Package ‘asremlPlus’

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**Title** Augments ‘ASReml-R’ in Fitting Mixed Models and Packages

Generally in Exploring Prediction Differences

**Depends** R (>= 3.5.0)

**Imports** dae, doParallel, dplyr, ggplot2, foreach, grDevices, graphics, methods, parallel, qplot, reshape2, RColorBrewer, rlang, stats, stringr, sticky, utils

**Suggests** testthat, lattice, emmeans, lmerTest, pbkrtest, R.rsp

**Enhances** asreml

**VignetteBuilder** R.rsp

**SystemRequirements** asreml-R 4.x

**LazyData** true

**Description** Assists in automating the selection of terms to include in mixed models when 'asreml' is used to fit the models. Also used to display, in tables and graphs, predictions obtained using any model fitting function and to explore differences between predictions. The content falls into the following natural groupings: (i) Data, (ii) Object manipulation functions, (iii) Model modification functions, (iv) Model testing functions, (v) Model diagnostics functions, (vi) Prediction production and presentation functions, (vii) Response transformation functions, and (viii) Miscellaneous functions (for further details see ‘asremlPlus-package’ in help). A history of the fitting of a sequence of models is kept in a data frame. Procedures are available for choosing models that conform to the hierarchy or marginality principle and for displaying predictions for significant terms in tables and graphs. The 'asreml' package provides a computationally efficient algorithm for fitting mixed models using Residual Maximum Likelihood. It is a commercial package that can be purchased from 'VSNi' [https://vsni.co.uk/] as 'asreml-R', who will supply a zip file for local installation/updating (see [https://asreml.kb.vsni.co.uk/]). It is not needed for functions that are methods for ‘alldiffs’ and ‘data.frame’ objects. The package ‘asremlPlus’ can also be installed from [http://chris.brien.name/rpackages/].

**License** MIT + file LICENSE

**URL** http://chris.brien.name

**NeedsCompilation** no

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Description

Assists in automating the selection of terms to include in mixed models when 'asreml' is used to fit the models. Also used to display, in tables and graphs, predictions obtained using any model fitting function and to explore differences between predictions. The content falls into the following natural groupings: (i) Data, (ii) Object manipulation functions, (iii) Model modification functions, (iv) Model testing functions, (v) Model diagnostics functions, (vi) Prediction production and presentation functions, (vii) Response transformation functions, and (viii) Miscellaneous functions (for further details see 'asremlPlus-package' in help). A history of the fitting of a sequence of models is kept in a data frame. Procedures are available for choosing models that conform to the hierarchy or marginality principle and for displaying predictions for significant terms in tables and graphs. The 'asreml' package provides a computationally efficient algorithm for fitting mixed models using Residual Maximum Likelihood. It is a commercial package that can be purchased from 'VSNi' <https://vsni.co.uk/> as 'asreml-R', who will supply a zip file for local installation/updating (see <https://asreml.kb.vsni.co.uk/>). It is not needed for functions that are methods for 'alldiffs' and 'data.frame' objects. The package 'asremPlus' can also be installed from <http://chris.brien.name/rpackages/>.

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(i) Data

Oats.dat  Data for an experiment to investigate nitrogen response of 3 oats varieties.
Wheat.dat  Data for an experiment to investigate 25 varieties of wheat.
WaterRunoff.dat  Data for an experiment to investigate the quality of water runoff over time

(ii) Object manipulation

as.alldiffs  Forms an alldiffs.object from the supplied predictions, along with those statistics, associated with the predictions and their pairwise differences, that have been supplied.
asrtests  Pseudonym for as.asrtests.
as.asrtests  Forms an asrtests.object that stores (i) a fitted asreml object, (ii) a pseudo-anova table for the fixed terms and (iii) a history of changes and hypothesis testing used in obtaining the model.
as.predictions.frame  Forms a predictions.frame from a data.frame, ensuring that the correct columns are present.
facCombine.alldiffs  Combines several factors into one in the components of an alldiffs.object.
facRecast.alldiffs
Reorders and/or revises the factor levels using the order of old levels in levels.order and the new labels for the levels given in newlabels.

facRename.alldiffs
Renames factors in the prediction component of an alldiffs.object.

getFormulae.asreml
 Gets the formulae from an asreml object.

is.alldiffs
A single-line function that tests whether an object is of class alldiffs.

is.asrtests
A single-line function that tests whether an object is of class asrtests.

is.predictions.frame
A single-line function that tests whether an object is of classes predictions.frame and data.frame.

makeTPPSplineMats.data.frame
Make the spline basis matrices and data needed to fit Tensor Product P-Splines.

print.alldiffs
Prints the values in an alldiffs.object in a nice format.

print.asrtests
Prints the values in an asrtests.object.

print.LSDdata
Prints the components of a list containing data on the LSDs for all pairwise differences of predictions.

print.predictions.frame
Prints the values in a predictions.frame, with or without title and heading.

print.test.summary
Prints a data.frame containing a test.summary.

print.wald.tab
Prints a data.frame containing a Wald or pseudoanova table.

sort.alldiffs
Sorts the components of an alldiffs.object according to the predicted values associated with a factor.

subset.alldiffs
Subsets the components in an alldiffs.object according to the supplied condition.

subset.list
Forms a list that contains a subset of the components of the supplied list.

validAlldiffs
Checks that an object is a valid alldiffs.object.

validAsrtests
Checks that an object is a valid asrtests.object.

validPredictionsFrame
Checks that an object is a valid predictions.frame.

(iii) Model modification

addSpatialModel.asrtests
Adds, to a supplied model, a spatial model that accounts for local spatial variation.

addSpatialModelOnIC.asrtests
Uses information criteria to decide whether to add a spatial model to account for local spatial variation.

changeTerms.asrtests
Adds and drops terms from one or both of the fixed or random model, replaces the residual (rcov) model with a new model and changes bounds or initial values of terms.

iterate.asrtests
Subject the fitted asreml.obj stored in an asrtests.object to further iterations of the fitting process.

newfit.asreml
Refits an asreml model with modified model formula using either a call to 'update.asreml' or a direct call to 'asreml'.

reparamSigDevn.asrtests
Reparamterizes each random (deviations) term involving 'devn.fac' to a fixed term and ensures that the same term, with 'trend.num' replacing 'devn.fac', is included if any other term with 'trend.num' is included in 'terms'.

rmboundary.asrtests
Removes any boundary or singular variance components from the fit stored in 'asreml.obj' and records their
setvarianceterms.call

Allows the setting of bounds and initial values for terms in the 'random' and 'residual' arguments of an 'asreml' call.

(iv) Model selection

addto.test.summary

 Adds a row to a test.summary data.frame.

changeModelOnIC.asrtests

Uses information criteria to decide whether to change an already fitted model.

chooseModel.asrtests

Determines and records the set of significant terms using an asrtests.object, taking into account the hierarchy or marginality relations of the terms.

chooseModel.data.frame

Determines the set of significant terms from results stored in a data frame, taking into account the marginality relations of terms and recording the tests used in a data frame.

chooseSpatialModelOnIC.asrtests

Uses information criteria to choose the best fitting spatial model for accounting for local spatial variation.

getTestPvalue.asrtests

Gets the p-value for a test recorded in the test.summary data.frame of an asrtests.object.

infoCriteria.asreml

Computes AIC and BIC for models.

infoCriteria.list

Computes AIC and BIC for models.

recalcWaldTab.asrtests

Recalculates the denDF, F.inc and P values for a table of Wald test statistics obtained using 'wald.asreml'.

REMLRT.asreml

Performs a REML ratio test.

bootREMLRT.asreml

Performs a REML ratio test using the parametric bootstrap.

testranfix.asrtests

Tests for a single fixed or random term in model fitted using 'asreml' and records the result in an asrtests.object.

testresidual.asrtests

Fits a new residual formula using 'asreml', tests whether the change is significant and records the result in an asrtests.object.

testswapran.asrtests

Tests, using a REMLRT, the significance of the difference between the current random model and one in which old terms are dropped and new terms are added. The result is recorded in an asrtests.object.

(v) Model diagnostics and simulation

plotVariofaces

Plots empirical variogram faces, including envelopes, from supplied residuals as described by Stefanova, Smith & Cullis (2009).

variofaces.asreml

Calculates and plots empirical variogram faces, including envelopes, as described by Stefanova, Smith & Cullis (2009).

estimateV.asreml

Forms the estimated variance, random or residual matrix for the observations from the variance parameter estimates.

simulate.asreml

Produce sets of simulated data from a multivariate normal distribution and save quantities related to the simulated data.

(vi) Prediction production and presentation
addBacktransforms.alldiffs
• Adds or recalculates the backtransforms component of an alldiffs.object.

allDifferences.data.frame
• Using supplied predictions and standard errors of pairwise differences or the variance matrix of predictions, forms all pairwise differences between the set of predictions, and p-values for the differences.

exploreLSDs
• Explores the computed LSD values for pairwise differences between predictions.

linTransform.alldiffs
• Calculates a linear transformation of the predictions stored in an alldiffs.object.

pairdiffsTransform.alldiffs
• Calculates the differences between nominated pairs of predictions stored in an alldiffs.object.

pickLSDstatistics
• Pick LSDstatistics whose values minimize the number of errors in pairwise comparisons of predictions.

plotLSDerrors.data.frame
• Plots a map of the supplied errors that occur in using the computed LSD values for pairwise differences between predictions.

plotLSDerrors.alldiffs
• Plots a map of the errors that occur in using the computed LSD values for pairwise differences between predictions.

plotLSDs.data.frame
• Plots a heat map of computed LSD values for pairwise differences between predictions.

plotLSDs.alldiffs
• Plots a heat map of computed LSD values for pairwise differences between predictions.

plotPredictions.data.frame
• Plots the predictions for a term, possibly with error bars.

plotPvalues.alldiffs
• Plots the p-values in the p.differences components of an alldiffs.object as a heat map.

plotPvalues.data.frame
• Plots the p-values in data.frame as a heat map.

predictPlus.asreml
• Forms the predictions and associated statistics for a term, using an asreml object and a wald.tab and taking into account that a numeric vector and a factor having parallel values may occur in the model. It stores the results in an object of class ‘alldiffs’ and may print the results. It can be when there are not parallel values.

predictPresent.asreml
• Forms the predictions for each of one or more terms and presents them in tables and/or graphs.

ratioTransform.alldiffs
• Calculates the ratios of nominated pairs of predictions stored in an alldiffs.object.

recalcLSD.alldiffs
• Adds or recalculates the LSD.frame that is a component of an alldiffs.object.

redoErrorIntervals.alldiffs
• Adds or replaces the error intervals stored in the prediction component of an alldiffs.object according to a new classify.

renewClassify.alldiffs
• Renews the components in an alldiffs.object according to a new classify.

sort.alldiffs
• Sorts the components in an alldiffs.object according to the predicted values associated with a factor.

subset.alldiffs
• Subsets the components in an alldiffs.object according to the supplied condition.

sort.predictions.frame
• Sorts a predictions.frame according to the predicted values. associated with a factor.
(vii) Response transformation

**angular**
Applies the angular transformation to proportions.

**angular.mod**
Applies the modified angular transformation to a vector of counts.

**powerTransform**
Performs a combination of a linear and a power transformation on a variable. The transformed variable is stored in the `data.frame` data.

