Package 'hhh4contacts'

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Description Meyer and Held (2017) <doi:10.1093 biostatistics="" kxw051=""> present an age-structured spatio-temporal model for infectious disease counts. The approach is illustrated in a case study on norovirus gastroenteritis in Berlin, 2011-2015, by age group, city district and week, using additional contact data from the POLYMOD survey. This package contains the data and code to reproduce the results from the paper, see 'demo(``hhh4contacts'')'.</doi:10.1093>				
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adaptP

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Adapt a Transition Matrix to a Specific Stationary Distribution

Description

Experimental Metropolis-Hastings algorithm, which tries to adjust a transition matrix such that its stationary distribution becomes approximately equal to a prespecified probability vector.

Usage

```
adaptP(P, target, niter = 1e+06)
```

Arguments

P a transition matrix, i.e., a square matrix where all rows sum to 1.

target the stationary probability vector to approximate.

niter the number of iterations of the MCMC algorithm

Value

the adjusted transition matrix.

Author(s)

Leonhard Held

See Also

C2pop for an alternative method.

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Examples

addGroups2WFUN

Group-Dependent Parametric Weights

Description

This function takes a specification of parametric weights and returns a modified version with group-dependent parameters. Only single-parameter functions are currently supported.

Usage

```
addGroups2WFUN(WFUN, groups, initial = rep.int(WFUN$initial, nlevels(groups)))
```

Arguments

WFUN	a list specification of parametric weights, e.g., as returned by the constructor functions $W_powerlaw$ and W_np .
groups	a vector of length nUnits determining to which group each unit belongs to. The supplied vector is converted to a factor using as.factor.

initial (named) vector of initial parameters.

Value

a list specifying group-dependent parametric weights for hhh4.

Author(s)

Sebastian Meyer

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Examples

```
data("measlesWeserEms")
WPLgroups <- addGroups2WFUN(
   W_powerlaw(maxlag = 5, normalize = FALSE, log = FALSE),
   groups = factor(sample(2, ncol(measlesWeserEms), replace = TRUE)))</pre>
```

 ${\tt aggregateC}$

Aggregate a Contact Matrix

Description

The (age) groups of a contact matrix can be joined together by the grouping argument, which first sums over contact groups (columns) and then averages over the corresponding participant groups (rows), optionally using weights such as the age distribution of the study participants.

Usage

```
aggregateC(C, grouping, ..., weights = NULL)
```

Arguments

С	a square numeric contact matrix such as contactmatrix_POLYMOD.
grouping	specification of how to aggregate groups (a named list or an integer vector of group sizes) using aggregateC with the "agedistri" attribute of the contact matrix as weights. If NULL, the original 5-year intervals are returned. The default setting produces the six age groups of Meyer and Held (2017).
• • •	specification of how to aggregate groups (alternative to using a named list as the grouping argument).
weights	a named numeric vector containing the weights for the rows of C, typically the age distribution of the participants. The names are matched against rownames (C). A value of NULL is interpreted as uniform weights.

Author(s)

Sebastian Meyer

aggregateCountsArray 5

Description

Aggregate an Array of Counts wrt One Dimension (Stratum)

Usage

```
aggregateCountsArray(counts, dim, grouping, ..., sort = TRUE)
```

Arguments

counts an (integer) array of counts with dimnames, e.g., counts or pop2011.

dim the dimension index of the stratum defining groups.

grouping, ... how the groups should be built.

sort logical indicating if the resulting array should be ordered by the grouping levels in the dim dimension.

Value

an array with similar dimensions as the input counts except for the dim dimension, which will be smaller due to the aggregation as specified by the grouping argument.

Author(s)

Sebastian Meyer

```
## works for matrices
aggregateCountsArray(pop2011, dim = 2, grouping = c(2,1,3,2,4))
aggregateCountsArray(pop2011, dim = 1, grouping = list(
    "a" = c("chwi","span","zehl"),
    "b" = c("neuk","scho")
))
## and of course for arrays
str(aggregateCountsArray(counts, dim = 3, grouping = c(1, 3, 4)))
```

C2pop

C2pop

Adapt a Contact Matrix to Population Fractions

Description

Experimental function, which tries to adjust a given contact matrix such that the stationary distribution of its row-normalized version (i.e., the transition matrix) becomes approximately equal to a prespecified probability vector.

