = "o", pch = 19)
points(zeri, xy[zeri], pch = 19, col = "green")
abline(h = 0, col = "grey")
par(mfrow = opar)
```

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Description

Performes EEMD

Usage

```
extricate(
  хy,
 dt,
 nimf,
  ini = NULL,
  repl = 1,
 comb = 100,
 mirror_noise = TRUE,
 factor_noise = 3,
  unit_noise = "1stdiff",
  sifting = 1,
  output_sifting = FALSE,
  remove = "lin.trend",
  bind = FALSE,
  speak = FALSE,
  plot_process = FALSE,
 pdf = TRUE,
  name = "extricate",
  ext = ".pdf",
  dir = tempdir(),
 width = 10,
 height = 20,
  track = TRUE,
 openfile = TRUE
)
```

Arguments

| xy | signal, maybe linearly interpolated to have regular sampling interval |
|--------------|---|
| dt | depth/time |
| nimf | number of modes/components/intrinsic mode functions to decompose the signal into |
| ini | an optional vector of length n of the eventual initial Intrinsic Mode Function xy would be a demodulation of, if it is a demodulation. In that case the mode indexes will start at 2. |
| repl | the amount of decompositions to output |
| comb | the amount of decompositions each output decomposition will be a combination of. Has to be a multiple of 2 (even and odd extension stacks have to be combined in any case) |
| mirror_noise | whether to generate a mirrored noise signal (for even and odd extension) that will cancel perfectly when combining the decompositions |

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a factor for the amplitude of white noise (finite amplitude obtained via runif).

bind whether to bind the removed linear trend or mean to the last component (T), or

to add it as another component (F)

speak whether to print a sentence at each sifting: it gives the stack (even or odd), the

mode number and sifting number

plot_process whether to have a plot of the entire sifting process. This slows down the algo-

rithm, use with low 'repl' and 'comb' values for visualisation purposes

pdf whether the plot be directly set as a pdf file

name, ext, dir, width, height, track, openfile

arguments to provide to pdfDisplay if plot_process and pdf are TRUE

Value

factor_noise

a list made of \$xy (original signal), \$dt (depth/time), \$m (a matrix of the decomposition), \$repl (the replicate id of each point) and \$mode (the mode id of each point). If output_sifting is TRUE, additional \$even_sifting and \$odd_sifting data.tables are provided, giving the condensed siftings for the even and odd extensions.

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```
integrity(xy, dec)
parsimony(dec)

plot_emd(dec, select = c(4, 6), pdf = FALSE)
## Not run:
plot_emd(dec, li = list(v = 0), dir = tempdir())
## End(Not run)
```

gzc

Calculates instantaneous frequency using the GZC method

Description

Calculates instantaneous frequency using the Generalised Zero-Crossing method from Huang et al., 2009. General wrapper for the gzc.algorithm function that does all the actual work.

Usage

```
gzc(
  emd = NULL,
  ini = NULL,
  m = NULL,
  dt = NULL,
  repl = 1,
  mode = NULL,
  dtout = NULL,
  output = 1,
  warn = TRUE
)
```

Arguments

| emd | emd-type object |
|-------|---|
| ini | an optional vector of length n of the eventual initial Intrinsic Mode Function xy would be a demodulation of, if it is a demodulation. It will be integrated to the results as mode 1. |
| m | a matrix of the amplitude values (xy) of the components, each column being a component. Each column should have the same number of non NA values. Vectors, for 1 component, are accepted. Is overridden by emd. |
| dt | the depth or time value. Is overridden by emd. |
| repl | the amount of replicates in m. Is overridden by emd. |
| mode | the mode sequence index to give to each replicated IMFs |
| dtout | the dt values to sample the frequency and amplitude from if output = 2. |

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output the style of the output, whether 0, 1 or 2. 0 provides the raw output of gzc.algorithm,

1 and 2 provides a matrix with \$dt (depth/time), \$f (frequency) and \$a ()amplitude, but with output = 1 the matrix provides the dt only at the extremas and zero-crossings, whereas with output = 2 the dt values are the ones provided with the dtout parameter. 1 is better for plots, 2 allows easier calculations to be

performed downstream.

warn whether to warn if the sampling interval defined by the dtout parameter is to

small (redirected from StratigrapheR::tie.lim)

Value

depending on the output parameter:

output = 0 provides the raw output of gzc.algorithm, with \$ldt and \$rdt (the left and right boundaries of the depth/time intervals), \$f (frequency) and \$a (amplitude). To that are added \$repl (the replicate id) and \$mode (the mode id)

output = 1 or 2 provides a matrix with \$dt, \$f and \$a, but with output = 1 the matrix provides the dt only at the extremas and zero-crossings, whereas with output = 2 the dt values are the ones provided with the out parameter. 1 is better for plots, 2 allows easier calculations to be performed downstream.

References

Huang, Norden E., Zhaohua Wu, Steven R. Long, Kenneth C. Arnold, Xianyao Chen, and Karin Blank. 2009. "On Instantaneous Frequency". Advances in Adaptive Data Analysis 01 (02): 177–229. https://doi.org/10.1142/S1793536909000096.

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gzc.algorithm

Calculates instantaneous frequency of simplified IMF using the GZC method

Description

Calculates instantaneous frequency of simplified IMF using the Generalised Zero-Crossing method from Huang et al., 2009.

Usage

```
gzc.algorithm(xy, dt)
```

Arguments

xy a matrix of amplitude

dt a vector of depth or time values

Details

the GZC method is precise to 1/4th of a period, so the results are provided between left and right points, i.e. either an extrema or a zero-crossing.

Value

```
a list of $ldt (left position), $rdt (right position), $f (frequency) and $a (amplitude)
```

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References

Huang, Norden E., Zhaohua Wu, Steven R. Long, Kenneth C. Arnold, Xianyao Chen, and Karin Blank. 2009. 'On Instantaneous Frequency'. Advances in Adaptive Data Analysis 01 (02): 177–229. https://doi.org/10.1142/S1793536909000096.

