Package ‘DendroSync’

July 16, 2018

Type Package
Title A Set of Tools for Calculating Spatial Synchrony Between Tree-Ring Chronologies
Version 0.1.2
Date 2018-07-16
Author Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas
Maintainer Josu G. Alday <josucham@gmail.com>
Description Provides functions for the calculation and plotting of synchrony in tree growth from tree-ring width chronologies (TRW index). It combines variance-covariance (VCOV) mixed modelling with functions that quantify the degree to which the TRW chronologies contain a common temporal signal. It also implements temporal trends in spatial synchrony using a moving window. These methods can also be used with other kind of ecological variables that have temporal autocorrelation corrected.
License GPL-2
Depends R (>= 3.1.2), nlme, ggplot2
LazyData TRUE
Imports stats, utils, gridExtra
RoxygenNote 6.0.1
NeedsCompilation no
Repository CRAN
Date/Publication 2018-07-16 18:20:03 UTC

R topics documented:

bet.aSE ................................................................. 2
bet.het.aSE ............................................................ 3
conifersIP ............................................................. 5
csbet.aSE ............................................................. 6
csbet.het.aSE .......................................................... 7
cswi.aSE .............................................................. 8
Description

The function calculates the between-group synchrony ($a^b$) and standard error (SE) for a homoscedastic unstructured model or full model (mUN).

Usage

```
bet.aSE(model)
```

Arguments

- `model` : a class "lme" unstructured model (mUN) produced by `dendro.varcov` with homoscedastic equals TRUE.

Details

The function calculates between-group synchrony for a homoscedastic unstructured model (mUN).

Value

The function returns a matrix containing between-group synchrony and SE for each combination of varGroup levels. This function is used internally in `sync`.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas
bet.het.aSE

References


See Also

`sync` for a clear description of synchrony evaluation.

Examples

```r
## Calculate between-group synchrony and SE
# For an unstructured homoscedastic model of conifersIP data:
data(conifersIP)

# Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using taxonomic grouping criteria (i.e. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = TRUE)
summary(ModHm)

# Obtain the unstructured model between-group synchrony and SE
# For each varGroup stratum.
bet.aSE(ModHm$mUN) # Unstructure
```

---

**bet.het.aSE**

*Between-group synchrony for a heteroscedastic unstructured model*

Description

The function calculates the between-group synchrony ($a^g$) and standard error (SE) for a heteroscedastic unstructured model (mHeUN).

Usage

`bet.het.aSE(model)`

Arguments

- `model` a class "lm" unstructured model (mHeUN) produced by `dendro.varcov` with homoscedastic equals FALSE.
Details

The function calculates between-group synchrony for a heteroscedastic unstructured model (mHeUN).

Value

The function returns a matrix containing between-group synchrony and SE for each combination of varGroup levels. This function is used internally in `sync`.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

References


See Also

`sync` for a clear description of synchrony evaluation.

Examples

```r
## Calculate between-group synchrony and SE
##for a heteroscedastic unstructured model for conifersIP data:
data(conifersIP)

# Fit the heteroscedastic set of varcov models (mBE, mHeNE, mHeCS, mHeUN)
# using taxonomic grouping criteria (ie. Species)
ModHt <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = FALSE)

# between-group synchrony and SE for each varGroup stratum combination
# in heteroscedastic unstructured models.
bet.het.aSE(ModHt$mHeUN)#Unstructured model
```
Iberian Peninsula tree-ring width dataset

Description

This dataset presents tree-ring width chronologies of residual indices published in Shestakova et al. (2016). The dataset contains 30 tree-ring width chronologies of conifer species from the Iberian Peninsula for the period 1950-1999. Tree species are represented by *Abies alba* Mill., *Pinus nigra* subsp. *salzmannii* (Dunal) Franco and *Pinus sylvestris* L. The sampling sites are distributed across three subregions (north, centre and south) of the Iberian Peninsula. Site chronologies are obtained by cross-dating tree-ring width series and posterior detrending and autocorrelation removal with the Friedman supersmoother spline and autoregressive modeling.