Usage

```
C2pop(C, target, eps = 0.001, iter.max = 100)
```

Arguments

C a square numeric (contact) matrix.

target the stationary probability vector to approximate.

eps the tolerated mean absolute difference between the target probabilities and the

stationary distribution of the adapted, normalized contact matrix.

iter.max maximum number of iterations (guard against infinite loop).

Value

the adapted, normalized contact matrix.

Author(s)

Leonhard Held (original) and Sebastian Meyer (this implementation)

See Also

adaptP for an alternative method.

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contactmatrix

POLYMOD Contact Matrices for Germany

Description

The function contactmatrix retrieves various social contact matrices for Germany from the POLY-MOD survey (Mossong et al., 2008). Such a matrix contains the average numbers of reported contacts by participant age group. The original age groups (5-year intervals) can be joined together by the grouping argument, which first sums over contact groups (columns) and then averages over the corresponding participant groups (rows) using the corresponding age distribution as weights.

Usage

```
contactmatrix(
  which = c("corrected", "mossong", "reciprocal"),
  type = c("all", "physical"),
  grouping = c(1, 2, 2, 4, 4, 2),
  normalize = FALSE
)

contactmatrix_mossong

contactmatrix_mossong_physical

contactmatrix_POLYMOD

contactmatrix_POLYMOD_physical

contactmatrix_wallinga

contactmatrix_wallinga
```

Arguments

which

character string indicating which contact matrix to return. "mossong" uses the average numbers of reported contacts as published in Table S5 of Mossong et al. (2008), available as contactmatrix_mossong or contactmatrix_mossong_physical. "corrected" (from contactmatrix_POLYMOD or contactmatrix_POLYMOD_physical) fixes an error in these numbers related to the age group 70+ (see the Examples) and is the default. If which="reciprocal" (corresponding to contactmatrix_wallinga or contactmatrix_wallinga_physical as used by Meyer and Held, 2017), the returned social contact matrix fulfils reciprocity of contacts with respect to the age distribution of Berlin, pop2011, via the method of Wallinga et al. (2006).

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type a character string to select the type of contacts to use: either "all" contacts,

i.e., count both physical and pure conversational contacts, or only "physical"

contacts.

grouping specification of how to aggregate groups (a named list or an integer vector of

group sizes) using aggregateC with the "agedistri" attribute of the contact matrix as weights. If NULL, the original 5-year intervals are returned. The default

setting produces the six age groups of Meyer and Held (2017).

normalize a logical indicating whether to normalize the matrix such that each row sums to

1.

Format

The dataset contactmatrix_POLYMOD and its variants are all square numeric matrices with 15 rows (participants) and 15 columns (contacts), labelled with the corresponding age groups. There is an attribute "agedistri", a named numeric vector of length 15, which for the "_mossong_" and "_POLYMOD_" variants gives the age distribution of the German POLYMOD sample, and for the _wallinga_ variants gives the age distribution of Berlin, i.e., prop.table(colSums(pop2011)).

Value

a square numeric matrix containing the average numbers of contact persons recorded per day per survey participant in Germany, potentially averaged over multiple *row* (participant) age groups and aggregated over the corresponding *column* (contact) age groups.

Author(s)

Sebastian Meyer

Source

contactmatrix_mossong and contactmatrix_mossong_physical are taken from the Supporting Information in Mossong et al. (2008): the matrices from Table S5 (8.2), and the attached age distribution from Table S2 (3.2).

The corrected versions contactmatrix_POLYMOD and contactmatrix_POLYMOD_physical were constructed from the raw POLYMOD data available at https://www.researchgate.net/publication/232701632_POLYMOD_contact_survey_for_researchers. The reciprocal contact matrices contactmatrix_wallinga and contactmatrix_wallinga_physical were estimated from these raw data via the method of Wallinga et al. (2006).