Examples

```
1,1,0,-1,-1,0,1,1,0,-1,-1,0,1,1,0,-1,-1
dti <- 1:length(xyi)</pre>
d \leftarrow simp.emd(m = xyi, dt = dti)
xy < - d$xy
dt <- d$dt
res <- gzc.algorithm(xy, dt)</pre>
opar <- par('mfrow')</pre>
par(mfrow = c(2,1))
plot(dti, xyi, pch = 19, type = "o", ylab = "xy", xlab = "dt")
points(dt, xy, pch = 19, col = "green")
points(res$ldt, res$a, pch = 19, col = "red")
points(res$rdt, res$a, pch = 19, col = "red")
plot(dt, rep(max(res$f, na.rm = TRUE), length(dt)), type = "n",
    ylab = "Frequency", xlab = "dt",
    ylim = c(0, 2 * max(res$f, na.rm = TRUE)))
points(res$ldt, res$f, pch = 19)
points(res$rdt, res$f, pch = 19)
par(mfrow = opar)
```

gzc.departure

departure of instantaneous frequency to generalized zero-crossing

Description

departure of instantaneous frequency to generalized zero-crossing of instantaneous frequency. The departure is calculated as the exponential of the absolute difference of logarithms of frequencies obtained using a robust generalized zero-crossing method through the gzc function (where the components are simplified into extrema separated by zero-crossings) and instantaneous frequency computed from another method

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Usage

```
gzc.departure(
  pulse = NULL,
  dt = NULL,
  m = NULL,
  f = NULL,
  repl = 1,
  mode = NULL,
  simplify = TRUE
)
```

Arguments

| pulse | a pulse object object |
|----------|--|
| dt | the depth or time. Is overridden by pulse. |
| m | a matrix of the modes to calculate the gzc frequency from. Is overridden by pulse. |
| f | a matrix of the frequencies to compare to gzc. |
| repl | the amount of replicates in m. Is overridden by emd. |
| mode | the mode sequence index to give to each replicated IMFs. Is overridden by emd. |
| simplify | whether to average the value for each component of each replicate |

Value

If simplify is TRUE, the function returns the average gzc departure as a data frame where the columns stand for the modes and the rows for the replicates. If simplify if FALSE, the function returns the functions returns local gzc departure.

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HilbertEnvelope

Instantaneous amplitude

Description

Generates the instantaneous amplitude of an analytic signal given by HilbertTransform

Usage

```
HilbertEnvelope(asig)
```

Arguments

asig

The analytic signal returned by HilbertTransform

Value

envelope Instantaneous amplitude

Author(s)

Daniel C. Bowman (in the hht package)

See Also

HilbertTransform, InstantaneousFrequency

22 HilbertTransform

Examples

```
tt <- seq(1000) * 0.01
sig <- sin(4 * pi * tt) + sin(3.4 * pi * tt)
asig <- HilbertTransform(sig)
env <- HilbertEnvelope(asig)
plot(tt, sig, type = "l")
lines(tt, env, col = "red")
lines(tt, -env, col = "red")</pre>
```

HilbertTransform

The Hilbert transform

Description

Creates the analytic signal using the Hilbert transform.

Usage

```
HilbertTransform(sig)
```

Arguments

sig

Signal to transform.

Details

Creates the real and imaginary parts of a signal.

Value

```
asig Analytic signal
```

Author(s)

Daniel C. Bowman (in the hht package)

See Also

HilbertEnvelope, InstantaneousFrequency

inst.pulse 23

Examples

inst.pulse

Computes instantaneous frequency using the Hilbert transform

Description

Calculates instantaneous frequency using the Hilbert transform (HT), normalised Hilbert transform (NHT) or the direct quadrature (DQ) methods. Normalisation is done for NHT and DQ using Huang et al., 2009 algorithm, but the empirical normalisation scheme can fail due to overshoot or undershoot of the spline. Additional research is necessary for that last feature.

Usage

```
inst.pulse(
  emd = NULL,
  imf = NULL,
 m = NULL,
 dt = NULL,
  ini = NULL,
  repl = 1,
 mode = NULL,
  last = FALSE,
  plot = TRUE,
 method = "HT",
  delta = NULL,
  tolerance = 8,
  relative = TRUE,
  breaks = 500,
  bins = 100,
  cut = 18,
  lines = NULL
)
```

24 inst.pulse

Arguments

| emd | an emd object |
|-------------------|--|
| imf | a matrix of same frequency modes to calculate the frequency from. Is overridden by emd. This allows to calculate and visualise the results for single IMFs more clearly than in a population plot. |
| m | a matrix of the modes to calculate the frequency from. Is overridden by emd and imf. |
| dt | the depth or time. Is overridden by emd. |
| ini | an optional vector of length n of the eventual initial Intrinsic Mode Function xy would be a demodulation of, if it is a demodulation. It will be integrated to the results as mode 1. |
| repl | the amount of replicates in m. Is overridden by emd. |
| mode | the mode sequence index to give to each replicated IMFs. Is overridden by emd. |
| last | whether to use the last mode (trend/residue). |
| plot | whether to have a plot summary of the output. |
| method | the IF calculation method: "HT" for Hilbert transform (default), "NHT" for normalised Hilbert transform, and "DQ" for direct quadrature. The two last require normalisation, which can sometimes fail. |
| delta, tolerance | e, relative |
| | parameters to feed to respace for interpolation |
| breaks, bins, cut | |
| | parameter for the plots: breaks is fed to plot_hist, bins is fed to plot_hex, and cut defines the number of color cuts for plot_hex. For better control use plot_hist and plot_hex directly. |
| lines | the period of lines to be added to the plots for better visualisation |

Value

a list made of \$dt (depth/time), \$f (instantaneous frequency), \$a (instantaneous amplitude),\$repl (the replicate id of each point) and \$mode (the mode id of each point)

References

Huang, Norden E., Zhaohua Wu, Steven R. Long, Kenneth C. Arnold, Xianyao Chen, and Karin Blank. 2009. "On Instantaneous Frequency". Advances in Adaptive Data Analysis 01 (02): 177–229. https://doi.org/10.1142/S1793536909000096.

```
set.seed(42)

n <- 600
t <- seq_len(n)

p1 <- 30
p2 <- 240</pre>
```

inst.ratio 25

inst.ratio

Computes instantaneous ratio of frequency

Description

Computes instantaneous ratio of frequency

Usage

```
inst.ratio(
 pulse = NULL,
 dt = NULL,
 f = NULL,
 a = NULL,
  repl = 1,
  plot = TRUE,
  sqrt.rpwr = TRUE,
  style = "b",
  select = NA,
 bins = 100,
  cut = 18,
  lines = NULL,
 width = 10,
 height = 10,
 name = "Ratio",
```

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```
ext = ".pdf",
dir = tempdir(),
track = TRUE,
openfile = TRUE
```

Arguments

```
pulse
                   a pulse object (created by inst.pulse for instance)
dt
                   depth/time. Is overridden by pulse.
f
                  instantaneous frequency. Is overridden by pulse.
                   instantaneous amplitude. Is overridden by pulse.
а
repl
                   number of replicates in f
                   whether to plot an output
plot
sqrt.rpwr, style, select, bins, cut, lines, width, height
                   parameters to feed to plot_ratio for the plots
name, ext, dir, track, openfile
                   parameters to feed to pdfDisplay in plot_ratio for pdf plot.
```

Value

a list of depth/time (\$dt), frequency (\$f), ratio of frequency (\$ratio), if a is provided; the ratio power (\$rpwr) i.e. the multiplication of the instantaneous amplitudes of the modes two by two, the replicates id (\$repl)and id for the first and second frequency modes used for the ratio (\$l for the first, \$r for the second, or \$lr for the two combined)

```
## End(Not run)
integrity(xy, dec)
parsimony(dec)
ht <- inst.pulse(dec, lines = c(30, 240))
ratio <- inst.ratio(ht, style = "s", lines = 8)</pre>
```

InstantaneousFrequency

Derive instantaneous frequency

Description

Calculates instantaneous frequency from an analytic signal.

Usage