(viii) Miscellaneous

**getASRemlVersionLoaded**
Finds the version of asreml that is loaded and returns the initial characters in version.

**loadASRemlVersion**
Ensures that a specific version of asreml is loaded.

**num.recode**
Recodes the unique values of a vector using the values in a new vector.

**permute.square**
Permutations the rows and columns of a square matrix.

**permute.to.zero.lowertri**
Permutates a square matrix until all the lower triangular elements are zero.

The functions whose names end in `alldiffs` utilize an `alldiffs.object` that stores: (i) a `predictions.frame`, being a data frame containing predicted values, variables indexing them and their standard errors and estimability status; the lower and upper limits of error intervals will be included when these are requested, (ii) optionally, square matrices containing all pairwise differences, the standard errors and p-values of the differences, and a `data.frame` containing LSD values and their summary statistics, (iii) optionally, the variance matrix of the predictions, and (iv) if the response was transformed for analysis, a data frame with backtransforms of the predicted values.

The functions whose names end in `asrtests`, which are most of the model functions, utilize an `asrtests.object` that stores: (i) the currently fitted model in `asreml.obj`, (ii) the table of test statistics for the fixed effects in `wald.tab`, and (iii) a data frame that contains a history of the changes made to the model in `test.summary`.

**Author(s)**

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**References**


**See Also**

asreml
Examples

## Not run:
## Analyse wheat dat using asreml and asremlPlus (see also the Wheat Vignette)
## Set up for analysis
library(dae)
library(asreml)
library(asremlPlus)
## use ?Wheat.dat for data set details
data(Wheat.dat)

# Fit initial model
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
  random = ~ Row + Column + units,
  residual = ~ ar1(Row):ar1(Column),
  data=Wheat.dat)
summary(current.asr)

# Intialize a testing sequence by loading the current fit into an asrtests object
current.asrt <- as.asrtests(current.asr, NULL, NULL)

# Check for and remove any boundary terms
current.asrt <- rmboundary(current.asrt)

# Unbind Rep, Row and Column components and reload into an asrtests object
current.asr <- setvarianceterms(current.asr$call,
  terms = c("Rep", "Rep:Row", "Rep:Column"),
  bounds = "U")
current.asr <- as.asrtests(current.asr, NULL, NULL)
current.asr <- rmboundary(current.asr)
summary(current.asr$asreml.obj)$varcomp
print(current.asr, which = "testsummary")
print(current.asr, which = "pseudoanova")

# Check term for within Column pairs (a post hoc covariate)
current.asrt <- testranfix(current.asr, "WithinColPairs", drop.fix.ns=TRUE)

# Test nugget term
current.asrt <- testranfix(current.asr, "units", positive=TRUE)

# Test Row autocorrelation
current.asrt <- testresidual(current.asr, "~ Row:ar1(Column)",
  label="Row autocorrelation", simpler=TRUE)

# Test Col autocorrelation (depends on whether Row autocorrelation retained)
(p <- getTestPvalue(current.asr, label = "Row autocorrelation retained"))
if (p <= 0.05)
  current.asr <- testresidual(current.asr, "~ ar1(Row):Column",
    label="Col autocorrelation",
    simpler=TRUE, update=FALSE)
else
  current.asrt <- testresidual(current.asr, "~ Row:Column",
    label="Col autocorrelation",
    simpler=TRUE, update=FALSE)

# Output the results
addBacktransforms.alldiffs

Adds or recalculates the backtransforms component of an alldiffs.object.

Description

Given an alldiffs.object, adds or recalculate its backtransforms component. The values of transform.power, offset, scale and transform.function from the backtransforms component will be used, unless this component is NULL when the values supplied in the call will be used.

Usage

## S3 method for class 'alldiffs'
addBacktransforms(alldiffs.obj,
transform.power = 1, offset = 0, scale = 1,
transform.function = "identity", ...)
Arguments

**alldiffs.obj**  
An *alldiffs.object*.

**transform.power**  
A **numeric** specifying the power of a transformation, if one has been applied to the response variable. Unless it is equal to 1, the default, back-transforms of the predictions will be obtained and presented in tables or graphs as appropriate. The back-transformation raises the predictions to the power equal to the reciprocal of *transform.power*, unless it equals 0 in which case the exponential of the predictions is taken.

**offset**  
A **numeric** that has been added to each value of the response after any scaling and before applying any power transformation.

**scale**  
A **numeric** by which each value of the response has been multiplied before adding any offset and applying any power transformation.

**transform.function**  
A **character** giving the name of a function that specifies the scale on which the predicted values are defined. This may be the result of a transformation of the data using the function or the use of the function as a link function in the fitting of a generalized linear (mixed) model (GL(M)M). The possible transform.functions are `identity`, `log`, `inverse`, `sqrt`, `logit`, `probit`, and `cloglog`. The predicted.values and error.intervals, if not `StandardError` intervals, will be back-transformed using the inverse function of the transform.function. The standard.error column will be set to NA, unless (i) `asreml` returns columns named `transformed.value` and `approx.se`, as well as those called `predicted.values` and `standard.error` (such as when a GLM is fitted) and (ii) the values in `transformed.value` are equal to those obtained by backtransforming the predicted.values using the inverse function of the transform.function. Then, the `approx.se` values will be saved in the `standard.error` column of the backtransforms component of the returned alldiffs.obj. Also, the `transformed.value` and `approx.se` columns are removed from both the predictions and backtransforms components of the alldiffs.obj. Note that the values that end up in the `standard.errors` column are approximate for the backtransformed values and are not used in calculating error.intervals.

Value

An *alldiffs.object* with components predictions, vcov, differences, p.differences, sed, LSD and backtransforms.

The backtransforms component will have the attributes (i) LSDtpe, LSDby and LSDstatistic added from the predictions component and (ii) transform.power, offset, scale, and link.

Author(s)

Chris Brien

See Also

`asremlPlus-package`, `as.alldiffs`, `sort.alldiffs`, `subset.alldiffs`, `print.alldiffs`, `renewClassify.alldiffs`, `redoErrorIntervals.alldiffs`, `plotPredictions.data.frame`, `predictPlus.asreml`, `predictPresent.asreml`
Examples

```r
##Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18" & Benches != "3")

##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = log.Turbidity ~ Benches + (Sources * (Type + Species)),
                    random = ~ Benches:MainPlots,
                    keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus(classify = "Sources:Type",
                        asreml.obj = current.asr,
                        wald.tab = current.asrt$wald.tab,
                        present = c("Sources", "Type", "Species"))

## End(Not run)

##Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &&
    requireNamespace("emmeans", quietly = TRUE))
{
m1.lmer <- lmerTest::lmer(log.Turbidity ~ Benches + (Sources * (Type + Species)) +
                         (1|Benches:MainPlots),
                         data=tmp)
TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Species)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
                                  se = "SE", interval.type = "CI",
                                  interval.names = c("lower.CL", "upper.CL"))

## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els,els]
TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Species",
                           vcov = TS.vcov, tdf = den.df)
validAlldiffs(TS.diffs)
}

## Recalculate the back-transforms of the predictions obtained using asreml or lmerTest
if (exists("TS.diffs"))
{
  TS.diffs <- addBacktransforms.alldiffs(TS.diffs, transform.power = 0)
}
```

```
addSpatialModel.asrtests

Adds, to a supplied model, a spatial model that accounts for local spatial variation.
```
addSpatialModel.asrtests

Description

Adds either a correlation, two-dimensional tensor-product natural cubic smoothing spline (TP-NCSS), or a two-dimensional tensor-product penalized P-spline model (TPPS) to account for the local spatial variation exhibited by a response variable measured on a potentially irregular grid of rows and columns of the units. The data may be arranged in sections, for each of which there is a grid and for which the model is to be fitted separately. Also, the rows and columns of a grid are not necessarily one observational unit wide.

No hypothesis testing or comparison of information criteria is made. To only change the terms based on a comparison of information criteria use chooseSpatialModelOnIC.asrtests.

The model fit supplied in the asrtests.obj should not include terms that will be included in the local spatial model. All spatial model terms are fitted as fixed or random. Consequently, the residual model does not have to be iid.

One or more rows is added for each section to the test.summary data.frame. Convergence and the occurrence of fixed correlations in fitting the model is checked and a note included in the action if there was not. All components of the asrtests.object are updated for the new model.