Usage

data(conifersIP)

Format

A `data.frame` containing 30 tree-ring chronologies of residual indices for the period 1950-1999. The dataset includes three species (*Abies alba*, *Pinus nigra*, *Pinus sylvestris*) that are distributed across three regions (north, centre and south) in accordance with the latitudinal position of each sampling site. `conifersIP` has following variables:

- **Year**: year of tree-ring formation (1950-1999)
- **Region**: geographical region for Iberian Peninsula: north, centre, south
- **Species**: tree species: *Abies alba*, *Pinus nigra*, *Pinus sylvestris*
- **Code**: specific name of tree-ring width chronology
- **TRW**: residual indices of tree-ring width chronologies

References

csbet.aSE  

**Between-group synchrony for homoscedastic compound symmetry model**

**Description**

The function calculates the between-group synchrony ($a^\circ$) and standard error (SE) for homoscedastic compound symmetry model (mCS).

**Usage**

```r
csbet.aSE(model)
```

**Arguments**

- `model`: a class "lme" compound symmetry model (mCS) produced by `dendro.varcov` with homoscedastic equals TRUE.

**Details**

The function calculates between-group synchrony for homoscedastic compound symmetry model (mCS).

**Value**

The function returns a matrix containing between-group synchrony and SE for each combination of varGroup levels. This function is used internally in `sync`.

**Author(s)**

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

**References**


**See Also**

`sync` for a clear description of synchrony evaluation.
Examples

```r
## Calculate between-group homoscedastic synchrony and SE
# for compound symmetry homoscedastic model of conifersIP data:
data(conifersIP)

# Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using taxonomic grouping criteria (ie. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = TRUE)
summary(ModHm)

# Obtain the compound symmetry model between-group synchrony and SE
# for each varGroup stratum.
csbet.aSE(ModHm$mCS)# compound symmetry
```

---

**csbet.het.aSE**

*Between-group synchrony for heteroscedastic compound symmetry model*

---

**Description**

The function calculates the between-group synchrony ($a^\gamma$) and standard error (SE) for heteroscedastic compound symmetry model (mHeCS).

**Usage**

```r
csbet.het.aSE(model)
```

**Arguments**

- `model` a class "lme" compound symmetry model (mHeCS) produced by `dendro.varcov` with homoscedastic equals FALSE.

**Details**

The function calculates between-group synchrony for heteroscedastic compound symmetry models (mHeCS).

**Value**

The function returns a matrix containing between-group synchrony and SE for each combination of varGroup levels. This function is used internally in `sync`.

**Author(s)**

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas
References


See Also

*sync* for a clear description of synchrony evaluation.

Examples

```r
## Calculate between-group synchrony and SE
## For heteroscedastic compound symmetry model for conifersIP data:
data(conifersIP)

# Fit the heteroscedastic set of varcov models (mBE, mHeNE, mHeCS, mHeUN)
# Using taxonomic grouping criteria (i.e. Species)
modHt <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = FALSE)

# Between-group synchrony and SE for each varGroup stratum combination
# In heteroscedastic compound symmetry models.
csbet.het.aSE(modHt$mHeCS)
```

---

**cswi.aSE**

Within-group synchrony for homoscedastic compound symmetry model

Description

The function calculates the within-group synchrony (α^) and standard error (SE) for homoscedastic compound symmetry models (mCS).

Usage

`cswi.aSE(model)`

Arguments

- `model` a class "lme" compound symmetry model (mCS) produced by `dendro.varcov` with homoscedastic equals TRUE.
Details

The function calculates within-group synchrony for homoscedastic compound symmetry model (mCS).

Value

The function returns a matrix containing within-group synchrony and SE for each combinations of varGroup levels. This function is used internally in sync.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

References


See Also

sync for a clear description of synchrony evaluation.

Examples

```R
# Calculate within-group homoscedastic synchrony and SE
# for compound symmetry homoscedastic model of conifersIP data:
data(conifersIP)

# Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using taxonomic grouping criteria (ie. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = TRUE)
summary(ModHm)

# Obtain the compound symmetry model within-group synchrony and SE
# for each varGroup stratum.
cswi.aSE(ModHm$mCS) # compound symmetry
```
cswi.het.aSE

Within-group synchrony for heteroscedastic compound symmetry mixed models

Description

The function calculates for each varGroup stratum the synchrony (a^) and standard error (SE) for heteroscedastic compound symmetry models (mHeCS).

Usage

cswi.het.aSE (model)

Arguments

model: a class "lme" model produced by dendro.varcov with homoscedastic equals FALSE.

Details

The function calculates the within-group synchrony values for each varGroup stratum based on the methodology described in Shestakova et al. (2014) and Shestakova et al. (2016). Note that this function is designed to work only in heteroscedastic compound symmetry mixed models (i.e. mHeCS).