```
InstantaneousFrequency(asig, tt, method = "arctan", lag = 1)
```

Arguments

asig Analytic signal produced by HilbertTransform

tt Sample times

method How the instantaneous frequency is calculated. "arctan" uses the arctangent

of the real and imaginary parts of the Hilbert transform, taking the numerical derivative of phase for frequency. "chain" uses the analytical derivative of the

arctangent function prior to performing the numerical calculation.

lag Differentiation lag, see the diff function in the base package.

Value

instfreq Instantaneous frequency in 1/time

Note

The "arctan" method was adapted from the hilbertspec function in the EMD package.

!!IMPORTANT!! The numeric differentiation may be unstable for certain signals. For example, high frequency sinusoids near the Nyquist frequency can give inaccurate results when using the "chain" method. When in doubt, use the PrecisionTester function to check your results!

Author(s)

Daniel C. Bowman (in the hht package)

See Also

PrecisionTester

28 integrity

| integrity Integrity of a decomposition |
|--|
|--|

Description

The function additions each component of a decomposition by depth/time, subtract it with the original signal, and provides the absolute of this subtraction. This is allows to verify if the decomposition is computed correctly.

The bulk value is the cumulated value of this proxy. If the decomposition is done right the value should be very small, but non-zero due to the floating-point arithmetics used by computers that generate tiny errors. Its actually interesting: the first computations of the orbital solutions were strongly affected by this error, as the chaotic behaviour of the equations enhanced the effect of these tiny tiny errors.

Usage

```
integrity(xy, emd = NULL, m = NULL, repl = 1, bulk = TRUE)
```

Arguments

| xy | the signal |
|------|--|
| emd | an emd object to test. The emd\$xy original signal is not used, to avoid confusion: you always have to provide the xy signal yourself. |
| m | a matrix with columns of same length that xy, made of the decomposition of the signal. Is overridden by emd. |
| repl | the replication of decompositions in m. Is overridden by emd. |
| bulk | whether to have a bulk value each decomposition replication, or for each dt of each replication |

Value

a matrix with each column being a replication, or a list of bulk values for each replication

```
set.seed(42)

n <- 600
t <- seq_len(n)

p1 <- 30
p2 <- 240

xy <- (1 + 0.6 * sin(t*2*pi/p2)) * sin(t*2*pi/p1) + 2 * sin(t*2*pi/p2) +
    rnorm(n, sd = 0.5)

inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
```

is.ratio 29

is.ratio

Check ratio objects

Description

Check ratio objects

Usage

```
is.ratio(ratio)
```

Arguments

ratio

a ratio object to check

30 mode.in

is.simp.emd

Tests for simplified EMD

Description

Tests whether each column of a matrix is an alternation of -minima zero-crossing maxima zero-crossing-

Usage

```
is.simp.emd(xy)
```

Arguments

ху

a vector or matrix of values to test

Examples

mode.in

Add / Remove / Bind modes in emd objects

Description

Add / Remove / Bind modes in emd objects

mode.in 31

Usage

```
mode.in(emd, xy, mode = NA, adjust = TRUE, name = "Added")
mode.out(obj, keep = NULL, lose = NULL, adjust = F, reorder = F)
mode.bind(emd, mode = NA, xy = NULL, adjust = T, name = "bound")
```

Arguments

emd emd-type object

xy an Instrinsic Mode Function to add

mode, keep, lose [mode.in] the position where to add the mode / [mode.out] the modes to keep or

lose / [mode.bind] the modes to merge

adjust whether to adapt the initial signal of an emd object (\$xy in the emd object) when

adding or removing a mode

name the name of the new mode obj emd or pulse type object

reorder whether to reinitialise the index of modes when suppressing one

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy < (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5)
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,
                  repl = 10, comb = 10, factor_noise = 10,
                  speak = TRUE)
opar <- par('mfrow')</pre>
par(mfrow = c(2,1))
integrity(xy, dec)
ht <- inst.pulse(dec, plot = FALSE)</pre>
plot_hist(x = 1/ht$f, breaks = 500, id = ht$mode,
```

32 n.extrema

n.extrema

Number of extrema/zero-crossings

Description

Computes the number of extrema and zero-crossings for different groups of data, by their id or separated by NA values

Usage

```
n.extrema(
   xy,
   id = NULL,
   use.names = TRUE,
   bound = FALSE,
   local = FALSE,
   zc = TRUE
)
```

Arguments

xy signal or decomposed signal

id the id for different groups. If any NA value is in xy, it will also separate two

groups of data

use.names whether to use the names in id bound, local, zc parameters to feed to extremist

Value

a list of the number of minima (\$n.min), maxima (\$n.max), and, if zc = TRUE, zero-crossings (\$n.cross)

normalise 33

Examples

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy < (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5)
xy <- xy - mean(xy)
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,</pre>
                repl = 1, comb = 40, factor_noise = 10,
                 speak = TRUE)
integrity(xy, dec)
parsimony(dec)
n.extrema(dec$m, dec$mode)
plot_emd(dec, select = c(6,8,9), pdf = FALSE, adapt.axis = TRUE)
## Not run:
plot_emd(dec, li = list(v = 0), adapt.axis = TRUE, dir = tempdir())
## End(Not run)
```

normalise

Empirical AM and FM decomposition

Description

Applies the normalisation scheme of Huang et al., 2009 to decompose any Intrinsic Mode Functions obtained (usually via Empirical Mode Decomposition) into an Frequency Modulated component of amplitude 1, also called carrier, and its Amplitude Modulated enveloppe. The carrier can then be used to compute the instantaneous frequency via the Normalised Hilbert Transform (NHT) or by calculating its Direct Quadrature (DQ) (Huang et al., 2009). HOWEVER THIS FUNCTION CAN FAIL due to overshoot or undershoot of the spline fitting. Additional research is necessary.

Usage

```
normalise(emd = NULL, m = NULL, dt = NULL, repl = 1, last = TRUE, speak = TRUE)
normalize(emd = NULL, m = NULL, dt = NULL, repl = 1, last = TRUE, speak = TRUE)
```

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Arguments

| emd | an emd object |
|-------|--|
| m | a matrix of the modes to calculate the amplitude and the frequency carrier from. Is overridden by emd. |
| dt | the depth or time. Is overridden by emd. |
| repl | the amount of replicates in m. Is overridden by emd. |
| last | whether to use the last mode (trend/residue). |
| speak | whether to print a sentence at each iteration |

Value

a list of two matrices: \$fc (frequency carrier) and \$a (instantaneous amplitude)

References

Huang, Norden E., Zhaohua Wu, Steven R. Long, Kenneth C. Arnold, Xianyao Chen, and Karin Blank. 2009. 'On Instantaneous Frequency'. Advances in Adaptive Data Analysis 01 (02): 177–229. https://doi.org/10.1142/S1793536909000096.

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy < (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
        rnorm(n, sd = 0.5)
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,</pre>
                repl = 1, comb = 100, factor_noise = 10,
                speak = TRUE)
plot_emd(dec, pdf = FALSE, select = 4)
integrity(xy, dec)
parsimony(dec)
m <- dec$m
res <- normalise(dt = dt, m = m, last = FALSE)</pre>
```

oscillate 35

oscillate

Modify a signal using a Van der Pol oscillator

Description

Modify a signal using a Van der Pol oscillator

Usage

```
oscillate(
    xy,
    dt,
    period,
    delta = 0.05,
    damp = 5e-05,
    f.noise = 5,
    f.signal = 0.95,
    dx = function(x, y, beta, damp) beta * y - x * (x^2 + y^2 - 1) * damp,
    dy = function(x, y, beta, damp) -beta * x - y * (x^2 + y^2 - 1) * damp,
    xi = if (length(xy) != 0) xy[1] else 0.5,
    yi = if (length(xy) != 0) xy[1] else 0.5,
    normalise = TRUE,
    limit = TRUE
)
```