Usage

```r
## S3 method for class 'asrtests'
addSpatialModel(asrtests.obj, spatial.model = "TPPS",
sections = NULL,
row.covar = "cRow", col.covar = "cCol",
row.factor = "Row", col.factor = "Col",
corr.funcs = c("ar1", "ar1"),
row.corrFitfirst = TRUE,
dropRowterm = NULL, dropColterm = NULL,
nsegs = NULL, nestorder = c(1,1),
degree = c(3,3), difforder = c(2,2),
asreml.option = "mbf", tpps4mbf.obj = NULL,
allow.unconverged = FALSE, allow.fixedcorrelation = FALSE,
checkboundaryonly = FALSE, update = FALSE,
IClikelihood = "full", ...)
```

Arguments

- **asrtests.obj**: An `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary`.
- **spatial.model**: A single character string nominating the type of spatial model to fit. Possible values are `corr`, `TPNCSS` and `TPPS`.
- **sections**: A single character string that species the name of the column in the data.frame that contains the factor that identifies different sections of the data to which separate spatial models are to be fitted.
- **row.covar**: A single character string nominating a numeric that contains the values of a centred covariate indexing the rows of a grid. The numeric must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.
- **col.covar**: A single character string nominating a numeric that contains the values of a centred covariate indexing the columns of a grid. The numeric must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.
row.factor
A single character string nominating a factor that indexes the rows of a grid that are to be one dimension of a spatial correlation model. The factor must a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

col.factor
A single character string nominating a factor that indexes the columns of a grid that are to be one dimension of a spatial correlation model. The factor must a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

corr.funcs
A single character string of length two that specifies the asreml one-dimensional correlation or variance model function for the row and column dimensions of a two-dimensional separable spatial correlation model; the two-dimensional model is fitted as a random term. If a correlation or variance model is not to be investigated for one of the dimensions, specify "" for that dimension.

row.corrFitfirst
If TRUE then the row correlation or variance function is fitted first, followed by the addition of the column correlation or variance function. If FALSE, the order of fitting is reversed.

dropRowterm
A single character string nominating a factor in the data.frame that has as many levels as there are unique values in row.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropRowterm and a random row deviations term based on row.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

dropColterm
A single character string nominating a factor in the data.frame that has as many levels as there are unique values in col.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropColterm and a random column deviations term based on col.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

nsegs
A pair of numeric values giving the number of segments into which the column and row ranges are to be split, respectively (each value specifies the number of internal knots + 1). If only one number is specified, that value is used in both dimensions. If not specified, then (number of unique values - 1) is used in each dimension; for a grid layout with equal spacing, this gives a knot at each data value.

nestorder
A character of length 2. The order of nesting for column and row dimensions, respectively; default=1 (no nesting). A value of 2 generates a spline with half the number of segments in that dimension, etc. The number of segments in each direction must be a multiple of the order of nesting.

degree
A character of length 2. The degree of polynomial spline to be used for column and row dimensions respectively; default=3.

difforder
A character of length 2. The order of differencing for column and row dimensions, respectively; default=2.

asreml.option
A single character string specifying whether the grp or mbf methods are to be used to supply externally formed covariate matrices to asreml. If the mbf methods is to be used, then makeTPPSplineMats.data.frame must be used before calling addSpatialModel.asrtests. Compared to the mbf method, the grp method creates large asrtests.objects, but is faster. The grp method adds columns to the data.frame containing the data; the mbf method adds only fixed covariate to data and stores the random covariates externally.
addSpatialModel.asrtests

tppsmbf.obj An object made with `makeTPPSplineMats.data.frame` and which contains the spline basis information, extra to that created by `makeTPPSplineMats.data.frame`, that is needed to fit a TPPS model using the mbf method of asreml.

allow.unconverged A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit of the new model does not converge, the supplied asrtests.obj is returned. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

allow.fixedcorrelation A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly If TRUE then boundary and singular terms are not removed by `rmboundary.asrtests`; a warning is issued instead. Note that, for correlation models, the fitting of each dimension and the test for a nugget term are performed with checkboundaryonly set to TRUE and its supplied setting only honoured using a call to `rmboundary.asrtests` immediately prior to returning the final result of the fitting.

update If TRUE then `update.asreml` is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified as specified and (ii) modifications specified via ... are made.

IClikelihood A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new asrtests.object. If none, none are included. Otherwise, if REML, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full, then the AIC and BIC based on the full likelihood are included. (See also `infoCriteria.asreml`.)

... Further arguments passed to `changeModelOnIC.asrtests`, asreml and tpsmmb.

Details

The model to which the spatial models is to be added is supplied in the asrtests.obj. It should not include terms that will be included in the local spatial model. All spatial model terms are fitted as fixed or random. Consequently, the residual model does not have to be iid. The improvement in the fit resulting from the addition of a spatial model to the supplied model is evaluated. Note that the data must be in the order that corresponds to the residual argument with a variable to the right of another variable changes levels in the data frame faster than those of the other variable e.g. Row:Column implies that all levels for Column in consecutive rows of the data.frame with a single Row level.

For the corr spatial model, the default model is an autocorrelation model of order one (ar1) for each dimension. However, any of the single dimension correlation/variance models from asreml can be specified for each dimension, as can no correlation model for a dimension; the models for the two dimensions can differ. A series of models are tried, without removing boundary or singular terms, beginning with the addition of row correlation and followed by the addition of column correlation or, if the row.corrFitfirst is set to FALSE, the reverse order. If the fitting of the first-fitted
correlation did not result in a model change because the fitting did not converge or correlations were fixed, but the fit of the second correlation was successful, then adding the first correlation will be retried. If one of the metric correlation functions is specified (e.g. exp), then the row.covar or col.covar will be used in the spatial model. However, because the correlations are fitted separately for the two dimensions, the row.factor and col.factor are needed for all models and is used for a dimension that does not involve a correlation/variance function for the fit being performed. Also, the correlation models are fitted as random terms and so the correlation model will include a variance parameter for the grid even when ar1 is used to specify the correlation model, i.e. the model fitted is a variance model and there is no difference between ar1 and ar1v in fitting the model. The variance parameter for this term represents the spatial variance and the fit necessarily includes a nugget term, this being the residual variance. The nugget term can be removed by fixing the residual variance to one using setvarianceterms.call, provided heterogeneous residual variances have not been specified. Once the fitting of the correlation model has been completed, the rmboundary function will be executed with the checkboundaryonly value supplied in the addSpatialModel.asrtests call.

The TPNCSS spatial model is as described by Verbyla et al. (2018) and the TPPS model is as described by Rodriguez-Alvarez et al. (2018). However, for the TPPS model, the degree of the polynomial and the order of differencing can be varied. The defaults of 3 and 2, respectively, fit a cubic spline with second order differencing, while setting both the degree and order of differencing to 1 will fit a type of linear variance model (Piepho, Boer and Williams, 2022) The fixed terms for these models are row.covar + col.covar + row.covar:col.covar and the random terms are spl(row.covar) + spl(col.covar) + dev(row.covar) + dev(col.covar) + spl(row.covar):col.covar + row.covar:spl(col.covar) + spl(row.covar):spl(col.covar). The supplied model should not include any of these terms. However, any fixed or random main-effect term for either dropRowterm or dropColterm will be removed from the fit.

The TPPS model is fitted using the function tpsmb from the R package TPSbits authored by Sue Welham (2022). There are two methods for supplying the spline basis information produced by tpsmb to asreml. The grp method adds the it to the data.frame holding the information for the analysis. The mbf method requires the spline basis information to be in the same environment as the function that is called to make a fit using asreml. To this end and prior to invoking the calling function, makeTPPSplineMats.data.frame must be used produce the data.frames.

All models utilize the function changeTerms.asrtests to fit the spatial model. Arguments from tpsmb and changeTerms.asrtests can be supplied in calls to addSpatialModel.asrtests and will be passed on to the relevant function through the ellipses argument (...).

The data for experiment can be divided sections and the same spatial model fitted separately to each. The fit over all of the sections is assessed.

Each combination of a row.coords and a col.coords does not have to specify a single observation; for example, to fit a local spatial model to the main units of a split-unit design, each combination would correspond to a main unit and all subunits of the main unit would have the same combination.

Value

An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary for the model that includes the spatial model, unless the spatial model fails to be fitted when allow.unconverged and/or allow.fixedcorrelation is set to FALSE.

Author(s)

Chris Brien
addSpatialModel.asrtests

References


See Also

`as.asrtests, rmboundary.asrtests, addSpatialModelOnIC.asrtests, testranfix.asrtests, testresidual.asrtests, newfit.asreml, reparamSigDevn.asrtests, changeTerms.asrtests, chooseSpatialModelOnIC.asrtests, changeTerms.asrtests, infoCriteria.asreml`

Examples

```r
## Not run:

data(Wheat.dat)

# Add row and column covariates
Wheat.dat <- within(Wheat.dat,
{
  cColumn <- dae::as.numfac(Column)
  cColumn <- cColumn - mean(unique(cColumn))
  cRow <- dae::as.numfac(Row)
  cRow <- cRow - mean(unique(cRow))
})

# Fit initial model
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                      random = ~ Row + Column,
                      data=Wheat.dat)

# Create an asrtests object, removing boundary terms
current.asrt <- as.asrtests(current.asr, NULL, NULL,
                            label = "Random Row and Column effects")

current.asrt <- rmboundary(current.asrt)

current.asrt <- addSpatialModel(current.asrt, spatial.model = "TPPS",
                                row.covar = "cRow", col.covar = "cColumn",
                                dropRowterm = "Row", dropColterm = "Column",
                                asreml.option = "grp")

infoCriteria(current.asrt$asreml.obj)

## End(Not run)```
addSpatialModelOnIC

Uses information criteria to decide whether to add a spatial model to account for local spatial variation.

Description

Adds either a correlation, two-dimensional tensor-product natural cubic smoothing spline (TP-NCSS), or a two-dimensional tensor-product penalized P-spline model (TPPS) to account for the local spatial variation exhibited by a response variable measured on a potentially irregular grid of rows and columns of the units. The data may be arranged in sections for each of which there is a grid and for which the model is to be fitted separately. Also, the rows and columns of a grid are not necessarily one observational unit wide. The spatial model is only added if the information criterion of the supplied model is decreased with the addition of the local spatial model.

A row is added for each section to the test.summary data.frame of the asrtests.object stating whether or not the new model has been swapped for a model in which the spatial model has been add to the supplied model. Convergence and the occurrence of fixed correlations in fitting the model is checked and a note included in the action if there was not. All components of the asrtests.object are updated to exhibit the differences between the supplied and the new model, if a spatial model is added.

Usage