Value

The function returns a matrix containing within-group synchrony and SE for each level of varGroup. This function is used internally in sync.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

References


See Also

sync for a clear description of synchrony evaluation.
Examples

```r
## Calculate within-group heteroscedastic synchrony and SE for conifersIP data:
data(conifersIP)

# Fit the heteroscedastic set of varcov models (mBE, mHeNE, mHeCS, mHeUN)
# using taxonomic grouping criteria (i.e. Species)
ModHt <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = FALSE)
summary(ModHt)

# Obtain the heteroscedastic compound symmetry
# within-group synchrony and SE for each varGroup stratum.
cswi.het.aSE(ModHt$mHeCS)
```

---

dendro.varcov  
Fit variance-covariance mixed models on tree-ring chronologies

Description

The function calculates variance-covariance (VCOV) mixed models from a data.frame with tree-ring width index and years for each chronology following the methodology described in Shestakova et al. (2014). The mixed models relate tree-ring width (Y) against specific names of tree-ring width chronologies (A), using years and grouping variable as random factors to characterize the strength of the common signal across the grouping variable. First, a linear mixed-effect model with null positive-definite matrix structure or broad evaluation is fitted and the subsequent models are consequently derived from it using the function update. When a data.frame with tree-ring width index has NAs the models are fitted with na.action = na.omit. Simultaneously, complete.cases is applied to guarantee that rows have no missing values across the entire data.frame.

Usage

```r
dendro.varcov(formula, varTime = "", varGroup = "", data,
               homoscedastic = TRUE, null.mod = FALSE, all.mod = FALSE)
```

Arguments

- **formula**: a model formula such as Y ~ A, where Y is usually tree-ring width and A will be a factor variable such as the specific names of tree-ring width chronologies (conifersIP) or ~1 for a null model.
- **varTime**: a character specifying the time variable to consider in calculating synchrony estimates. Models with varTime variable with less than 10 different time-points produce unreliable results.
- **varGroup**: a character grouping variable. In dendrochronological studies different grouping strategies can be used. We used here two strategies following taxonomic (i.e. species) or geographic (i.e. region) criteria.
data a data.frame with tree-ring chronologies, years and grouping variables as columns.
homoscedastic logical if TRUE models do not included an optional varFunc object. If FALSE models will include a one-sided formula describing the within-group heteroscedasticity structure (varIdent).
null.mod logical if TRUE only broad evaluation model will be fitted. Default FALSE.
all.mod logical if TRUE all homoscedastic and heteroscedastic model types will be fitted. Default FALSE.

Details

The function fits a set of variance-covariance mixed models following Shestakova et al. (2014). A total of 7 different variance-covariance mixed models can be fitted: a null positive-definite matrix structure (mBE), and the homoscedastic and heteroscedastic versions of a diagonal positive-definite matrix structure (mNE, mHeNE), a positive-definite matrix with compound symmetry structure (mCS, mHeCS) and a general positive-definite matrix structure (mUN, mHeUN). Note that if null.mod is TRUE the function only fits broad evaluation model (mBE), this is set to FALSE by default. If all.mod is TRUE the function fits heteroscedastic and homoscedastic versions of all models. This is set to FALSE by default, because for large-datasets it may take a long time to converge.

Value

The function returns a list containing the following components:

• for null.mod = TRUE:
  
mBE an object of class "lme" representing the linear mixed-effects model fit of null positive-definite matrix structure or broad evaluation. See lmeObject for the components of the fit.

• for homoscedastic = TRUE:
  
mNE an object of class "lme" representing the linear mixed-effects model fit of a diagonal positive-definite matrix structure or narrow evaluation. See lmeObject for the components of the fit.
mCS an object of class "lme" representing the linear mixed-effects model fit of a positive-definite matrix with compound symmetry structure. See lmeObject for the components of the fit.
mUN an object of class "lme" representing the linear mixed-effects model fit of a general positive-definite matrix structure or unstructured. See lmeObject for the components of the fit.

• for homoscedastic = FALSE:
  
mHeNE an object of class "lme" representing the linear mixed-effects model fit of the heteroscedastic variant of a diagonal positive-definite matrix structure or narrow evaluation. See lmeObject for the components of the fit.
mHeCS an object of class "lme" representing the linear mixed-effects model fit of the heteroscedastic variant of a positive-definite matrix with compound symmetry structure. See lmeObject for the components of the fit.
mHeUN

an object of class "lme" representing the linear mixed-effects model fit of the heteroscedastic variant of a general positive-definite matrix structure or unstructured. See lmeObject for the components of the fit.