Arguments

```
xy initial signal (vector or matrix)

dt depth/time (same length than length/rows of xy)

period the period of the oscillator (length 1 or n)

delta the sampling interval for iteration (length 1 or n)
```

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| damp | damping parameter |
|-----------|---|
| f.noise | a factor of the amount of noise (length 1 or n) |
| f.signal | a factor of the amount of signal (length 1 or n) |
| dx, dy | the differentials used in the oscillator. They should be provided as functions needing x , y , beta (2*pi/period) and damp (damping) parameters |
| xi | the initial x value |
| yi | the initial y value |
| normalise | whether to recenter the output signal on the initial signal |
| limit | whether to warn when parameters are irrealistic (subjective) |

Examples

parsimony

Parsimony of a decomposition

Description

The function additions the absolute values of each component of a decomposition by depth/time, and computes the ratio of that with the absolute values of the signal. This is done either by depth/time or on the time/depth-cumulated signal (i.e. the bulk signal).

This is a proxy for parsimony: it is the factor of amplitude added by the decomposition. A perfect decomposition, that does not 'invent' wiggles, should approach 1, but will logically always be

parsimony 37

higher. However it is influenced by the absolute value of the initial signal: if the original signal is not centered around 0, the parsimony is not significative (it will artificially be closer to 1). To correct for that, the residue (part of the decomposition that is not centered around zero) has to be removed from the original signal.

Usage

```
parsimony(
  emd = NULL,
  xy = NULL,
  m = NULL,
  mode = NULL,
  repl = 1,
  bulk = TRUE,
  correct = NA
)
```

Arguments

| emd | an emd object |
|---------|--|
| xy | the signal |
| m | a matrix with columns of same length that xy, made of the decomposition of the signal |
| mode | the mode sequence index to give to each replicated IMFs |
| repl | the replication of decompositions in m |
| bulk | whether to have a bulk value each decomposition replication, or for each dt of each replication |
| correct | the modes to remove from the original signal and decomposition for a significative parsimony calculation. If NA, it removes the last mode, considered as the residue. Can be a vector of several integers, standing for the columns of m. If |

Value

a matrix with each column being a replication, or a list of bulk values for each replication

Examples

NULL, no mode is removed

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```
plot_emd(dec, dir = tempdir())
## End(Not run)

parsimony(dec, correct = NULL)

parsimony(dec)
```

pile.down

Destacks a pile.up() signal

Description

Destacks a signal stacked by pile.up by averaging each repetition back to n multiples.

Usage

```
pile.down(x, stack, even, n = length(unique(stack$id)) - 2)
```

Arguments

| X | Treated signal |
|-------|---|
| stack | Initial stack from which the x signal is from |
| even | Whether the x signal comes from even extension part of the initial stack (if FALSE, it would come from the odd extension part) |
| n | The multiple of destacking (has to be a multiple of n/2 (n being the parameter used in pile.up), in other words a multiple of length(unique(stack\$id)) - 2 (minus 2 as the upper an lower extension are to be removed) |

Value

a matrix or a vector of the destacked signal

```
set.seed(42)
n <- 200
t <- seq_len(n)

p1 <- 25
p2 <- 75

xy <- (1 + 0.6 * sin(t*2*pi/p2)) * sin(t*2*pi/p1) + 2 * sin(t*2*pi/p2) +
    rnorm(n, sd = 0.5)

inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)
inter_dt[20] <- 20</pre>
```

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```
dt <- cumsum(inter_dt)</pre>
opar <- par()$mfrow
par(mfrow = c(1,1))
res <- pile.up(xy, dt, 4)
par(mfrow = c(2,1))
plot(res$ndt, res$even, type = "1", col = "blue")
plot(res$ndt, res$odd, type = "1", col = "red")
par(mfrow = c(opar))
# Small number of repetitions ----
opar <- par("mfrow")</pre>
par(mfrow = c(1,2))
stack <- pile.up(xy, dt, 10)</pre>
signal <- stack$even + runif(length(stack$even), -3, 3)</pre>
res <- pile.down(signal, stack, even = TRUE, n = 5)
plot(xy, dt, type = "1", lwd = 2, main = "Low number of repetitions")
lines(res, dt, type = "l", lty = 5, col = "red")
# High number of repetitions ----
stack <- pile.up(xy, dt, 1000)</pre>
signal <- stack$even + runif(length(stack$even), -3, 3)</pre>
res <- pile.down(signal, stack, even = TRUE, n = 500)
plot(xy, dt, type = "1", lwd = 2, main = "High number of repetitions")
lines(res, dt, type = "1", lty = 5, col = "red")
par(mfrow = c(opar))
```

pile.up

Repeat and stack a signal in central and line symmetry

Description

Repeats and stacks a signal duplicated in central (even) and line (odd) symmetry to apply Ensemble Empirical Mode Decomposition (EEMD) on one single vector following the simple boundary rule of Zeng and He (2004). This allows to avoid the iterations that are typical of EEMD. A complete

40 pile.up

set of signal is added by default at the upper and lower part of the stack, to be removed in the end process.