```r
## S3 method for class 'asrtests'
addSpatialModelOnIC(asrtests.obj, spatial.model = "TPPS",
sections = NULL,
row.covar = "cRow", col.covar = "cCol",
row.factor = "Row", col.factor = "Col",
corr.funcs = c("ar1", "ar1"),
row.corrFitfirst = TRUE,
dropRowterm = NULL, dropColterm = NULL,
nsegs = NULL, nestorder = c(1,1),
degree = c(3,3), difforder = c(2,2),
asreml.option = "mbf", tpps4mbf.obj = NULL,
allow.unconverged = FALSE, allow.fixedcorrelation = FALSE,
checkboundaryonly = FALSE, update = FALSE,
IClikelihood = "full", which.IC = "AIC", ...)
```

Arguments

- **asrtests.obj** An *asrtests.object* containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.
- **spatial.model** A single character string nominating the type of spatial model to fit. Possible values are corr, TPNCSS and TPPS.
- **sections** A single character string that specifies the name of the column in the data.frame that contains the factor that identifies different sections of the data to which separate spatial models are to be fitted.
- **row.covar** A single character string nominating a numeric that contains the values of a centred covariate indexing the rows of a grid. The numeric must be a column in
the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

col.covar  A single character string nominating a numeric that contains the values of a centred covariate indexing the columns of a grid. The numeric must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

row.factor  A single character string nominating a factor that indexes the rows of a grid that are to be one dimension of a spatial correlation model. The factor must a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

col.factor  A single character string nominating a factor that indexes the columns of a grid that are to be one dimension of a spatial correlation model. The factor must a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

corr.funcs  A single character string of length two that specifies the asreml one-dimensional correlation or variance model function for the row and column dimensions of a two-dimensional separable spatial correlation model; the two-dimensional model is fitted as a random term. If a correlation or variance model is not to be investigated for one of the dimensions, specify "" for that dimension.

row.corrFitfirst  If TRUE then the row correlation or variance function is fitted first, followed by the addition of the column correlation or variance function. If FALSE, the order of fitting is reversed.

dropRowterm  A single character string nominating a factor in the data.frame that has as many levels as there are unique values in row.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropRowterm and a random row deviations term based on row.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

dropColterm  A single character string nominating a factor in the data.frame that has as many levels as there are unique values in col.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropColterm and a random column deviations term based on col.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

nsegs  A pair of numeric values giving the number of segments into which the column and row ranges are to be split, respectively (each value specifies the number of internal knots + 1). If only one number is specified, that value is used in both dimensions. If not specified, then (number of unique values - 1) is used in each dimension; for a grid layout with equal spacing, this gives a knot at each data value.

nestorder  A character of length 2. The order of nesting for column and row dimensions, respectively; default=1 (no nesting). A value of 2 generates a spline with half the number of segments in that dimension, etc. The number of segments in each direction must be a multiple of the order of nesting.

degree  A character of length 2. The degree of polynomial spline to be used for column and row dimensions respectively; default=3.

difforder  A character of length 2. The order of differencing for column and row dimensions, respectively; default=2.
addSpatialModelOnIC.asrtests

asreml.option  A single character string specifying whether the grp or mbf methods are to be used to supply externally formed covariate matrices to asreml. If the mbf methods is to be used, then makeTPPSplineMats.data.frame must be used before calling addSpatialModelOnIC.asrtests. Compared to the mbf method, the grp method creates large asreml objects, but is faster. The grp method adds columns to the data.frame containing the data; the mbf method adds only fixed covariate to data and stores the random covariates externally.

tpps4mbf.obj  An object made with makeTPPSplineMats.data.frame and which contains the spline basis information, extra to that created by makeTPPSplineMats.data.frame, that is needed to fit a TPPS model using the mbf method of asreml.

allow.unconverged  A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit of the new model does not converge, the supplied asrtests.obj is returned. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

allow.fixedcorrelation  A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly  If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead. Note that, for correlation models, the fitting of each dimension and the test for a nugget term are performed with checkboundaryonly set to TRUE and its supplied setting only honoured using a call to rmboundary.asrtests immediately prior to returning the final result of the fitting.

update  If TRUE then update.asreml is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified as specified and (ii) modifications specified via ... are made.

which.IC  A character specifying the information criterion to be used in selecting the best model. Possible values are AIC and BIC. The values of the criterion for supplied model must exceed that for changed model for the changed model to be returned.

IClikelihood  A character specifying whether Restricted Maximum Likelihood (REML) or the full likelihood (full) are to be used in calculating the information criteria.

...  Further arguments passed to changeModelOnIC.asrtests, asreml and tpsmmb.

Details

A fitted spatial model is only returned if it improves the fit over and above that of achieved with the model fit supplied in the asrtests.obj. To fit the spatial model without any hypotheses testing or comparison of information criteria use addSpatialModel.asrtests. The model fit supplied in the asrtests.obj should not include terms that will be included in the local spatial model. All spatial model terms are fitted as fixed or random. Consequently, the residual model does not have to be iid. Note that the data must be in the order that corresponds to the residual argument with a variable to
the right of another variable changes levels in the data frame faster than those of the other variable e.g. Row:Column implies that all levels for Column in consecutive rows of the data frame with a single Row level.

For the corr spatial model, the default model is an autocorrelation model of order one (ar1) for each dimension. However, any of the single dimension correlation/variance models from asreml can be specified for each dimension, as can no correlation model for a dimension; the models for the two dimensions can differ. Using a forward selection procedure, a series of models are tried, without removing boundary or singular terms, beginning with the addition of row correlation and followed by the addition of column correlation or, if the row.corrFitfirst is set to FALSE, the reverse order. If the fitting of the first-fitted correlation did not result in a model change because the fitting did not converge or correlations were fixed, but the fit of the second correlation was successful, then adding the first correlation will be retried. If one of the metric correlation functions is specified (e.g. exp), then the row.covar or col.covar will be used in the spatial model. However, because the correlations are fitted separately for the two dimensions, the row.factor and col.factor are needed for all models and is used for a dimension that does not involve a correlation/variance function for the fit being performed. Also, the correlation models are fitted as random terms and so the correlation model will include a variance parameter for the grid even when ar1 is used to specify the correlation model, i.e. the model fitted is a variance model and there is no difference between ar1 and ar1v in fitting the model. The variance parameter for this term represents the spatial variance and the fit necessarily includes a nugget term, this being the residual variance. If any correlation is retained, the need for a nugget term is assessed by fixing the residual variance to one, which will have no effect if heterogeneous residual variances have been specified. Once the fitting of the correlation model has been completed, the rmboundary function will be executed with the checkboundaryonly value supplied in the addSpatialModelOnIC.asrtests call.

The TPNSS spatial model is as described by Verbyla et al. (2018) and the TPPS model is as described by Rodriguez-Alvarez et al. (2018). However, for the TPPS model, the degree of the polynomial and the order of differencing can be varied. The defaults of 3 and 2, respectively, fit a cubic spline with second order differencing, while setting both the degree and order of differencing to 1 will fit a type of linear variance model (Piepho, Boer and Williams, 2022) The fixed terms for these models are row.covar + col.covar + row.covar:col.covar and the random terms are spl(row.covar) + spl(col.covar) + dev(row.covar) + dev(col.covar) + spl(row.covar):col.covar + row.covar:spl(col.covar) + spl(row.covar):spl(col.covar). The supplied model should not include any of these terms. However, any fixed or random main-effect term for either dropRowterm or dropColterm will be removed from the fit.

The TPPS model is fitted using the function tpsmb from the R package TPSbits authored by Sue Welham (2022). There are two methods for supplying the spline basis information produced by tpsmb to asreml. The grp method adds the it to the data.frame holding the information for the analysis. The mbf method requires the spline basis information to be in the same environment as the function that is called to make a fit using asreml. To this end and prior to invoking the calling function, makeTPPSplineMats.data.frame must be used produce the data.frames.

All models utilize the function changeModelOnIC.asrtests to assess the model fit, the information criteria used in assessing the fit being calculated using infoCriteria. Arguments from tpsmb and changeModelOnIC.asrtests can be supplied in calls to addSpatialModelOnIC.asrtests and will be passed on to the relevant function through the ellipses argument (...).

The data for experiment can be divided sections and the same spatial model fitted separately to each. The fit over all of the sections is assessed.

Each combination of a row.coords and a col.coords does not have to specify a single observation; for example, to fit a local spatial model to the main units of a split-unit design, each combination would correspond to a main unit and all subunits of the main unit would have the same combination.
addSpatialModelOnIC.asrtests

Value

An `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary` for the model whose fit has the smallest information criterion between the supplied and spatial model. The values of the degrees of freedom and the information criteria in the `test.summary` are differences between those of the changed model and those of the model supplied to `addSpatialModelOnIC`.

Author(s)

Chris Brien

References


Welham, S. J. (2022) **TPSbits**: Creates Structures to Enable Fitting and Examination of 2D Tensor-Product Splines using ASReml-R. Version 1.0.0 [https://mmade.org/tpsbits/](https://mmade.org/tpsbits/)

See Also

- `as.asrtests`, `rmboundary.asrtests`, `testranfix.asrtests`, `testresidual.asrtests`, `newfit.asreml`, `reparamSigDevn.asrtests`, `addSpatialModel.asrtests`, `changeModelOnIC.asrtests`, `chooseSpatialModelOnIC.asrtests`, `changeModelOnIC.asrtests`, `infoCriteria.asreml`

Examples

```r
## Not run:

data(Wheat.dat)

#Add row and column covariates
Wheat.dat <- within(Wheat.dat,
{
  cColumn <- dae::as.numfac(Column)
  cColumn <- cColumn - mean(unique(cColumn))
  cRow <- dae::as.numfac(Row)
  cRow <- cRow - mean(unique(cRow))
})

#Fit initial model
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
  random = ~ Row + Column,
  data=Wheat.dat)

#Create an asrtests object, removing boundary terms
current.asrt <- as.asrtests(current.asr, NULL, NULL,
  label = "Random Row and Column effects")
current.asrt <- rmboundary(current.asrt)
```
addto.test.summary

```r
current.asrt <- addSpatialModelOnIC(current.asrt, spatial.model = "TPPS",
  row.covar = "cRow", col.covar = "cColumn",
  dropRowterm = "Row", dropColterm = "Column",
  asreml.option = "grp")
infoCriteria(current.asrt$asreml.obj)
## End(Not run)
```

addto.test.summary  

**Description**

A row that summarizes the result of a proposed change to a model is added to a test.summary data.frame. Only the values of those arguments for which there are columns in test.summary will be included in the row.

**Usage**