References

Meyer S and Held L (2017): Incorporating social contact data in spatio-temporal models for infectious disease spread. *Biostatistics*, **18** (2), 338-351. doi: 10.1093/biostatistics/kxw051

Mossong et al. (2008): Social contacts and mixing patterns relevant to the spread of infectious diseases. *PLoS Medicine*, **5** (3), e74. doi: 10.1371/journal.pmed.0050074

Wallinga J, Teunis P and Kretzschmar M (2006): Using data on social contacts to estimate age-specific transmission parameters for respiratory-spread infectious agents. *American Journal of Epidemiology*, **164** (10), 936-944. doi: 10.1093/aje/kwj317

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```
## contact matrix reported in Mossong et al (2008, Table S5)
(C_original <- contactmatrix(which = "mossong", grouping = NULL))</pre>
## this simply returns the dataset 'contactmatrix_mossong'
stopifnot(identical(C_original, contactmatrix_mossong))
## with corrected numbers for the 70+ age group (the default)
C_corrected <- contactmatrix(which = "corrected", grouping = NULL)</pre>
## this simply returns the dataset 'contactmatrix_POLYMOD'
stopifnot(identical(C_corrected, contactmatrix_POLYMOD))
## check for differences
C_original == round(C_corrected, 2)
## compare entries of last row and last column
round(rbind(original = C_original[15,], corrected = C_corrected[15,]), 2)
round(cbind(original = C_original[,15], corrected = C_corrected[,15]), 2)
## contact matrix estimated to be reciprocal on the population level
C_reciprocal <- contactmatrix(which = "reciprocal", grouping = NULL)</pre>
## this simply returns the dataset 'contactmatrix_wallinga'
## (without its "overdisp" attribute)
stopifnot(all.equal(C_reciprocal, contactmatrix_wallinga, check.attributes=FALSE))
## check reciprocity
agedistriBE <- attr(C_reciprocal, "agedistri")</pre>
stopifnot(identical(agedistriBE, prop.table(colSums(pop2011))))
stopifnot(isSymmetric(C_reciprocal * agedistriBE, check.attributes=FALSE))
## visually compare raw to reciprocal contact matrix
if (require("gridExtra"))
    grid.arrange(plotC(C_corrected, main = "raw"),
                 plotC(C_reciprocal, main = "reciprocal"),
                 nrow = 1
## select physical contacts and aggregate into 5 age groups
contactmatrix(type = "physical", grouping = c(1, 2, 7, 3, 2))
## the default 6 age groups, normalized to a transition matrix
contactmatrix(normalize = TRUE)
## reciprocity also holds for this grouping
(C6 <- contactmatrix(which = "reciprocal"))</pre>
stopifnot(isSymmetric(C6 * attr(C6, "agedistri"), check.attributes=FALSE))
```

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Description

The expectation and variance of aggregated predictions is just a sum if the predictions are (conditionally) independent. This function computes the DSS for a matrix of observations and a matrix of predictions where the columns are to be summed according to a given factor.

Usage

```
dssAggregate(observed, pred, psi, groups)
```

Arguments

observed a numeric matrix of observed counts.

pred a numeric matrix of predicted counts.

psi a numeric vector or matrix of overdispersion parameters such that pred * (1 +

pred/exp(psi)) is the prediction's variance. Alternatively, psi = NULL indi-

cated Poisson predictions.

groups a factor variable of length ncol(observed) indicating which columns should

be aggregated.

Value

a matrix of DSS values

expandC

Expand the Contact Matrix over Regions

Description

This is simply the Kronecker product of the contact matrix C with a matrix of ones of dimension n x n

Usage

```
expandC(C, n)
```

Arguments

C a contactmatrix.

n the size of the secondary dimension to expand to.

Value

a square matrix with n*ncol(C) rows and columns.