Usage

```
pile.up(xy, dt, n, warn = TRUE)
```

Arguments

the signal
 the depth/time positions of each xy
 the number of replicates you want. It has to be a multiple of two, as you will generate two stacks: the even and the odd one.
 warn whether you want to be annoyed

Value

a dataframe of the original dt (odt), the stack-modified dt (ndt), the inversion factor to change the even stack into the odd one and vice-versa (invert), the even xy stack (even) and the odd one (odd)

```
set.seed(42)
n <- 200
t <- seq_len(n)
p1 <- 25
p2 <- 75
xy \leftarrow (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5)
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
inter_dt[20] <- 20
dt <- cumsum(inter_dt)</pre>
opar <- par()$mfrow
par(mfrow = c(1,1))
res <- pile.up(xy, dt, 4)</pre>
par(mfrow = c(2,1))
plot(res$ndt, res$even, type = "1", col = "blue")
plot(res$ndt, res$odd, type = "1", col = "red")
par(mfrow = c(opar))
```

plot_emd 41

plot_emd

Plot a decomposition

Description

General plot for a complete decomposition (that can be summed back to the original signal)

Usage

```
plot_emd(
  emd = NULL,
  xy = NULL,
  ini = NULL,
  dt = NULL,
 m = NULL
 mode = NULL,
  repl = 1,
  size.xy = 5,
  size.dt = 25,
  style = 2,
  xylim = NULL,
  dtlim = NULL,
  inilim = NULL,
  vertical = TRUE,
  adapt.axis = FALSE,
  adapt.last = TRUE,
  select = NULL,
  over = NULL,
  s = list(type = "o", pch = 19, cex = 0.5),
 o = list(type = "1", col = "blue", lwd = 2),
i = list(type = "o", pch = 19, cex = 0.5),
  e = list(type = "1", col = "red", lwd = 2),
  la = list(h = c(), v = c(), col = "red", xpd = FALSE),
  ls = list(),
  li = list(col = "grey", lty = 5),
  box = TRUE,
  ax = list(),
  ay = list(),
  parg = list(),
  title = TRUE,
  t1 = "Signal",
  t2 = "Mode",
  pdf = TRUE,
  name = "EMD",
  ext = ".pdf",
  dir = tempdir(),
  track = TRUE,
```

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```
openfile = TRUE
)
```

Arguments

| emd | an emd object |
|------------------|--|
| xy | the original signal. Is overridden by emd. |
| ini | an optional vector of length n of the eventual initial Intrinsic Mode Function xy would be a demodulation of, if it is a demodulation. |
| dt | the depth/time. Is overridden by emd. |
| m | a matrix with columns of same length that xy, made of the decomposition of the signal. Is overridden by emd. |
| mode | which modes/decompositions to plot |
| repl | the replication of decompositions in m. Is overridden by emd. |
| size.xy, size.d | t |
| | the size i inches of each individual plot in pdf |
| style | whether to not plot the original signal (style = 0), to plot it as the first signal (style = 1), or to plot it before each individual mode (style = 2 , is the default) |
| xylim, dtlim, in | |
| | the boundaries for the plots (inilim stands for the xy boundaries of the plot of the initial IMF xy is a demodulation of, if applicable) |
| vertical | whether to have the depth/time [dt] axis vertically (geologist convention) or horizontaly (climatologist convention) |
| adapt.axis | whether to let the plot adapt the axis to see the variability of the decompositions. The default os to have a comparable x axis for each plots |
| adapt.last | whether to adapt the last plot as a residue (if TRUE the x axis will be identical to the one of the signal, not centered on 0) |
| select | the components to plot |
| over | which modes/decompositions will be cumulated and added to the signal plotted at their left or above them (if style = 2) |
| s, o, i, e | lists of parameters to feed lines, for the original signal, the cumulated modes/decompositions overlapping it, the modes/decompositions themselves, and the enveloppe of the initial signal used for demodulation if it applies, respectively. |
| la, ls, li | lists of parameters to provide the abline function (makes personalised lines for you to have a better grasp of the data). la will plot on all panels, ls on the signal ones, and li on the modes ones. |
| box | whether to draw boxes around the plots |
| ax, ay | lists of parameters to feed minorAxis, the function making the axes, for the x and y axes |
| parg | list of parameters to feed par |
| title | whether to write titles |
| t1 | the title for the signal |

```
t2 the title for the modes
pdf whether to plot as a pdf
name, ext, dir, track, openfile
```

parameters for the pdfDisplay function, namely the name of the pdf file, its extension (if you want to make a .svg file you can), the directory of the file, whether to track the changes (if you use sumatrapdf as a default pdf reader you can set it to F and it will avoid creating too many pdf files), and whether to directly open the file

Examples

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy \leftarrow (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5) + 0.01 * t
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7,</pre>
                  repl = 10, comb = 10, factor_noise = 10,
                  speak = TRUE)
plot_{emd}(dec, select = c(4,6), pdf = FALSE)
## Not run:
plot_emd(dec, dir = tempdir())
## End(Not run)
```

plot_hex

Group and/or log-scale hexagonal binning

Description

Group and/or log-scale hexagonal binning. Provides a legend indicating the count representations. USES THE GRID GRAPHICAL SYSTEM, BASE GRAPHICS NOT SUPPORTED. To add lines, polygons or text, use the l, g and t arguments.

Usage

```
plot_hex(
 х,
 у,
  id = NA,
  select = NA,
  uniform = TRUE,
 bins = 60,
  xbnds = range(x, na.rm = TRUE),
  ybnds = range(y, na.rm = TRUE),
  xlim = xbnds,
  ylim = ybnds,
  log = "",
  shape = 1,
 mincnt = 1,
 maxcnt = NA,
  colorcut = seq(0, 1, length = 17),
  colramp = function(n) matlab.like(length(colorcut) - 1),
  trans = NULL,
  inv = NULL,
  border = NULL,
  1wd = 0.1,
  cex = 1,
 main = ""
  xlab = "x",
  ylab = "y",
  xaxis = TRUE,
 yaxis = TRUE,
  xaxs = "r",
  yaxs = "r",
  box = TRUE,
  mar = c(0.15, 0.125, 0.15, 0.2),
  legend = TRUE,
  leg_sep = 0.1,
  xpd_hex = 0.75,
  xpd_leg = 1.5,
  1 = list(x = NULL, y = NULL, default.units = "native"),
  g = list(x = NULL, y = NULL, default.units = "native"),
  t = list(label = NULL, default.units = "native"),
 plot = TRUE
)
```

Arguments

x, y vectors giving the coordinates of the bivariate data points to be binned.

id a vector of ids for each x value, to separate different groups of data

select the groups of ids to plot

uniform whether to keep the creaks defined by the entire matrixes when selecting only a

part of it

bins the number of bins partitioning the range of xbnds.

xbnds, ybnds horizontal and vertical limits of the binning region in x or y units respectively;

must be numeric vector of length 2.

xlim, ylim the limits of the plot

log a character string which contains "x" if the x axis is to be logarithmic, "y" if the

y axis is to be logarithmic and "xy" or "yx" if both axes are to be logarithmic.

shape the theoretical shape = yheight/xwidth of the plotting. This adapts the form of

the hexagons accordingly.

mincnt, maxcnt fraction of cell area for the lowest and largest count, respectively

colorcut vector of values covering [0, 1] that determine hexagon color class boundaries

and hexagon legend size boundaries. Alternatively, an integer (<= maxcnt) spec-

ifying the number of equispaced colorcut values in [0,1].

colramp function accepting an integer n as an argument and returning n colors.

trans a transformation function for the counts such as log10

inv the inverse transformation function (if trans = log10, inv should for instance

be function(x) 10^x .

border the color of the border of the hexagons. By default it will be the color of the

filling

1wd the width of the border of the hexagons.

cex the magnification of text.

main main title.

box

xlab, ylab x and y axis labels respectively.