```r
addto.test.summary(test.summary, terms, DF = 1, denDF = NA, 
  p = NA, AIC = NA, BIC = NA, 
  action = "Boundary")
```

**Arguments**

- **test.summary**: A data.frame whose columns are a subset of terms, DF, denDF, p, AIC, BIC and action. Each row summarizes the results of proposed changes to the fitted model. See asreml.object for more information.
- **terms**: A character giving the name of a term that might be added to or removed from the model or a label indicating a change that might be made to the model.
- **DF**: A numeric giving the numerator degrees of freedom for a Wald F-statistic or the number of variance parameters in the current model minus the number in the proposed model.
- **denDF**: A numeric giving the denominator degrees of freedom for a Wald F-statistic.
- **p**: A numeric giving the p-value for a Wald F-statistic or REML ratio test.
- **AIC**: A numeric giving Akaike Information Criterion (AIC) for a model or the difference between the AIC values for the current and proposed models.
- **BIC**: A numeric giving Bayesian (Schwarz) Information Criterion for a model or the difference between the AIC values for the current and proposed models.
- **action**: A character giving what action was taken with respect to the proposed change. See asrtests.object for more information.

**Value**

A data.frame.

**Author(s)**

Chris Brien
See Also

asremlPlus-package, asrtests.object, print.test.summary

Examples

```r
## Not run:
data(Wheat.dat)

## Fit an autocorrelation model
ar1.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                   random = ~ Row + Column + units,
                   residual = ~ ar1(Row):ar1(Column),
                   data=Wheat.dat)
ar1.asrt <- as.asrtests(ar1.asr, NULL, NULL,
                   label = "Autocorrelation model")
ar1.asrt <- rmboundary.asrtests(ar1.asrt)

## Fit a tensor spline
Wheat.dat <- within(Wheat.dat,
                   {
                     cRow <- dae::as.numfac(Row)
                     cRow <- cRow - mean(unique(cRow))
                     cColumn <- dae::as.numfac(Column)
                     cColumn <- cColumn - mean(unique(cColumn))
                   })
ts.asr <- asreml(yield ~ Rep + cRow + cColumn + WithinColPairs +
                 Variety,
                 random = ~ spl(cRow) + spl(cColumn) +
                 dev(cRow) + dev(cColumn) +
                 spl(cRow):cColumn + cRow:spl(cColumn) +
                 spl(cRow):spl(cColumn),
                 residual = ~ Row:Column,
                 data=Wheat.dat)
ts.asrt <- as.asrtests(ts.asr, NULL, NULL,
                   label = "Tensor spline model")
ts.asrt <- rmboundary.asrtests(ts.asrt)
ar1.ic <- infoCriteria(ar1.asr$asreml.obj)
ts.ic <- infoCriteria(ts.asr$asreml.obj)
if (ar1.ic$AIC < ts.ic$AIC)
{
ic.diff <- ar1.ic - ts.ic
new.asrt <- ar1.asrt
new.asrtest$test.summary <- addto.test.summary(new.asrtest$test.summary,
                                                  terms = "Compare ar1 to ts",
                                                  DF = ic.diff$varDF,
                                                  AIC = ic.diff$AIC, BIC = ic.diff$BIC,
                                                  action = "Chose ar1")
}
else
{
ic.diff <- ts.ic - ar1.ic
new.asrt <- ts.asrt
new.asrtest$test.summary <- addto.test.summary(new.asrtest$test.summary,
                                                  terms = "Compare ar1 to ts",
                                                  DF = ic.diff$varDF,
                                                  AIC = ic.diff$AIC, BIC = ic.diff$BIC,
                                                  action = "Chose ts")
}
```
allDifferences.data.frame

Using supplied predictions and standard errors of pairwise differences or the variance matrix of predictions, forms all pairwise differences between the set of predictions, and p-values for the differences.

Description

Uses supplied predictions and standard errors of pairwise differences, or the variance matrix of predictions to form, in an `alldiffs.object`, for those components not already present, (i) a table of all pairwise differences of the predictions, (ii) the p-value of each pairwise difference, and (iii) the minimum, mean, maximum and accuracy of LSD values. Predictions that are aliased (or non-estimable) are removed from the predictions component of the `alldiffs.object` and standard errors of differences involving them are removed from the `sed` component.

If necessary, the order of the columns of the variables in the predictions component are changed to be the initial columns of the `predictions.frame` and to match their order in the `classify`. Also, the rows of predictions component are ordered so that they are in standard order for the variables in the `classify`. That is, the values of the last variable change with every row, those of the second-last variable only change after all the values of the last variable have been traversed; in general, the values of a variable are the same for all the combinations of the values to the variables to its right in the `classify`. The `sortFactor` or `sortOrder` arguments can be used to order of the values for the `classify` variables, which is achieved using `sort.alldiffs`.

Each p-value is computed as the probability of a t-statistic as large as or larger than the absolute value of the observed difference divided by its standard error. The p-values are stored in the `p.differences` component. The degrees of freedom of the t-distribution is the degrees of freedom stored in the `tdf` attribute of the `alldiffs.object`. This t-distribution is also used in calculating the LSD statistics stored in the `LSD` component of the `alldiffs.object`.

Usage

```r
## S3 method for class 'data.frame'
allDifferences(predictions, classify, vcov = NULL,
differences = NULL, p.differences = NULL, sed = NULL,
LSD = NULL, LSDtype = "overall", LSDsupplied = NULL,
LSDby = NULL, LSDstatistic = "mean",
LSDaccuracy = "maxAbsDeviation",
retain.zerolSDs = FALSE,
zero.tolerance = .Machine$double.eps ^ 0.5,
backtransforms = NULL,
response = NULL, response.title = NULL,
term = NULL, tdf = NULL,
x.num = NULL, x.fac = NULL,
level.length = NA,
pairwise = TRUE, alpha = 0.05,
transform.power = 1, offset = 0, scale = 1,
transform.function = "identity",
)```
```
inestimable.rm = TRUE,
sortFactor = NULL, sortParallelToCombo = NULL,
sortNestingFactor = NULL, sortOrder = NULL,
decreasing = FALSE, ...)

Arguments

predictions A predictions.frame, or a data.frame, beginning with the variables classifying the predictions and also containing columns named predicted.value, standard.error and est.status; each row contains a single predicted value. It may also contain columns for the lower and upper limits of error intervals for the predictions. Note that the names standard.error and est.status have been changed to std.error and status in the pvals component produced by asreml-R4; if the new names are in the data.frame supplied to predictions, they will be returned to the previous names.

classify A character string giving the variables that define the margins of the multiway table that has been predicted. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator.

cov A matrix containing the variance matrix of the predictions; it is used in computing the variance of linear transformations of the predictions.

differences A matrix containing all pairwise differences between the predictions; it should have the same number of rows and columns as there are rows in predictions.

ee differences A matrix containing p-values for all pairwise differences between the predictions; each p-value is computed as the probability of a t-statistic as large as or larger than the observed difference divided by its standard error. The degrees of freedom of the t distribution for computing it are computed as the denominator degrees of freedom of the F value for the fixed term, if available; otherwise, the degrees of freedom stored in the attribute tdf are used; the matrix should be of the same size as that for differences.

esed A matrix containing the standard errors of all pairwise differences between the predictions; they are used in computing the p-values.

LSD An LSD.frame containing the mean, minimum and maximum LSD for determining the significance of pairwise differences, as well as an assigned LSD and a measure of the accuracy of the LSD. If LSD is NULL then the LSD.frame stored in the LSD component will be calculated and the values of LSDtype, LSDby and LSDstatistic added as attributes of the alldiffs.object. The LSD for a single prediction assumes that any predictions to be compared are independent; this is not the case if residual errors are correlated.

LSDtype A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparison2, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations.
See `LSD.frame` for further information on the values in a row of this `data.frame` and how they are calculated.

LSDsupplied
A `data.frame` or a named `numeric` containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the `predictions.frame` or a single LSD value that is an overall LSD. If a `data.frame`, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; `assignedLSD` is sensible, but not obligatory. Otherwise, a `numeric` containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function `dae::fac.combine` to the `predictions` component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into `assignedLSD` column of the `LSD.frame` stored as the LSD component of the `alldiffs.object`.

LSDby
A `character` (vector) of variables names, being the names of the `factors` or `numeros` in the `classify`; for each combination of their levels and values, there will be or is a row in the `LSD.frame` stored in the LSD component of the `alldiffs.object` when LSDtype is `factor.combinations`.

LSDstatistic
A `character` nominating one or more of `minmum`, `q10`, `q25`, `mean`, `median`, `q75`, `q90` or `maximum` as the value(s) to be stored in the `assignedLSD` column in an `LSD.frame`; the values in the `assignedLSD` column are used in computing halfLeas Significant error.intervals. Here `q10`, `q25`, `q75` and `q90` indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function `quantile` is used to obtain them. The `mean` LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The `median` is calculated using the `median` function. Multiple values are only produced for LSDtype set to `factor.combination`, in which case LSDby must not be `NULL` and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is `NULL`, it is reset to `mean`.

LSDaccuracy
A `character` nominating one of `maxAbsDeviation`, `maxDeviation`, `q90Deviation` or `RootMeanSqDeviation` as the statistic to be calculated as a measure of the accuracy of `assignedLSD`. The option `q90Deviation` produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named `accuracyLSD` in an `LSD.frame`.

retain.zeroLSDs
A `logical` indicating whether to retain or omit LSDs that are zero when calculating the summaries of LSDs.

zero.tolerance
A `numeric` specifying the value such that if an LSD is less than it, it will be considered to be zero.

backtransforms
A `data.frame` containing the backtransformed values of the predicted values that is consistent with the `predictions` component, except that the column named `predicted.value` is replaced by one called `backtransformed.predictions`. Any `error.interval` values will also be the backtransformed values. Each row contains a single predicted value.

response
A `character` specifying the response variable for the predictions. It is stored as an attribute to the `alldiffs.object`. 
response.title  A character specifying the title for the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

term  A character string giving the variables that define the term that was fitted using asreml and that corresponds to classify. It only needs to be specified when it is different to classify; it is stored as an attribute of the alldiffs.object. It is likely to be needed when the fitted model includes terms that involve both a numeric covariate and a factor that parallel each other; the classify would include the covariate and the term would include the factor.

tdf  an integer specifying the degrees of freedom of the standard error. It is used as the degrees of freedom for the t-distribution on which p-values and confidence intervals are based. It is stored as an attribute to the alldiffs.object.

x.num  A character string giving the name of the numeric covariate that corresponds to x.fac, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of unique values as the number of levels in x.fac.

x.fac  A character string giving the name of the factor that corresponds to x.num, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of levels as the number of unique values in x.num. The levels of x.fac must be in the order in which they are to be plotted - if they are dates, then they should be in the form yyyyymmdd, which can be achieved using as.Date. However, the levels can be non-numeric in nature, provided that x.num is also set.

level.length  The maximum number of characters from the levels of factors to use in the row and column labels of the tables of pairwise differences and their p-values and standard errors.

pairwise  A logical indicating whether all pairwise differences of the predictions and their standard errors and p-values are to be computed and stored. If FALSE, the components differences and p.differences will be NULL in the returned alldiffs.object.

alpha  A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

transform.power  A numeric specifying the power of a transformation, if one has been applied to the response variable. Unless it is equal to 1, the default, back-transforms of the predictions will be obtained and presented in tables or graphs as appropriate. The back-transformation raises the predictions to the power equal to the reciprocal of transform.power, unless it equals 0 in which case the exponential of the predictions is taken.

offset  A numeric that has been added to each value of the response after any scaling and before applying any power transformation.

scale  A numeric by which each value of the response has been multiplied before adding any offset and applying any power transformation.

transform.function  A character giving the name of a function that specifies the scale on which the predicted values are defined. This may be the result of a transformation of the data using the function or the use of the function as a link function in the fitting of a generalized linear (mixed) model (GL(M)M). The possible transform.functions are identity, log, inverse, sqrt, logit, probit, and cloglog. The predicted.values and error.intervals, if not StandardError
intervals, will be back-transformed using the inverse function of the transform.function. The standard.error column will be set to NA, unless (i) asreml returns columns named transformed.value and approx.se, as well as those called predicted.values and standard.error (such as when a GLM is fitted) and (ii) the values in transformed.value are equal to those obtained by backtransforming the predicted.values using the inverse function of the transform.function. Then, the approx.se values will be saved in the standard.error column of the backtransforms component of the returned alldiffs.obj. Also, the transformed.value and approx.se columns are removed from both the predictions and backtransforms components of the alldiffs.obj. Note that the values that end up in the standard.errors column are approximate for the backtransformed values and are not used in calculating error.intervals.

inestimable.rm A logical indicating whether rows for predictions that are not estimable are to be removed from the components of the alldiffs.object.

sortFactor A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.

sortParallelToCombo A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the supplied list is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.

sortNestingFactor A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

sortOrder A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the desired order of the levels in the reordered components of the alldiffs.object. The argument sortParallelToCombo is ignored.

The following creates a sortOrder vector lev for factor f based on the values in x:
lev <- levels(f)[order(x)].

decreasing A logical passed to order that determines whether the order for sorting the components of the alldiffs.object is for increasing or decreasing magnitude of the predicted values.
... provision for passing arguments to functions called internally - not used at present.

Value

An `alldiffs.object` with components `predictions, vcov, differences, p.differences, sed,` and LSD.

The name of the response, the response.title, the term, the classify, tdf, alpha, sortFactor and the sortOrder will be set as attributes to the object. Note that the classify in an `alldiffs.object` is based on the variables indexing the predictions, which may differ from the classify used to obtain the original predictions (for example, when the `alldiffs.objects` stores a linear transformation of predictions.

Also, see `predictPlus.asreml` for more information.

Author(s)

Chris Brien

See Also

`asremlPlus-package, as.alldiffs, as.predictions.frame, sort.alldiffs, subset.alldiffs, print.alldiffs, renewClassify.alldiffs, redoErrorIntervals.alldiffs, recalclSD.alldiffs, pickLSDstatistics.alldiffs, plotPredictions.data.frame, predictPlus.asreml, predictPresent.asreml`

Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety, random=~Blocks/Wplots, data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety", sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- Var.pred$pvals
Var.sed <- Var.pred$sed
Var.vcov <- NULL
wald.tab <- current.asrt$wald.tab
den.df <- wald.tab[match("Variety", rownames(wald.tab)), "denDF"]

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE)) {
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots), data=Oats.dat)
  Var.emm <- emmeans::emmeans(m1.lmer, specs = ~ Nitrogen:Variety)
```r
Var.preds <- summary(Var.emm)
den.df <- min(Var.preds$df)
## Modify Var.preds to be compatible with a predictions.frame
Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                   se = "SE", interval.type = "CI",
                                   interval.names = c("lower.CL", "upper.CL"))
Var.vcov <- vcov(Var.emm)
Var.sed <- NULL
}