- for all.mod = TRUE:

  The function returns the homoscedastic and heteroscedastic versions of all fitted models.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

References


See Also

lmeObject, na.action, complete.cases

Examples

```r
## Calculate variance-covariance models on Iberian Peninsula conifers
# chronologies using two different grouping strategies.
# Tree-ring width chronologies are grouped according to taxonomic (i.e. Species)
# or geographic (i.e. Region) criteria.
# User-defined homoscedastic or heteroscedastic variances can be fitted.
data(conifersIP)

#Chop the data from 1960 to 1989.
conif.30 <- conifersIP[conifersIP$Year>1959 & conifersIP$Year<1990,]
summary(conif.30$Year)

##Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using taxonomic grouping criteria (ie. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
                      data = conif.30, homoscedastic = TRUE)

summary(ModHm)# Class and length of list elements
ModHm
ModHm[2]#mNE fitted model results

##Fit the heteroscedastic set of varcov models (mBE, mHeNE, mHeCS, mHeUN)
# using geographic grouping criteria (ie. Region)
```
Description

The function calculates for each varGroup stratum the within-group synchrony (\(a^g\)) and standard error (SE). However, it only works for homoscedastic broad evaluation, narrow evaluation and unstructured models (mBE, mNE, mUN).

Usage

```r
gen.aSE(model)
```

Arguments

- `model`: a class "lme" model produced by `dendro.varcov` with homoscedastic equals TRUE.

Details

The function calculates the within-group synchrony values for each varGroup stratum based on the methodology described in Shestakova et al. (2014) and Shestakova et al. (2016). Note that this function is designed to work only in 3 homoscedastic models (mBE, mNE, mUN).

Value

The function returns a matrix containing the synchrony and SE for each level of varGroup. This function is used internally in `sync`.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas
References

See Also
`sync` for a clear description of synchrony evaluation.

Examples
```r
# Calculate within-group homoscedastic synchrony and SE for conifersIP data:
data(conifersIP)

# Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN) # using taxonomic grouping criteria (ie. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
                      data = conifersIP, homoscedastic = TRUE)
supmary(ModHm)

# Obtain the within-group synchrony and SE for each varGroup stratum.
gen.aSE(ModHm$mBE)# Broad evaluation
ngen.aSE(ModHm$mUN)# Unstructured
```

**Description**
The function calculates for each varGroup stratum the synchrony (a^) and standard error (SE), but only for heteroscedastic unstructured and narrow evaluation mixed models (mHeNE, mHeUN).

**Usage**
`gen.het.aSE (model)`

**Arguments**
- `model` a class "lme" model produced by `dendro.varcov` with homoscedastic equals FALSE.
Details

The function calculates the within-group synchrony values for each varGroup stratum based on the methodology described in Shestakova et al. (2014) and Shestakova et al. (2016). Note that this function is designed to work only in heteroscedastic narrow evaluation and unstructured mixed models (i.e. mHeNE and mHeUN).

Value

The function returns a matrix containing within-group synchrony and SE for each level of varGroup. This function is used internally in sync.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

References


See Also

sync for a clear description of synchrony evaluation.

Examples

```r
## Calculate within-group heteroscedastic synchrony and SE for conifersIP data:
data(conifersIP)

## Fit the heteroscedastic set of varcov models (mBE, mHeNE, mHeCS, mHeUN)
## using taxonomic grouping criteria (i.e. Species)
ModHt <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
                        data = conifersIP, homoscedastic = FALSE)
summary(ModHt)

## Obtain the heteroscedastic within-group synchrony and SE for each varGroup stratum.
gen.het.aSE(ModHt$mHeNE)#Narrow evaluation model
gen.het.aSE(ModHt$mHeUN)#Unstructured model
```
het.var

Variance per varGroup stratum for heteroscedastic models

Description
The function obtains the heteroscedastic variances for each varGroup elements for a selected model (mHeCS, mHeUN, mHeNE).

Usage
het.var(model)

Arguments

model a class "lme" model produced by dendro.varcov with homoscedastic equals FALSE.