```
expandC(contactmatrix(), 2)
```

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fitC

Estimate the Power of the Contact Matrix in a "hhh4" Model

Description

The profile log-likelihood of the log(power) parameter of the contact matrix (see powerC) is maximized using optim. The hhh4 fit for the optimal power value is returned with an additional element logpower which holds information on the result of the optimization.

Usage

```
fitC(object, C, normalize = TRUE, truncate = TRUE, optim.args = list(), ...)
```

Arguments

Value

an object of class "fitC", which is an "hhh4" object with an additional element logpower.

Author(s)

Sebastian Meyer

noroBE

Create "sts" Objects from the Berlin Norovirus Data

Description

The function noroBE() creates an "sts" object based on the array of norovirus surveillance counts, the map of Berlin's city district, and the pop2011 data stored in the package. This is the data analysed by Meyer and Held (2017).

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Usage

```
noroBE(
   by = c("districts", "agegroups", "all", "none"),
   agegroups = c(1, 2, 2, 4, 4, 2),
   timeRange = c("2011-w27", "2015-w26"),
   flatten = FALSE
)

counts
map
```

Arguments

by

character string determining the stratification, i.e., which units the resulting "sts" object should represent:

"districts": aggregates counts and pop2011 over the age groups and stores the matrix of adjacency orders from the map in the neighbourhood slot. The latter is obtained via nbOrder(poly2adjmat(map), maxlag = 5).

"agegroups": aggregates counts and pop2011 over the districts and stores the contactmatrix() in the neighbourhood slot, potentially also combining some age groups via the agegroups argument.

"all": retains both dimensions, either as a list of spatial "sts" objects per age group, or in a single "sts" object (see flatten below).

"none": creates the overall (univariate) time series of rowSums(counts).

agegroups

how the age groups in counts (and pop2011) should be aggregated. Will be used as the grouping argument in aggregateCountsArray and contactmatrix. The default setting uses the six age groups of Meyer and Held (2017).

timeRange

character vector of length two determining the time range of the "sts" object to generate. The two strings are matched against dimnames(counts)[[1]], which ranges from "2011-w01" until "2016-w30". The default value extracts four seasons (years) starting at "2011-w27".

flatten

logical indicating whether for by = "all" a single "sts" object should be returned where the observation unit is the interaction of district and age group ("flattened" counts array, see as.data.frame.array). By default (flatten = FALSE), a list of district-based "sts" objects is returned, one for each age group.

Format

counts: an integer-valued array of norovirus surveillance counts with labelled dimensions of size 290 ("week") x 12 ("district") x 15 ("agegroup").

map: a "SpatialPolygonsDataFrame" of length 12 with row.names(map) matching colnames(counts), representing Berlin's city districts in longlat coordinates (WGS84). The data slot contains the full "NAME"s of the city districts as well as their "POPULATION", i.e., rowSums(pop2011).

The function noroBE() returns an "sts" object generated from these data (and pop2011).

noroBE

Author(s)

Sebastian Meyer

Source

counts: based on norovirus surveillance counts retrieved from the SurvStat@RKI 2.0 online service (https://survstat.rki.de) of Germany's public health institute, the Robert Koch Institute, as of 2016-09-08.

map: based on a KML file of Berlin's 97 local centres ("Ortsteile") downloaded from the Berlin Open Data repository at https://daten.berlin.de/datensaetze/geometrien-der-ortsteile-von-berlin-jul as of 2014-11-12, published by Amt fuer Statistik Berlin-Brandenburg (Statistical Office of Berlin-Brandenburg) under the 'CC BY 3.0 DE' license (https://creativecommons.org/licenses/by/3.0/de/). The map included here is a gUnaryUnion of these local centres by city district.

References

Meyer S and Held L (2017): Incorporating social contact data in spatio-temporal models for infectious disease spread. *Biostatistics*, **18** (2), 338-351. doi: 10.1093/biostatistics/kxw051

```
## the raw data
str(counts)
summary(map)
## district-specific time series
noroBEr <- noroBE(by = "districts")</pre>
plot(noroBEr)
## age group-specific time series
noroBEg <- noroBE(by = "agegroups")</pre>
plot(noroBEg)
## list of spatio-temporal surveillance counts, one for each age group
noroBErbyg <- noroBE(by = "all", flatten = FALSE)</pre>
plot(noroBErbyg[[1L]], par.list = list(oma=c(0,0,2,0)))
title(main = names(noroBErbyg)[1], outer = TRUE, line = -1)
## flattened "sts" object (the 'neighbourhood' only reflects spatial info)
noroBEall <- noroBE(by = "all", flatten = TRUE)</pre>
dev.new(width = 16, height = 7)
plot(noroBEall, par.list = list(
    xaxt = "n", mar = c(1,4,1,1), mfrow = c(ncol(noroBEg), ncol(noroBEr))
))
```

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plotC

Generate an Image of a Contact Matrix

Description

Generate an Image of a Contact Matrix

Usage