xaxis, yaxis whether to plot the x and y axes respectively.

xaxs, yaxs The style of axis interval calculation to be used for the axes. By default the style

"r" (regular) first extends the data range by 4 percent at each end and then finds an axis with pretty labels that fits within the extended range. Style "i" (internal) just finds an axis with pretty labels that fits within the original data range.

whether to plot a box.

mar a numerical vector of the form c(bottom, left, top, right) which gives the room

the give to the margins in Normalised Parent Coordinates (see grid package for

more information)

legend whether to plot the legend.

leg_sep the distance between hexagons and text f the legend in Normalised Parent Coor-

dinates left on the right margin

xpd_hex factor to expand the legend hexagons xpd_leg factor to expand the height of the legend

a list of arguments to feed to grid::grid.polyline ATTENTION the grid

package has to be loaded

```
g a list of arguments to feed to grid::grid.polygon ATTENTION the grid package has to be loaded
t a list of arguments to feed to grid::grid.text ATTENTION the grid package has to be loaded
plot whether to plot. If FALSE, returns a grob.
```

```
library(grid) # To use the gpar function
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy < (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
        rnorm(n, sd = 0.5)
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,</pre>
                repl = 10, comb = 10, factor_noise = 10,
                 speak = FALSE)
## Not run:
plot_emd(dec, dir = tempdir())
## End(Not run)
integrity(xy, dec)
parsimony(dec)
ht <- inst.pulse(dec, plot = FALSE)</pre>
plot_hex(x = 1/htf, y = htfa, bins = 100, ybnds = c(0,2),
         \log = x^n, trans = \log 10, inv = function(x) 10^x,
         main = "Spectral Population", xlab = "Period", ylab = "Amplitude")
plot_hex(x = 1/htf, y = htfa, bins = 100, ybnds = c(0,2),
         log = "x", trans = log10, inv = function(x) 10^x,
         main = "Spectral Population", xlab = "Period", ylab = "Amplitude",
         id = ht \mod e, select = c(4,6,7),
         1 = list(x = c(30, 30, 240, 240), y = unit(c(0,1,0,1), "npc"),
                 id = c(1,1,2,2), gp = gpar(col = c("red", "blue"), lwd = 2)),
        g = list(x = c(18, 50, 50, 18, 18, 50, 50, 18),
                 y = c(0, 0, 1.9, 1.9, 2.05, 2.05, 1.95, 1.95),
                 id = c(1,1,1,1,2,2,2,2),
                 gp = gpar(col = c("red", NA), fill = c(NA, "white"), lwd = 2)),
```

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```
t = list(label = "Mode 4", x = 30, y = 2, gp = gpar(col = "red")))
```

plot_hist

Group and/or log-scale histogram

Description

Specialised histogram: allows to work in log-scale (for x) and to distinguish different groups of data

Usage

```
plot_hist(
 х,
 breaks = 100,
 id = NA,
  select = NA,
 pile = TRUE,
 line = FALSE,
 mids = FALSE,
 xlim = NA,
 ylim = NA,
  xlog = FALSE,
 axes = TRUE,
 xa = list(),
 ya = list(),
 main = "",
 xlab = "X",
 ylab = "Counts",
  col = NA,
 border = NA,
  text = FALSE,
 labels = NA,
  t = list(adj = c(0.5, -2), font = 2),
  add = FALSE
)
```

Arguments

x vector or matrix

breaks one of:

- a vector giving the breakpoints between histogram cells,
- a function to compute the vector of breakpoints,
- a single number giving the number of cells for the histogram,
- a character string naming an algorithm to compute the number of cells (see 'Details' in hist),

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• a function to compute the number of cells.

In the last three cases the number is a suggestion only; as the breakpoints will be set to pretty values, the number is limited to 1e6 (with a warning if it was larger). If breaks is a function, the x vector is supplied to it as the only argument (and the number of breaks is only limited by the amount of available memory).

id a vector of ids for each x value, to separate different groups of data

select a vector of id values idenifying the groups of data to plot and their order

pile whether to cumulate the different one on the other

line whether to plot as lines or rectangles

mids if lines is TRUE, whether the nodes of the lines are the middle positions or the

upper corner of the rectangles.

xlim, ylim the boundaries for the plots. If ylim = NA the upper ylim will be increased by

10% to allow for text (see 'text' parameter)

xlog whether to set the x axis in log scale

axes whether to plot the axes

xa, ya list of arguments to feed minorAxis for the x and y axes respectively

main, xlab, ylab the main title and the labels of the x and y axes

col a function or a character vector defining the colors of the different modes

border the colour of the borders, by default identical to col

text if there are different groups, whether to add a number above each of them to

distinguish them

labels the labels to put on top of each group t a list of parameters to feed text()

add whether to add the plot to a preexisting plot

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```
## Not run:
 plot_emd(dec, dir = tempdir())
 ## End(Not run)
 integrity(xy, dec)
parsimony(dec)
ht <- inst.pulse(dec, plot = FALSE)</pre>
opar <- par('mfrow')</pre>
 par(mfrow = c(2,1))
 plot_hist(x = 1/htf, breaks = 500,
                                                                                xlog = TRUE, xlab = "Period")
 plot_hist(x = 1/htf, breaks = 500, id = htf, breaks = 500, id = htf, breaks, id = 
                                                                                  xlog = TRUE, text = TRUE, add = TRUE, line = TRUE, pile = FALSE)
 abline(v = c(p1, p2), col = "red", lwd = 2, lty = 5)
 plot_hist(x = 1/htf, breaks = 500, id = htf, breaks = 500, id = htf, breaks, id = 
                                                                                xlog = TRUE, text = TRUE, xlab = "Period")
 abline(v = c(p1, p2), col = "red", lwd = 2, lty = 5)
 par(mfrow = opar)
```

plot_imf

Plot IMFs characteristics

Description

General plot for the envelope, instantaneous frequency (period) and identity tuning of an intrinsic mode function (IMF)

Usage

```
plot_imf(
  pulse,
  dtlim = NULL,
  xylim = NULL,
  flim = NULL,
  fclim = NULL,
  dtline = NULL,
  fline = NULL,
  rcline = NULL,
  vertical = FALSE,
```

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```
n = 10,
at.maj = NULL,
ls = list(type = "o", pch = 19),
le1 = list(lwd = 2),
le2 = list(lty = 2),
lid = list(type = "p", pch = 19),
lcos = list(),
ldt = list(lty = 5, lwd = 2),
lf = list(lty = 5),
lfc = list(lty = 5),
box = TRUE
)
```

Arguments

```
pulse
                  a pulse object
dtlim, xylim, flim, fclim
                  the boundaries for the plots, respectively for the depth/time, amplitude, fre-
                   quency and frequency carrier
dtline, fline, fcline
                  coordinates to add vertical/horizontal lines
vertical
                   whether to have the depth/time [dt] axis vertically
                   the the number of intervals defined by minor ticks (geologist convention) or
                  horizontaly (climatologist convention)
at.maj
                   the positions at which major tick-marks are to be drawn.
ls, le1, le2, lid, lcos
                  lists of parameters to feed lines, for the original signal, the upper and lower
                   envelope, the identity tuning, and the cosine line in the identity tuning
ldt, lf, lfc
                  lists of parameters to provide the abline function (makes personalised lines for
                   you to have a better grasp of the data).
box
                   whether to draw boxes around the plots
```

Details

the line in the identity tuning plot is a genuine cosine, independent from the signal. This is evident when riding waves generate dephasing.