Details
The function extracts the variances for each varGroup stratum using the within-group heteroscedastic structure of the fitted models (varIdent constant variance per stratum). Note that this function only works for heteroscedastic models: mHeCS, mHeNE, mHeUN.

Value
The function returns a numeric vector containing the variance per each level of varGroup. They are used internally to calculate synchrony (sync).

Author(s)
Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

Examples
```r
## Calculate within-group heteroscedastic variances for conifersIP data:
data(conifersIP)

# Fit the heteroscedastic set of models (mBE, mHeCS, mHeNE, mHeUN) # using taxonomic grouping criteria (i.e. Species)
ModHt <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, homoscedastic = FALSE)

# Obtain the within-group variances for the model of interest
het.var(ModHt$mHeCS)# Heterogeneous variant of compound symmetry model
het.var(ModHt$mHeUN)# Heterogeneous unstructured model
```
mod.table

Function to calculate goodness-of-fit statistics for variance-covariance models

Description

The function provides a table to compare fitted variance-covariance (VCOV) mixed models by AIC, AICc, BIC and LogLik. The restricted log-likelihood (LogLik) statistics for different models can be compared by Chi-square test, while Akaike information criterion (AIC), corrected Akaike information criterion (AICc) and Bayesian information criterion (BIC) are in the smaller-is-better form.

Usage

mod.table(modelList)

Arguments

- modelList: a list of variance-covariance (VCOV) mixed models of the type as produced by dendro.varcov.

Details

The function returns a table to compare the fitted variance-covariance (VCOV) mixed models to the same data based on information criteria. The smaller AIC, AICc or BIC, the better fit. Also, LogLik value is included. AICc is calculated according to the formula AIC + 2*npar*(nobs/(nobs-npar-2)), where npar represents the number of parameters and nobs the number of observations in the fitted model.

Value

The function returns a data.frame with rows corresponding to the objects and columns containing the following components:

- n: the number of observations used in the model fit.
- df: the number of parameters in the fitted model.
- AIC: Akaike’s Information Criterion of the fitted model.
- AICc: corrected Akaike’s Information Criterion of the fitted model.
- BIC: Bayesian Information Criterion of the fitted model.
- LogLik: log-likelihood of the fitted model

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Volta
References


See Also

AIC, BIC, logLik

Examples

```r
## Compare homoscedastic variance-covariance models on Iberian Peninsula
# conifer ring chronologies using taxonomic grouping criteria (i.e. Species).

data(conifersIP)
ModHmSp <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
                           data = conifersIP, homoscedastic = TRUE)

mod.table(ModHmSp)# a data.frame containing information criterion values

## Compare homoscedastic variance-covariance models on Iberian Peninsula conifers
# ring chronologies using geographic criteria (i.e. Region).

ModHmGoe <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Region",
                           data = conifersIP, homoscedastic = TRUE)

mod.table(ModHmGoe)
```

---

**sync**

Calculate within- and between-group synchrony

Description

The function calculates spatial synchrony from a list of fitted mixed models with variance-covariance structures of the type as produced by `dendro.varcov`. Within- and between-group synchrony are calculated, quantifying the degree to which the values of N chronologies contain a common temporal signal. Different models allow for the estimation of intraclass correlations either at the intragroup or intergroup level. The underlying idea is to split the mean correlation estimated between all possible pairs of chronologies drawn from the whole dataset into: (i) a mean correlation between pairs of chronologies for every group; and (ii) a mean correlation between pairs of chronologies for pairs of groups.

Usage

```r
sync (modellist, modname = c("mBE", "mNE", "mCS", "mUN", "mHeNE",
                           "mHeCS", "mHeUN"), trend.mBE = FALSE)
```
Arguments

modelist a list of the type as produced by `dendro.varcov`.

modname a character string of "mBE", "mNE", "mCS", "mUN", "mHeNE", "mHeCS" or "mHeUN", specifying the variance-covariance structures selected for synchrony evaluation.

trend.mBE a logical specifying if a broad evaluation model (mBE) output for each grouping level is reported. This is a special mBE output to plot synchrony trends with `sync.trend.plot`. Default FALSE.