```
plotC(
   C,
   grouping = NULL,
   xlab = "age group of contact",
   ylab = "age group of participant",
   at = 15,
   col.regions = rev(heat.colors(length(at) - 1)),
   ...,
   contour = FALSE
)
```

Arguments

С a square numeric matrix. grouping numeric vector of sizes of aggregated groups, e.g., grouping = c(1,3), to draw separation lines after the first and the forth subgroup. This is ignored if contour = TRUE. axis labels. xlab, ylab numeric vector of break points of the color levels, or a single integer specifying at the number of cuts (which defaults to 15 as in levelplot). col.regions vector of color levels. further arguments passed to levelplot or filled. contour (if contour = TRUE). logical indicating if a filled. contour should be drawn instead of a levelplot contour (the default).

```
plotHHH4_fitted_groups
```

Plot Mean Components of a hhh4 Fit by Group

Description

Fitted mean components for age-structured, areal time series hhh4 models can be aggregated over districts or age groups.

Usage

```
plotHHH4_fitted_groups(x, groups, total = FALSE, decompose = NULL, ...)
```

Arguments

```
x an object of class "hhh4".
```

groups a factor of grouping the units in the model, i.e., it must be of length x\$nUnit.

There will be one plot for each factor level.

total a logical indicating if the group-wise mean components should be subsequently

summed up over all groups for an overall plot.

decompose, ... see plotHHH4_fitted.

Value

```
see plotHHH4_fitted.
```

Description

This is a wrapper for plotHHH4_maps with prior aggregation over different (age) groups.

Usage

```
plotHHH4_maps_groups(x, map, districts, ...)
```

Arguments

```
x an object of class "hhh4".
```

map an object inheriting from "SpatialPolygons".

districts a factor of length x\$nUnit with as many levels as there are districts and names

according to row.names(map).

... arguments passed to plotHHH4_maps.

Value

```
see plotHHH4_maps
```

```
plotHHH4_season_groups
```

Plot Seasonality of a hhh4 Fit by Group

Description

A plot method for models with group-specific seasonality terms that are not handled correctly by plotHHH4_season.

Usage

```
plotHHH4_season_groups(
    x,
    component = "end",
    seasonStart = 1,
    conf.level = 0.95,
    conf.B = 999,
    col = 1:6,
    xlab = "time",
    ylab = "multiplicative effect",
    ...,
    refline.args = list(),
    yearline.args = list(),
    legend.args = list()
)
```

Arguments

x an object of class "hhh4".

component character string indicating from which component seasonality terms should be

extracted.

seasonStart an integer defining the epochInYear that starts a new season (by default the

first).

conf.level, conf.B

a confidence level for the pointwise confidence intervals around the group-specific seasonal effects. The confidence intervals are based on quantiles of conf.B samples from the asymptotic multivariate normal distribution of the maximum likelihood estimate. Alternatively, if conf.level = NA, the individual samples are drawn instead of the confidence lines. Set conf.level = NULL to disable

confidence intervals.

col a vector of group-specific colors, recycled as necessary and passed to matplot.

xlab, ylab, ... arguments passed to matplot.

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refline.args a list of arguments for abline to change the style of the horizontal reference

line at 1. This line is omitted if refline.args is not a list.

yearline.args a list of arguments for abline to change the style of the line marking the end

of the year at x\$stsObj@freq if seasonStart is not 1. This line is omitted if

yearline.args is not a list.

legend.args a list of arguments for legend modifying the internal defaults. If legend.args

is not a list, the legend is omitted.

Value

a matrix of the plotted point estimates of the multiplicative seasonal effect by group.

pop2011

Berlin and German Population by Age Group, 2011

Description

Population numbers from Berlin are available in the city district x age group (5-year intervals) matrix pop2011. The corresponding age distribution for whole Germany is stored in the vector popDE.

Usage