```
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 40 * 21
am <- sin(t*2*pi/p2 + 50) + 0.03</pre>
```

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plot_pulse

Visualise the instantaneous frequencies and amplitudes of a decomposition

Description

Visualise the instantaneous frequencies and amplitudes of a decomposition

Usage

```
plot_pulse(
  pulse,
  style = "b",
  breaks = 500,
  bins = 100,
  cut = 18,
  lines = NULL,
  keep = NULL,
  lose = NULL)
```

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Arguments

pulse a pulse object (created by inst.pulse or as.pulse)

style whether to plot the distribution of frequency ('d'), the spectral population ('p') or both ('b', is the default)

breaks, bins, cut

parameter for the plots: breaks is fed to plot_hist, bins is fed to plot_hex, and cut defines the number of color cuts for plot_hex. For better control use plot_hist and plot_hex directly.

lines the period of lines to be added to the plots for better visualisation which modes to plot or to not (keep overrides lose)

Examples

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy < (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5) + t * 0.01
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10, repl = 10, comb = 10,</pre>
                  factor_noise = 10, speak = TRUE)
## Not run:
plot_emd(dec, dir = tempdir())
## End(Not run)
integrity(xy, dec)
parsimony(dec)
     <- inst.pulse(dec, plot = FALSE)
plot_pulse(ht, lines = c(30, 240))
```

plot_ratio

Visualise the instantaneous frequencies ratios of a decomposition

Description

Visualise the instantaneous frequencies ratios of a decomposition

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Usage

```
plot_ratio(
  ratio,
  sqrt.rpwr = TRUE,
  style = "b",
  select = NA,
 bins = 100,
  cut = 18,
  lines = NULL,
 plot = TRUE,
 width = 10,
 height = 10,
 name = "Ratio",
  ext = ".pdf",
  dir = tempdir(),
  track = TRUE,
 openfile = TRUE
)
```

Arguments

| | ratio | a ratio object (created by inst.ratio |
|---------------------------------|---------------|--|
| | sqrt.rpwr | whether to use the square root of ratio power (i.e. the square root of the multiplication of the instantaneous amplitudes of the modes two by two) rather than the ratio power itself. |
| | style | whether to plot a single plot in the graphics device ('s'), the to plot an ensemble of all the ratios combinations in a pdf ('e'), or both ('b', is the default) |
| | select | the groups of ratios combinations to plot in the single plot (in the "1/2" form) |
| | bins, cut | parameter for the plots: bins is fed to plot_hex, and cut defines the number of color cuts for plot_hex. For better control use plot_hex directly. |
| | lines | the ratio of lines to be added to the plots for better visualisation |
| | plot | whether to plot. Otherwise output a grob of the single plot. |
| | width, height | the width and height in inches of each separate plot in the ensemble of all the ratios combinations |
| name, ext, dir, track, openfile | | |
| | | parameters for the pdfDisplay function, namely the name of the pdf file, its |

parameters for the pdfDisplay function, namely the name of the pdf file, its extension (if you want to make a .svg file you can), the directory of the file, whether to track the changes (if you use sumatrapdf as a default pdf reader you can set it to F and it will avoid creating too many pdf files), and whether to directly open the file

```
set.seed(42)

n <- 600
t <- seq_len(n)</pre>
```

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```
p1 <- 30
p2 <- 240
xy \leftarrow (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5) + t * 0.01
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,</pre>
                  repl = 10, comb = 10,
                  factor_noise = 10, speak = TRUE)
## Not run:
plot_emd(dec, dir = tempdir())
## End(Not run)
integrity(xy, dec)
parsimony(dec)
      <- inst.pulse(dec, plot = FALSE)
ratio <- inst.ratio(ht, plot = FALSE)</pre>
plot_ratio(ratio, lines = c(8), style = "s")
plot_ratio(ratio, lines = c(8), style = "s", select = c("4/6"))
## Not run:
plot_ratio(ratio, lines = c(8), style = "e", dir = tempdir())
## End(Not run)
```

PrecisionTester

Test numerically determined instantaneous frequency against exact instantaneous frequency

Description

This function compares the performance of InstantaneousFrequency against signals of known instantaneous frequency. The known signal is of the form

$$x(t) = a\sin(\omega_1 + \varphi_1) + b\sin(\omega_2 + \varphi_2) + c$$

One can create quite complicated signals by choosing the various amplitude, frequency, and phase constants.

Usage

```
PrecisionTester(
  tt = seq(0, 10, by = 0.01),
  method = "arctan",
```

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```
lag = 1,
a = 1,
b = 1,
c = 1,
omega.1 = 2 * pi,
omega.2 = 4 * pi,
phi.1 = 0,
phi.2 = pi/6,
plot.signal = TRUE,
plot.instfreq = TRUE,
new.device = TRUE,
...
)
```

Arguments

| tt | Sample times. |
|---------------|--|
| method | How the numeric instantaneous frequency is calculated, see InstantaneousFrequency |
| lag | Differentiation lag, see the diff function in the base package |
| а | Amplitude coefficient for the first sinusoid. |
| b | Amplitude coefficient for the second sinusoid. |
| С | DC shift |
| omega.1 | Frequency of the first sinusoid. |
| omega.2 | Frequency of the second sinusoid. |
| phi.1 | Phase shift of the first sinusoid. |
| phi.2 | Phase shift of the second sinusoid. |
| plot.signal | Whether to show the time series. |
| plot.instfreq | Whether to show the instantaneous frequencies, comparing the numerical and analytical result. |
| plot.error | Whether to show the difference between the numerical and analytical result. |
| new.device | Whether to open each plot as a new plot window (defaults to TRUE). However, Sweave doesn't like dev.new(). If you want to use PrecisionTester in Sweave, be sure that new.device = FALSE |
| | Plotting parameters |

Value

The numerically-derived instantaneous frequency from InstantaneousFrequency

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Author(s)

Daniel C. Bowman (in the hht package)