## Use the predictions obtained with either asreml or lmerTest
if (exists("Var.preds"))
{
## Order the Varieties in decreasing order for the predictions values in the
## first N level
Var.diffs <- allDifferences(predictions = Var.preds,
                            classify = "Nitrogen:Variety",
                            sed = Var.sed, vcov = Var.vcov, tdf = den.df,
                            sortFactor = "Variety", decreasing = TRUE)
print.alldiffs(Var.diffs, which="differences")

## Change the order of the factors in the alldiffs object and reorder components
Var.reord.diffs <- allDifferences(predictions = Var.preds,
                                   classify = "Variety:Nitrogen",
                                   sed = Var.sed, vcov = Var.vcov, tdf = den.df)
print.alldiffs(Var.reord.diffs, which="predictions")
}
```

---

### alldiffs.object

**Description of an alldiffs object**

An object of S3-class `alldiffs` that stores the predictions for a model, along with supplied statistics for all pairwise differences. While `alldiffs.object` can be constructed by defining a list with the appropriate components, it can be formed by passing the components to `as.alldiffs`, or from a predictions data.frame using `allDifferences.data.frame`. `as.alldiffs` is a function that assembles an object of this class from supplied components.

`is.alldiffs` is the membership function for this class; it tests that an object is of class `alldiffs`. `validAlldiffs(object)` can be used to test the validity of an object with this class.

`allDifferences.data.frame` is the function that constructs an object of this class by calculating components from statistics supplied via its arguments and then using `as.alldiffs` to make the object.

### Value

A list of class `alldiffs` containing the following components: `predictions`, `vcov`, `differences`, `p.differences`, `sed`, `LSD` and `backtransforms`. Except for predictions, the components are optional and can be set to NULL.

An `alldiffs.object` also has attributes `response`, `response.title`, `term`, `classify`, `tdf`, `alpha`, `sortFactor` and `sortOrder`, which may be set to NULL.

The details of the components are as follows:
1. predictions: A `predictions.frame`, being a data.frame beginning with the variables classifying the predictions, in the same order as in the `classify`, and also containing columns named `predicted.value`, `standard.error` and `est.status`; each row contains a single predicted value. The number of rows should equal the number of unique combinations of the classify variables and will be in standard order for the classify variables. That is, the values of the last variable change with every row, those of the second-last variable only change after all the values of the last variable have been traversed; in general, the values of a variable are the same for all the combinations of the values to the variables to its right in the classify. The data.frame may also include columns for the lower and upper values of error intervals, either standard error, confidence or half-LSD intervals. The names of these columns will consist of three parts separated by full stops: 1) the first part will be `lower` or `upper`; 2) the second part will be one of `Confidence`, `StandardError` or `halfLeastSignificant`; 3) the third component will be `limits`. Note that the names `standard.error` and `est.status` have been changed to `std.error` and `status` in the `pvals` component produced by `asreml-R4`; if the new names are in the data.frame supplied to `predictions`, they will be returned to the previous names.

2. differences: A matrix containing all pairwise differences between the predictions; it should have the same number of rows and columns as there are rows in `predictions`.

3. p.differences: A matrix containing p-values for all pairwise differences between the predictions; each p-value is computed as the probability of a t-statistic as large as or larger than the observed difference divided by its standard error. The degrees of freedom of the t distribution for computing it are computed as the denominator degrees of freedom of the F value for the fixed term, if available; otherwise, the degrees of freedom stored in the attribute `tdf` are used; the matrix should be of the same size as that for `differences`.

4. sed: A matrix containing the standard errors of all pairwise differences between the predictions; they are used in computing the p-values in `p.differences`.

5. vcov: A matrix containing the variance matrix of the predictions; it is used in computing the variance of linear transformations of the predictions.

6. LSD: An LSD.frame containing (i) c, the number of pairwise predictions comparisons for each LSD value and the mean, minimum, maximum and assigned LSD, (ii) the column accuracyLSD that gives a measure of the accuracy of the assigned LSD, given the variation in LSD values, and (iii) the columns `false.pos` and `false.neg` that contain the number of false positives and negatives if the assignedLSD value(s) is(are) used to determine the significance of the pairwise predictions differences. The LSD values in the assignedLSD column is used to determine the significance of pairwise differences that involve predictions for the combination of levels given by a row name. The value in the assignedLSD column is specified using the `LSDstatistic` argument.

7. backtransforms: When the response values have been transformed for analysis, a data.frame containing the backtransformed values of the predicted values is added to the alldiffs.object. This data.frame is consistent with the predictions component, except that the column named `predicted.value` is replaced by one called backtransformed.predictions. Any error.interval values will also be the backtransformed values. Each row contains a single predicted value.

The details of the attributes of an alldiffs.object are:

1. response: A character specifying the response variable for the predictions.
2. response.title: A character specifying the title for the response variable for the predictions.
3. term: A character giving the variables that define the term that was fitted using asreml and that corresponds to classify. It is often the same as classify.
4. classify: A character giving the variables that define the margins of the multiway table used in the prediction. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator.
5. tdf: An integer specifying the degrees of freedom of the standard error. It is used as the degrees of freedom for the t-distribution on which p-values and confidence intervals are based.
6. alpha: An integer specifying the significance level. It is used as the significance level calculating LSDs.
7. LSDtype: If the LSD component is not NULL then LSDtype is added as an attribute. A character nominating the type of grouping of seds to be used in combining LSDs.
8. LSDby: If the LSD component is not NULL then LSDby is added as an attribute. A character vector containing the names of the factors and numerics within whose combinations the LSDs are to be summarized.
9. LSDstatistic: If the LSD component is not NULL then LSDstatistic is added as an attribute. A character nominating what statistic to use in summarizing a set of LSDs.
10. LSDaccuracy: If the LSD component is not NULL then LSDaccuracy is added as an attribute. A character nominating the method of calculating a measure of the accuracy of the LSDs stored in the assignedLSD column of the LSD.frame.
11. sortFactor: factor that indexes the set of predicted values that determined the sorting of the components.
12. sortOrder: A character vector that is the same length as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the order of the levels in the reordered components of the alldiffs.object.

The following creates a sortOrder vector levs for factor f based on the values in x:
levs <- levels(f)[order(x)].
See predictPlus.asreml for more information.

Author(s)
Chris Brien

See Also
is.alldiffs, as.alldiffs, validAlldiffs, allDifferences.data.frame

Examples

data(Oats.dat)
## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety,
                   random=Blocks/Wplots,
                   data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety",
                                   sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- Var.pred$pvals
Var.sed <- Var.pred$sed
## End(Not run)

### Use lmerTest and emmmeans to get predictions and associated statistics