```
## Berlin population by city district and age group, 2011
pop2011
## German population by age group, 2011
popDE
```

Format

pop2011: a named, integer-valued 12 (city districts) x 15 (age groups) matrix. **popDE:** a named integer vector of length 15 (age groups).

Author(s)

Sebastian Meyer

Source

pop2011: numbers extracted from https://www.statistik-berlin-brandenburg.de/webapi/ opendatabase?id=BevBBBE as of 2011-12-31 (before census), published by Amt fuer Statistik Berlin-Brandenburg (Statistical Office of Berlin-Brandenburg) under the 'CC BY 3.0 DE' license (https://creativecommons.org/licenses/by/3.0/de/).

popDE: numbers extracted from https://www-genesis.destatis.de/genesis/online/link/tabellen/12411-0005 as of 2010-12-31, published by *Statistisches Bundesamt* (Destatis, Federal Statistical Office of Germany) under the 'Data licence Germany - attribution - Version 2.0' (https://www.govdata.de/dl-de/by-2-0).

18 powerC

powerC

Exponentiate a Matrix via Eigendecomposition

Description

Based on a (contact) matrix C, the function make_powerC generates a function with a single argument power that returns the input matrix raised to that power. Matrix exponentiation is thereby defined via the eigendecomposition of C as $C^{power} := E\Lambda^{power}E^{-1}$.

Usage

```
make_powerC(C, normalize = FALSE, truncate = FALSE)
```

Arguments

C a square numeric matrix.

normalize a logical indicating if C should be normalized in advance such that all rows sum

to 1 (becomes a transition matrix).

truncate a logical indicating whether to force entries in the resulting matrix to be non-

negative (by truncation at 0).

Value

a function of the power that returns the exponentiated matrix.

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stationary

Stationary Distribution of a Transition Matrix

Description

This auxiliary function determines the stationary distribution from a transition matrix.

Usage

```
stationary(P)
```

Arguments

Ρ

a transition matrix, i.e., a square matrix where all rows sum to 1.

Value

the stationary probability vector.

Author(s)

Leonhard Held

Examples

```
Cgrouped_norm <- contactmatrix(normalize = TRUE)
Cgrouped_norm
(p <- stationary(Cgrouped_norm))
(Cpowered <- make_powerC(Cgrouped_norm)(1e6))
stopifnot(all.equal(Cpowered[1,], p))</pre>
```

stratum

Extract Strata

Description

Methods to extract strata information from an object. Here we only define a method for class "sts".

Usage

```
stratum(x, ...)
## S4 method for signature 'sts'
stratum(x, which = NULL, ...)
```

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Arguments

```
x an object of class "sts".... further arguments passed to methods.which an integer (strata dimension) or NULL (to get the plain colnames, the default).
```

Value

a character vector of strata names of length ncol(x).

Methods (by class)

• sts: Extract the names of the units, i.e., the colnames, from a multivariate "sts" object. If the units result from the interaction of multiple strata separated by dots, e.g., "region.group", the function can also extract the names corresponding to a specific strata dimension, e.g., which = 2 to get the group names.

Examples

```
noroBEall <- noroBE(by = "all", flatten = TRUE)
stratum(noroBEall)  # just colnames(noroBEall)
stratum(noroBEall, which = 2)  # the age groups</pre>
```

stsplothook

Hook functions for stsplot_time1

Description

Hook functions can be passed to stsplot_time1, which are evaluated after all the plotting has been done, and with the hook function environment set to the evaluation environment of stsplot_time1 such that local variables can be accessed. They are not intended to be called directly.

Usage

```
stsplothook_highlight(christmas = FALSE, epochInYear = NULL, col = 2, lwd = 2)
```

Arguments

christmas logical indicating if Christmas should be highlighted.

epochInYear integer vector of epochs to highlight.

col, lwd graphical parameters for the highlighting lines.

Author(s)

Sebastian Meyer

```
plot(noroBE("agegroups"), hookFunc = stsplothook_highlight(epochInYear=51))
```

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