See Also

InstantaneousFrequency

```
#Simple signal
tt <- seq(0, 10, by = 0.01)
a <- 1
b <- 0
c <- 0
omega.1 <- 30 * pi
omega.2 <- 0
phi.1 <- 0
phi.2 <- 0
PrecisionTester(tt, method = "arctan", lag = 1, a, b, c,
                omega.1, omega.2, phi.1, phi.2, new.device = FALSE)
#That was nice - what happens if we use the "chain" method...?
PrecisionTester(tt, method = "chain", lag = 1, a, b, c,
                omega.1, omega.2, phi.1, phi.2, new.device = FALSE)
#Big problems! Let's increase the sample rate
tt <- seq(0, 10, by = 0.0005)
PrecisionTester(tt, method = "chain", lag = 1, a, b, c,
                omega.1, omega.2, phi.1, phi.2, new.device = FALSE)
#That's better
#Frequency modulations caused by signal that is not symmetric about {\tt 0}
tt <- seq(0, 10, by = 0.01)
a <- 1
b <- 0
c <- 0.25
omega.1 <- 2 * pi
omega.2 <- 0
phi.1 <- 0
phi.2 <- 0
PrecisionTester(tt, method = "arctan", lag = 1, a, b, c,
                omega.1, omega.2, phi.1, phi.2, new.device = FALSE)
#Non-uniform sample rate
set.seed(628)
```

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ratios

Computes ratios of numerical values

Description

Computes ratios of numerical values

Usage

```
ratios(x)
```

Arguments

Х

values to compute the ratio from

Value

```
a dataframe of $ratio, $x1 and $x2
```

Examples

```
ratios(c(20,40,100,400))
```

repl.out

Remove / Bind replicates in emd objects

Description

Remove / Bind replicates in emd objects

Usage

```
repl.out(emd, keep = NULL, lose = NULL, reorder = FALSE)
repl.bind(emd, comb)
```

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Arguments

emd emd-type object
keep, lose the modes to keep or lose
reorder whether to reinitialise the index of replicates when suppressing one
comb the number of replicates that have to be bound together

Examples

```
set.seed(42)
n <- 600
t <- seq_len(n)
p1 <- 30
p2 <- 240
xy \leftarrow (1 + 0.6 * \sin(t*2*pi/p2)) * \sin(t*2*pi/p1) + 2 * \sin(t*2*pi/p2) +
  rnorm(n, sd = 0.5) + t * 0.01
inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)</pre>
dt <- cumsum(inter_dt)</pre>
dec <- extricate(xy, dt, nimf = 7, sifting = 10,</pre>
                  repl = 20, comb = 2, factor_noise = 10,
                  speak = TRUE, output_sifting = TRUE)
reduced <- repl.out(dec, keep = c(3,4))
parsimony(reduced)
plot_emd(reduced, pdf = FALSE, select = c(4,6))
combined <- repl.bind(dec, 10)</pre>
parsimony(combined)
plot_emd(combined, pdf = FALSE, select = c(4,6))
```

respace

Interpolate with even spacing

Description

Interpolate with even spacing. Can determine on its own the most conservative sampling interval (using the Greatest Common Rational Divisor)

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Usage

```
respace(
  dt,
  xy = NULL,
  delta = NULL,
  tolerance = 8,
  relative = TRUE,
  n.warn = 100
)
```

Arguments

dt depth/time (same length than length/rows of xy)

xy signal (vector or matrix)

delta the new sampling interval. If NULL, uses the Greatest Common Rational Divi-

sor

tolerance, relative

parameters for the divisor function (StratigrapheR package), to compute the

Greatest Common Rational Divisor

n.warn the amount of interpolated points in between the largest interval above which

a warning is provided. This warning can be useful to avoid needlessly long outputs, which might make any subsequent computation take too much time.

Value

a list of interpolated xy and dt values (\$xy and \$dt), plus a vector of logicals indicating whether each point was part of the initial input or was added by interpolation

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simp.emd

Simplifies the components of an EMD

Description

Simplifies the component of an EMD to only extremas and zero-crossings, and outputs problematic extrema: multiple extrema (extrema not separated by zero-crossings) and crossing extrema (extrema at zero).

Usage

```
simp.emd(emd = NULL, m = NULL, dt = NULL, repl = 1, use.names = FALSE)
```

Arguments

| emd | emd-type object |
|-----------|---|
| m | a matrix of the amplitude values (xy) of the components, each column being a component. Each column should have the same number of non NA values. Vectors, for 1 component, are accepted. Is overridden by emd. |
| dt | the depth or time value. Is overridden by emd. |
| repl | the amount of replicates in m. Is overridden by emd. |
| use.names | whether to use the column names to identify problematic extrema |

Value

a list of the depth or time values (\$dt) of the simplified IMF (Intrinsic Mode Function), of their amplitude (\$xy), and of the position and component of problematic multiple extrema (\$multiple_extrema) and crossing extrema (\$crossing_extrema)

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```
opar <- par("mfrow")

par(mfrow = c(1,1))

plot(dt, xytest, type = "o", pch = 19)
abline(h = 0, col = "grey")

me <- res$multiple_extrema$dt[res$multiple_extrema$repl == 1]
ce <- res$crossing_extrema$dt[res$multiple_extrema$repl == 1]
abline(v = me, col = "orange")
abline(v = ce, col = "darkred")

points(res$dt[,1], res$xy[,1], col = "red", pch = 19)

par(mfrow = opar)</pre>
```

simple.ssa

Simple SSA decomposition

Description

Simple wrapper for Singular Spectrum Analysis, using the functions of the Rssa package (which is not installed by default by the DecomposeR package, you should install it independently). This function allows unevenly sampled data.

Usage

```
simple.ssa(xy, dt, n = 10, remove = "trend", groups = list(), plot = T, ...)
```

Arguments

| xy | signal to be decomposed |
|--------|---|
| dt | depth/time |
| n | maximum amount of components |
| remove | whether to remove a linear trend ("trend", is the default), a mean value ("mean"), or to decompose as is (any other value) |
| groups | which components to regroup (list of the indices of elementary components to be regrouped, the entries of the list can be named, see the reconstruct() function in the Rssa package for more information) |
| plot | whether to show a visualisation of the importance of each component |
| ••• | any arguments to by given to the ssa() function (see Rssa package for more information) |

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Value

a list made of \$xy (original signal), \$dt (depth/time), \$m (a matrix of the decomposition), \$repl (the replicate id of each point) and \$mode (the mode id of each point).

Examples

```
set.seed(42)
n <- 600
t <- seq_len(n)

p1 <- 30
p2 <- 240

xy <- (1 + 0.6 * sin(t*2*pi/p2)) * sin(t*2*pi/p1) + 2 * sin(t*2*pi/p2) +
    rnorm(n, sd = 0.5) + 0.01 * t

inter_dt <- round(runif(length(xy), min = 0.5, max = 1.5),1)

dt <- cumsum(inter_dt)

res <- simple.ssa(xy, dt, groups = list(c(1,2), c= 3:10))

parsimony(res)

integrity(xy, res)

## Not run:
plot_emd(res, style = 1)
## End(Not run)</pre>
```

symmetry

Symmetry of components

Description

The function returns the highest factor of amplitude either in negative or positive values. This quantifies the symmetry of components.

Usage

```
symmetry(xy, names = "num")
```

Arguments

xy signal (vector or matrix)

names the names to use for the resulting vector. If NULL no names are provided, if NA

its the names of the columns of the xy matrix, if "num" it the column index of

the matrix xy

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