```r
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE)) {
  ml.1mer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
                           data=Oats.dat)
  Var.emm <- emmeans::emmeans(ml.1mer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  den.df <- min(Var.preds$df)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                    se = "SE", interval.type = "CI",
                                    interval.names = c("lower.CL", "upper.CL"))

  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
}
```

### Use the predictions obtained with either asreml or lmerTest

```r
if (exists("Var.preds")) {
  ## Form an all.diffs object
  Var.diffs <- as.alldiffs(predictions = Var.preds, classify = "Nitrogen:Variety",
                           sed = Var.sed, vcov = Var.vcov, tdf = den.df)

  ## Check the class and validity of the alldiffs object
  is.alldiffs(Var.diffs)
  validAlldiffs(Var.diffs)
}
```

---

**angular**

Applies the angular transformation to proportions.

### Description

Applies the angular transformation to numeric values. It is given by $\sin^{-1}(\sqrt{\text{proportions}})$

### Usage

```r
angular(proportions, n)
```

### Arguments

- `proportions` The proportions.
- `n` The divisor(s) for each proportion

### Value

A numeric.
**angular.mod**

**Author(s)**

Chris Brien

**See Also**

`angular.mod`, `powerTransform`.

**Examples**

```r
n <- 25
y <- rbinom(10, n, 0.5)
y <- c(y, 0, n)
p <- y/n
p.ang <- angular(p, n)
```

---

**angular.mod**

Applies the modified angular transformation to a vector of counts.

**Description**

Applies the angular transformation to a vector of counts. A modified transformation is used that is appropriate when N < 50 and the proportion is not between 0.3 and 0.7. The transformation is given by

\[
\sin\left(1 - \frac{\text{count} + 0.375}{\text{n} + 0.75}\right) = \arcsin\left(\sqrt{\frac{(\text{count} + 0.375)}{(\text{n} + 0.75)}}\right).
\]

**Usage**

`angular.mod(count, n)`

**Arguments**

- `count` The numeric vector of counts.
- `n` The number(s) of observations from which the count(s) were obtained.

**Value**

A numeric vector.

**Author(s)**

Chris Brien

**See Also**

`angular`, `powerTransform`.

**Examples**

```r
n <- 25
y <- rbinom(10, n, 0.5)
y <- c(y, 0, n)
p.ang.mod <- angular.mod(y, n)
```
as.alldiffs  

Forms an alldiffs.object from the supplied predictions, along with 
those statistics, associated with the predictions and their pairwise dif- 
terences, that have been supplied.

Description

Creates an alldiffs.object that consists of a list containing the following components: predictions, 
vcov, differences, p.differences, sed, LSD and backtransforms. Predictions must be sup- 
plied to the function while the others will be set only if they are supplied; those not supplied are 
set to NULL. It also has attributes response, response.title, term, classify, tdf, tdf, alpha, 
sortFactor and sortOrder. which will be set to the values supplied or NULL if none are supplied.

Usage

as.alldiffs(predictions, vcov = NULL, differences = NULL, 
p.differences = NULL, sed = NULL, LSD = NULL, 
backtransforms = NULL, 
response = NULL, response.title = NULL, 
term = NULL, classify = NULL, 
tdf = NULL, alpha = 0.05, 
sortFactor = NULL, sortOrder = NULL)

Arguments

predictions    A predictions.frame, being a data.frame beginning with the variables class- 
sifying the predictions and also containing columns named predicted.value, 
standard.error and est.status; each row contains a single predicted value. 
It may also contain columns for the lower and upper limits of error intervals for 
the predictions. Note that the names standard.error and est.status have 
been changed to std.error and status in the pvals component produced by 
asreml-R4; if the new names are in the data.frame supplied to predictions, 
they will be returned to the previous names.

differences    A matrix containing all pairwise differences between the predictions; it should 
have the same number of rows and columns as there are rows in predictions.

p.differences  A matrix containing p-values for all pairwise differences between the predic- 
tions; each p-value is computed as the probability of a t-statistic as large as or 
larger than the observed difference divided by its standard error. The degrees of 
freedom of the t distribution for computing it are computed as the denominator 
degrees of freedom of the F value for the fixed term, if available; otherwise, the 
degrees of freedom stored in the attribute tdf are used; the matrix should be of 
the same size as that for differences.

sed            A matrix containing the standard errors of all pairwise differences between the 
predictions; they are used in computing the p-values.

vcov           A matrix containing the variance matrix of the predictions; it is used in com- 
puting the variance of linear transformations of the predictions.

LSD            An LSD.frame containing the mean, minimum and maximum LSD for deter- 
mining the significance of pairwise differences, as well as an assigned LSD and 
a measure of the accuracy of the LSD. If LSD is NULL then the LSD.frame stored
in the LSD component will be calculated and the values of LSDtype, LSDby and LSDstatistic added as attributes of the alldiffs.object. The LSD for a single prediction assumes that any predictions to be compared are independent; this is not the case if residual errors are correlated.

backtransforms A data.frame containing the backtransformed values of the predicted values that is consistent with the predictions component, except that the column named predicted.value is replaced by one called backtransformed.predictions. Any error.interval values will also be the backtransformed values. Each row contains a single predicted value.

response A character specifying the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

response.title A character specifying the title for the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

term A character string giving the variables that define the term that was fitted using asreml and that corresponds to classify. It only needs to be specified when it is different to classify; it is stored as an attribute of the alldiffs.object. It is likely to be needed when the fitted model includes terms that involve both a numeric covariate and a factor that parallel each other; the classify would include the covariate and the term would include the factor.

classify A character string giving the variables that define the margins of the multiway table used in the prediction. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator. It is stored as an attribute to the alldiffs.object.

tdf an integer specifying the degrees of freedom of the standard error. It is used as the degrees of freedom for the t-distribution on which p-values and confidence intervals are based. It is stored as an attribute to the alldiffs.object.

alpha A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

sortFactor A character containing the name of the factor that indexes the set of predicted values that determined the sorting of the components.

sortOrder A character vector that is the same length as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the order of the levels in the reordered components of the alldiffs.object. The following creates a sortOrder vector levs for factor f based on the values in x: levs <- levels(f)[order(x)].

Value
An S3-class alldiffs.object. Also, see predictPlus.asreml for more information.

Author(s)
Chris Brien

See Also
asremlPlus-package, alldiffs.object, is.alldiffs, as.alldiffs, print.alldiffs, sort.alldiffs, subset.alldiffs, allDifferences.data.frame, renewClassify.alldiffs, redoErrorIntervals.alldiffs, recalclSD.alldiffs, predictPlus.asreml, plotPredictions.data.frame, predictPresent.asreml
Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety,
                 random=~Blocks/Wplots,
                 data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety",
                  sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- Var.pred$pvals
Var.sed <- Var.pred$sed
Var.vcov <- NULL

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
     requireNamespace("emmmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
                 data=Oats.dat)
  Var.emm <- emmmeans::emmmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  den.df <- min(Var.preds$df)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                    se = "SE", interval.type = "CI",
                                    interval.names = c("lower.CL", "upper.CL"))
  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
}

## Use the predictions obtained with either asreml or lmerTest
if (exists("Var.preds"))
{
  ## Form an all.diffs object
  Var.diffs <- as.alldiffs(predictions = Var.preds, classify = "Nitrogen:Variety",
                           sed = Var.sed, vcov = Var.vcov, tdf = den.df)

  ## Check the class and validity of the alldiffs object
  is.alldiffs(Var.diffs)
  validAlldiffs(Var.diffs)
}

as.asrtests
Forms an asrtests object that stores (i) a fitted asreml object, (ii) a
pseudo-anova table for the fixed terms and (iii) a history of changes
and hypothesis testing used in obtaining the model.
as.asrtests

Description

An asrtests.object that is a list consisting of the components asreml.obj, wald.tab and test.summary.

A call to as.asrtests with test.summary = NULL re-initializes the test.summary data.frame.

If there is no wald.tab, wald.asreml is called. In all cases, recalcWaldTab is called and any changes made as specified by the the recalcWaldTab arguments supplied via ....

The label argument can be used to include an entry in test.summary for the starting model. If a label is included, (i) the information criteria calculated using the asreml.obj will be added to the test.summary, if IClikelihood is not set to none and (ii) the number of variance parameters is included in the denDF column, if IClikelihood is set to none.

Usage

as.asrtests(asreml.obj, wald.tab = NULL, test.summary = NULL,
            denDF = "numeric", label = NULL,
            IClikelihood = "none", bound.exclusions = c("F","B","S","C"), ...)

Arguments

asreml.obj an asreml object for a fitted model.
wald.tab A data.frame containing a pseudo-anova table for the fixed terms produced by wald.asreml; it should have 4 or 6 columns. Sometimes wald.asreml returns a data.frame and at other times a list. For example, it may return a list when denDF is used. In this case, the Wald component of the list is to be extracted and stored. It is noted that, as of asreml version 4, wald.asreml has a kenadj argument.
test.summary A data.frame with columns term, DF, denDF, p and action containing the results of previous hypothesis tests.
denDF Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.
label A character to use as an entry in the terms column in test.summary to indicate as far as is possible the nature of the model that has been fitted. The action column in test.summary will be Starting model.
IClikelihood A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new asrtests.object. If none, none are included. Otherwise, if REML, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full, then the AIC and BIC based on the full likelihood are included. (See also infoCriteria.asreml.)
bound.exclusions A character specifying the bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in calculating information criteria. If set to NULL then none will be excluded.
... further arguments passed to wald.asreml and recalcWaldTab.
as.predictions.frame

Forms a predictions.frame from a data.frame, ensuring that the correct columns are present.

Description

Creates a predictions.frame from a data.frame by adding the class predictions.frame to it, and renaming the columns containing the predictions, se, est.status and error.intervals.

Usage

as.predictions.frame(data, classify = NULL,
    predictions = NULL, se = NULL, est.status = NULL,
    interval.type = NULL, interval.names = NULL)
Arguments

data

A `data.frame` containing columns giving the variables that uniquely index the predicted values and columns with the predicted values, their standard errors and, optionally, their estimation status (`est.status`).

classify

A `character` string giving the variables that define the margins of the multiway table that was predicted. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the `:` operator. For predicting the overall mean, the classify is set to "(Intercept)".

predictions

A `character` giving the name of the column in `data` that contains the predicted values. This column will be renamed to `predicted.value`.

se

A `character` giving the name of the column in `data` that contains the standard errors of the predicted values. This column will be renamed to `standard.error`.

est.status

A `character` giving the name of the column in `data` that contains the estimation status of the predicted values. It will have a value `Estimable` for predicted values that have been estimated and a value `Aliased` for predicted values that are `NA`. If a column named `est.status` is not present in `data` and `est.status` is `NULL`, a column `est.status` will be generated.

interval.type

A `character` specifying the type of `error.intervals` stored in `data` that require renaming. If `NULL`, `error.intervals` will not be renamed, even if they are present. Otherwise, `interval.type` should be set to one of "CI", "SE" or "halfLSD".

interval.names

A `character` specifying the column names of the lower and upper limits stored in `data` that are to be renamed. The character must be of length two, with the first element being the name of the 'lower' limit and the second element being the name of the 'upper' limit.

Value

An S3-class `predictions.frame`.

Author(s)

Chris Brien

See Also

`asremlPlus-package`, `predictions.frame`, `is.predictions.frame`, `predictions.frame`, `validPredictionsFrame`

Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics

## Not run:
ml.asr <- asreml(Yield ~ Nitrogen*Variety,
random=~Blocks/Wplots,
data=Oats.dat)
current.asrt <- as.asrtests(ml.asr)
Var.pred <- asreml::predict.asreml(ml.asr, classify="Nitrogen:Variety",
## asremlPlus-deprecated

### Deprecated Functions in the Package asremlPlus

These functions have been renamed and deprecated in asremlPlus:

1. `addrm.terms.asreml` and `addrm.terms.asrtests` -> `changeTerms.asrtests`,
2. `alldiffs` -> `as.alldiffs`,
3. `asrtests` -> `as.asrtests`,
4. `choose.model.asreml` and `choose.model.asrtests` -> `chooseModel.asrtests`,
5. `facRecode` and `facRecode.alldiffs` -> `facRecast.alldiffs`,
6. `info.crit` and `info.crit.asreml` -> `infoCriteria.asreml`,
7. `newrcov.asrtests` -> `changeTerms.asrtests`,
8. `plotvariofaces.asreml` -> `plotVariofaces.data.frame`,
9. `power.transform` -> `powerTransform`,
10. `predictiondiffs.asreml` -> `allDifferences.data.frame`,
11. `predictionplot.asreml` -> `plotPredictions.data.frame`,
12. `predictparallel.asreml` -> `predictPlus.asreml`,
13. `pred.present.asreml` -> `predictPresent.asreml`,
14. `recalc.wald.tab.asreml` and `recalc.wald.tab.asrtests` -> `recalcWaldTab.asrtests`.
15. `reorderClassify` and `reorderClassify.alldiffs` -> `renewClassify.alldiffs`,
16. `reml.lrt` and `reml.lrt.asreml` -> `REMLRT.asreml`,
17. `rmboundary.asreml` -> `rmboundary.asrtests`,
18. `setvarianceterms.asreml` -> `setvarianceterms.call`,
19. `sig.devn.reparam.asreml` and `sig.devn.reparam.asrtests` -> `reparamSigDevn.asrtests`,
20. `testranfix.asreml` -> `testranfix.asrtests`,
21. `testrcov.asreml` and `testrcov.asrtests` -> `testresidual.asrtests`,
22. `testswapran.asreml` -> `testswapran.asrtests`

Usage