Details

The function calculates the within- and between-group synchrony. For the more general (unstructured) model, the correlation of pairs of chronologies \(i\) and \(i^*\) belonging to group \(r\) is:

\[
\rho(W_i, W_i^*) = \frac{\text{cov}(W_i, W_i^*)}{\sqrt{\text{Var}(W_i) \times \text{Var}(W_i^*)}} = \frac{\sigma_{yr}^2}{\sqrt{\sigma_{yr}^2 + \sigma_e^2}}
\]

Where \(W_i\) is tree-ring width of \(i\)th chronology, \(\sigma_{yr}^2\) is a covariance between observations \(W_i\) and \(W_i^*\) belonging to a group \(r\), \(\sigma_e^2\) is a random deviation within the \(r\)th group. Conversely, the correlation of pairs of chronologies \(i\) and \(i^*\) belonging to groups \(r\) and \(r^*\) is:

\[
\rho(W_i, W_i^*) = \frac{\text{cov}(W_i, W_i^*)}{\sqrt{\text{Var}(W_i) \times \text{Var}(W_i^*)}} = \frac{\sigma_{yr}^*}{\sqrt{\left(\sigma_{yr}^2 + \sigma_e^2\right) + \left(\sigma_{yr}^2 + \sigma_e^2\right)}}
\]

Note that if no `modname` is provided a warning message appears indicating that synchrony will be only calculated for the first `modname` vector element, i.e. broad evaluation model (mBE).

Value

The function returns a list containing the following components:

- for within-group synchrony:

  a column indicating the variance-covariance mixed models fit type:

  - mBE: null (or broad evaluation) structure.
  - mNE: homoscedastic variant of banded main diagonal (or narrow evaluation) structure.
  - mCS: homoscedastic variant of compound symmetry structure.
  - mUN: homoscedastic variant of unstructured (or full) structure.
  - mHeNE: heteroscedastic variant of banded main diagonal (or narrow evaluation) structure.
  - mHeCS: heteroscedastic variant of compound symmetry structure.
  - mHeUN: heteroscedastic variant of unstructured (or full) structure.

- a_group a column representing the within-group synchrony.

- SE_Group standard error of each observation.
for between-group synchrony:

- `ModName` a column indicating the model fit type. See previous description.
- `Groupname` a column indicating between-group varGroup pairwise combinations r and r*.
- `a_betw_Grp` a column indicating between-group varGroup synchrony.
- `SE_betw_Grp` standard error of each observation.

**Author(s)**

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

**References**


**See Also**

dendro.varcov for models details.

**Examples**

```r
# Calculate synchrony for null.model (broad evaluation, mBE) and homoscedastic variant
# of unstructured model (or full, mUN) for conifersIP data,
# and heteroscedastic variant for 1970-1999 period.
data(conifersIP)

# Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using taxonomic grouping criteria (i.e. Species)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
                                      data = conifersIP, homoscedastic = TRUE)
summary(ModHm)# Class and length of list elements

# Synchrony for mBE and mUN models
sync(ModHm, modname = "mBE")
sync(ModHm, modname = "mUN")

# Chop the data from 1970 to 1999.
conif.30 <- conifersIP[conifersIP$Year>1969 & conifersIP$Year<2000,]
summary(conif.30$Year)

# Fit the heteroscedastic set of variance covariance mixed models (mBE, mHeNE, mHeCS, mHeUN)
# using taxonomic grouping criteria (i.e. Species)
ModHt30 <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Species",
```
sync.plot

Description

The function creates dot plots of within- and between-group synchrony as produced by `sync` from a selected model produced by `dendro.varcov`. Note that broad evaluation model (mBE) can not be plotted since it produces only one value per model.

Usage

```r
sync.plot(syncList)
```

Arguments

- `syncList`: a list of the type as produced by `sync`.

Details

The function makes a dot plots for within- and between-group synchrony for a user defined `varGroup` and `varTime` period in `dendro.varcov`.

Value

Dotplot

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

Examples

```r
## Plot homoscedastic narrow evaluation (mNE) and unstructured model (mUN)
# synchronies for conifersIP data:
data(conifersIP)

##Fit the homoscedastic set of varcov models (mBE, mNE, mCS, mUN)
# using geographic grouping criteria (ie. Region)
ModHm <- dendro.varcov(TRW ~ Code, varTime = "Year", varGroup = "Region",
data = conifersIP, homoscedastic = TRUE)

sync.plot(sync(ModHm, modname = "mNE"))
sync.plot(sync(ModHm, modname = "mUN"))
```
**Description**

The function calculates temporal trends of spatial synchrony from a `data.frame` with tree-ring width chronologies using a moving window as described in Shestakova et al. (2016). This method splits the time variable (`varTime`) in 30 years windows plus a 5 years lag, and in each window the within- or between-group level (`varGroup`) synchronies are calculated. The function can also be used to find synchrony with similar time series `data.frame` from other fields.

**Usage**

```r
sync.trend(formula, varTime='', varGroup='', data, window = 30, lag = 5,
            null.mod = TRUE, selection.method = c("AIC", "AICc", "BIC"),
            all.mod = FALSE, homoscedastic = TRUE, between.group = FALSE)
```

**Arguments**

- `formula`: a formula a typical model formula such as `Y ~ A`, where `Y` is usually tree-ring width and `A` may be a grouping factor such as the specific names of tree-ring width chronologies (`conifersIP`).
- `varTime`: a character specifying the time variable to consider in calculating synchrony estimates. Models with less than 10 different time-points may produce unreliable results.
- `varGroup`: a character grouping variable. In dendrochronological studies different grouping strategies can be used. We used here two strategies following taxonomic (i.e. species) or geographic (i.e. region) criteria.
- `data`: a `data.frame` with tree-ring chronologies, years and grouping variables as columns.
- `window`: an integer specifying the window size (i.e. number of years) to be used to calculate synchrony. Must be greater than 20 (>=20). Defaults to 20.
- `lag`: an integer specifying the lag that the window is moving (i.e. number of vrTirs moving window) to be used to calculate synchrony. Must be greater than 1 (>=1). Defaults to 5.
- `null.mod`: a logical specifying if only the null model for general synchrony is fitted (broad evaluation, mBE). Default TRUE.
- `selection.method`: a character string of "AIC", "AICc" or "BIC", specifying the information criterion used for model selection.
- `all.mod`: a logical specifying if all homoscedastic and heteroscedastic models should be fitted. Default FALSE.
- `homoscedastic`: a logical specifying if models should be an optional varFunc object or one-sided formula describing the within-group heteroscedasticity structure. Default TRUE.
- `between.group`: a logical specifying if between-group synchrony is displayed instead of within-group synchrony. Default FALSE.
Details

The function fits by default ("null.mod=T") the null model for general synchrony (broad evaluation, mBE) for a specified time window size and lag. If "null.mod=F" the function calculates homoscedastic or heteroscedastic versions of variance-covariance (VCOV) mixed models available (mBE, mNE, mCS, mUN, mHeNE, mHeCS, mHeUN; dendro.varcov) for each time window size and lag selected. In each window the best model is chosen based on the minimum information criterion selected between "AIC", "AICc" or "BIC". When no selection.method is defined by default AIC is used. If "all.mod=T" the functions fits the homoscedastic and heteroscedastic versions of the 7 models (this is a highly time consuming process).

Value

The function returns a data.frame containing the following components:

- for null.mod TRUE:

  a_Group a column representing the within-group synchrony (mBE).
  SE standard error of each observation.
  Windlag a column representing the lag of the window used to split the time variable. A 0 value means that lag is 0, and then the defined time window starts from minimum varTime value.
  varTime a column representing the varTime variable.

- for null.mod FALSE:

  Modname a column indicating the best model fit and the information criterion used.
  GroupName a column indicating levels of the varGroup for each time-window selected.
  a_Group a column indicating within-group synchrony for each varGroup level at time-window selected.
  a_betw_Grp a column indicating between-group synchrony for each varGroup level at time-window selected. Only if between.group is set to TRUE.
  SE standard error of each observation.
  Windlag a column representing the lag of the window used to split the time variable. A 0 value means that lag is 0, and then the defined time window starts from minimum varTime value.
  varTime a column representing the varTime variable window mean point.

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas
References


Examples

```r
## Calculate temporal trends of spatial synchrony for conifersIP data:
data(conifersIP)

## Fit the null model temporal trend (mBE)
# using taxonomic grouping criteria (i.e. Species)
mBE.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, null.mod = TRUE, window = 30, lag = 5)
mBE.trend# it returns a data.frame

## Not run:
## Fit homoscedastic within-group trends (mBE, mNE, mCS, mUN)
# using geographic grouping criteria (i.e. Region)
geo.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Region",
data = conifersIP, window = 30, lag = 5, null.mod = FALSE, homoscedastic = TRUE)
geo.trend# a data.frame with varGroup synchrony for each time window.

## Fit heteroscedastic between-group trends (mBE, mNE, mCS, mUN)
# using geographic grouping criteria (i.e. Region) and BIC
geo.het.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Region",
data = conifersIP, window = 30, lag = 5, null.mod = FALSE,
selection.method = c("BIC"), homoscedastic = FALSE,
between.group = TRUE)
geo.het.trend

## Fit homoscedastic and heteroscedastic within-group trends
# using taxonomic grouping criteria (i.e. Species) and BIC
geo.tot.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, window = 30, lag = 5,
selection.method = c("BIC"), all.mod = TRUE)
geo.tot.trend

## End(Not run)
```
Description

The function creates a line chart showing temporal trends of spatial synchrony from data.frame of the type as produced by `sync.trend`.

Usage

```r
sync.trend.plot (sync.trend.data)
```

Arguments

- `sync.trend.data`: a data.frame of the type as produced by `sync.trend`.

Details

The function makes a line chart showing synchrony trends across years from a data.frame produced by `sync.trend`. Within- or between-group synchrony and SE are indicated for a selected time window. If synchrony is defined using `null.mod = TRUE (sync.trend)` only general synchrony is plotted. If synchrony is defined using `null.mod = FALSE (sync.trend)` different synchronies for each group variable (`varGroup`) are fitted with different colours for each stratum.

Value

Line chart

Author(s)

Josu G. Alday, Tatiana A. Shestakova, Victor Resco de Dios, Jordi Voltas

Examples

```r
## Calculate temporal trends of synchrony for conifersIP data:
data(conifersIP)

## Fit the null.model temporal trend (mBE) using taxonomic grouping criteria (i.e. Species)
mBE.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Species",
                       data = conifersIP, null.mod = TRUE, window = 30, lag = 5)
mBE.trend# it returns a data.frame
sync.trend.plot(mBE.trend)# Broad evaluation synchrony linechart

## Not run:
## Fit homoscedastic within-group trends (mBE, mNE, mCS, mUN)
# using geographic grouping criteria (i.e. Region)
geo.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Region",
                       data = conifersIP, null.mod = FALSE, window = 30, lag = 5)
geo.trend# it returns a data.frame
sync.trend.plot(geo.trend)# Evaluation synchrony linechart
```
data = conifersIP, window = 30, lag = 5,
null.mod = FALSE, homoscedastic = TRUE)

geo.trend# a data.frame with varGroup synchrony for each time window.
sync.trend.plot(geo.trend)# Selected heteroscedastic between-group trends by AIC

## Fit heteroscedastic between-group trends (mBE, mHeNE, mHeCS, mHeUN)
# using geographic grouping criteria (i.e. Region) and AICc
geo.het.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Region",
data = conifersIP, window = 30, lag = 5, null.mod = FALSE,
selection.method = c("AICc"), homoscedastic = FALSE, between.group = TRUE)

geo.het.trend
sync.trend.plot(geo.het.trend)# Selected heteroscedastic between-group trends by AICc

## Fit homoscedastic and heteroscedastic within-group trends
# using taxonomic grouping criteria (i.e. Species) and BIC
geo.tot.trend <- sync.trend(TRW ~ Code, varTime = "Year", varGroup = "Species",
data = conifersIP, window = 30, lag = 5, selection.method = c("BIC"),
null.mod = F, all.mod = TRUE)

geo.tot.trend
# Selected homoscedastic and heteroscedastic within-group trends by BIC
sync.trend.plot(geo.tot.trend)

## End(Not run)
## Index

*Topic **datasets**

  conifersIP, \textit{5}

AIC, \textit{19}

bet.aSE, \textit{2}
bet.het.aSE, \textit{3}

BIC, \textit{19}

complete.cases, \textit{11, 13}
conifersIP, \textit{5, 11, 23}
csbet.aSE, \textit{6}
csbet.het.aSE, \textit{7}
cswi.aSE, \textit{8}
cswi.het.aSE, \textit{10}

dendro.varcov, \textit{2, 3, 6–8, 10, 11, 14, 15, 17–22, 24}

gen.aSE, \textit{14}
gen.het.aSE, \textit{15}

het.var, \textit{17}

lmeObject, \textit{12, 13}
logLik, \textit{19}

mod.table, \textit{18}

na.action, \textit{11, 13}

sync, \textit{2–4, 6–10, 14–17, 19, 22}
sync.plot, \textit{22}
sync.trend, \textit{23, 26}
sync.trend.plot, \textit{20, 26}

varFunc, \textit{12}

varIdent, \textit{12}