```r
addrm.terms.asreml(...)  
addrm.terms.asrtests(...)  
alldiffs(...)  
asrtests(...)  
choose.model.asreml(...)  
choose.model.asrtests(...)  
facRecode(...)  
facRecode.alldiffs(...)  
info.crit(...)  
info.crit.asreml(...)  
newrcov.asrtests(...)  
plotvariofaces.asreml(...)  
power.transform(...)  
predictiondiffs.asreml(...)  
predictionplot.asreml(...)  
predictparallel.asreml(...)  
pred.present.asreml(...)  
recalc.wald.tab.asreml(...)  
recalc.wald.tab.asrtests(...)  
reml.lrt(...)  
reml.lrt.asreml(...)  
```

## S3 method for class 'alldiffs'
reorderClassify(...)

## S3 method for class 'asreml'
rmboundary(...)

setvarianceterms.asreml(...)

`sig.devn.reparam.asreml(...)

`sig.devn.reparam.asrtests(...)

`testranfix.asreml(...)

`testrcov.asreml(...)

## S3 method for class 'asreml'
testswapran(...)

Arguments

... 
absorbs arguments passed from the old functions of the style foo.bar().
asremlPlusTips

*The randomly-presented, startup tips.*

Description

The intermittent, randomly-presented, startup tips.

Startup tips

Need help? The manual is a vignette and is in the vignettes subdirectory of the package’s install directory.

Find out what has changed in asremlPlus: enter `news(package = 'asremlPlus')`.

Need help getting started? Enter `vignette(package = 'asremlPlus')`.

To avoid start-up message that ASReml-R is needed, load asreml before asremlPlus.

The methods for alldiffs and data.frame do not require asreml

Use `suppressPackageStartupMessages()` to eliminate all package startup messages.

To see all the intermittent, randomly-presented, startup tips enter `?asremlPlusTips`.

To install the latest version: go to [http://chris.brien.name/rpackages](http://chris.brien.name/rpackages).

For versions between CRAN releases (and more) go to [http://chris.brien.name/rpackages](http://chris.brien.name/rpackages).

Author(s)

Chris Brien

---

asrtests.object

*Description of an asrtests object*

Description

An object of S3-class `asrtests` that contains information derived from the fits of a mixed model using `asreml`.

`as.asrtests` is function that makes an object of this class.

`is.list` is the membership function for this class; it tests that an object is of class `list`.

`validAsrtests` can be used to test the validity of an `asrtests.object`.
**Value**

A list that contains three components:

1. `asreml.obj`: an object of class `asreml` that contains the fit of a model;
2. `wald.tab`: A data.frame containing a pseudo-anova table for the fixed terms produced by `wald.asreml`. It has rownames that correspond to the fixed terms that were fitted and four columns. If denominator degrees of freedom were calculated then the columns are `DF`, `denDF`, `F.inc`, `Pr`; otherwise the columns are `Df`, `Sum of Sq`, `Wald statistic`, and `Pr(Chisq)`.
3. `test.summary`: A data.frame with columns `terms`, `DF`, `denDF`, `p`, `AIC`, `BIC` and action, each row of which summarizes the results of proposed changes to the fitted model. Possible codes for action are: Dropped, Retained, Swapped, Unswapped, Unchanged, Significant, Nonsignificant, Absent, Added, Removed and Boundary. If the either of the models did not converge, unconverged will be added to the code. Unchanged is used when `allow.unconverged` is `FALSE`. Note that the logical `asreml.obj$converge` also reflects whether there is convergence.

A row is added to the `test.summary` for each term that is dropped, added or tested or a note that several terms have been added or removed. When values for the AIC and BIC are included in the row, then the `DF` are the number of fixed parameters in the model and `denDF` are the numbers of variance parameters. When `changeModelOnIC` adds a row then the values of the degrees of freedom and information criteria are differences between those for the model that is supplied and the model changed by `changeModelOnIC`.

**Author(s)**

Chris Brien

**See Also**

`as.asrtests`, `as.asrtests`, `validAsrtests`

### bootREMLRT.asreml

**Uses the parametric bootstrap to calculate the p-value for a REML ratio test to compare two models.**

**Description**

Extracts the REML log likelihood for two `asreml` objects and forms the observed REML ratio statistic. It assumes that the second `asreml` object is the result of fitting a model that is a reduced version of the model for the first object and is considered to the null model. Using the `mean` and `V`, `nboot` bootstrap samples of simulated response values are generated in parallel; that is, `ncores` cores are used and each is used to generate and analyse a sample. The full and reduced models are fitted to the data and if either analysis fails to converge another sample is generated and analysed using the current core, with a maximum of `max.retries` attempts to obtain a sample that converges for both analysis. Thus the maximum number of data sets that will be generated is `nboot` * `max.retries`. If a bootstrap sample converges for both analyses, the REML ratio test statistic is formed for it. The p-value is then calculated as `(k + 1)/(b + 1)` where `k` is the number of simulated ratio test statistics greater than the observed test statistic and `s` is the number of bootstrap samples that were returned. The function checks that the models do not differ in either their fixed or sparse models. It also check the difference in the number of variance parameters between the two fits to the models, taking into account the `bound.exclusions`. 
Usage

## S3 method for class 'asreml'
bootREMLRT(h0.asreml.obj, h1.asreml.obj,
  nboot = 100, max.retries = 5, seed = NULL,
  means=NULL, V = NULL, extra.matrix = NULL, ignore.terms = NULL,
  fixed.spline.terms = NULL,
  bound.exclusions = c("F","B","S","C"),
  tolerance = 1E-10, update = TRUE, trace = FALSE,
  ncores = detectCores(), ...)

Arguments

h0.asreml.obj  asreml object containing the fit under the model for the null hypothesis.
h1.asreml.obj  asreml object containing the fit under the model for the alternative hypothesis.
nboot          The number of bootstrap samples to be generated.
max.retries    The maximum number of attempts to generate a sample whose analyses converge for both models.
seed           A single value, interpreted as an integer, that specifies the starting value of the random number generator. The "L'Ecyter-CMRG" random generator is used and nextRNGStream is used to seed each core from the original seed.
means          The vector of means to be used in generating simulated bootstrap samples. If it is NULL, the fitted values based on object are used. It must be the same length as the response variable for object.
V               The fitted variance matrix, i.e. having the pattern and values that conform to the model fit stored in the supplied object. If it is NULL, estimateV.asreml is used to estimate the variance matrix for the observations from the variance parameter estimates from the reduced.asreml.obj.
extra.matrix   A matrix of order equal to the number of observations that is to be added to the variance matrix, the latter based on the information in asreml.obj. It is assumed that the sigma-parameterized values of the variance parameter estimates, such as is given in the varcomp component of summary.asreml, have been used in calculating extra.matrix; the values in the vparameters component of G.param and R.param may be either gamma- or sigma-parameterized. The argument extra.matrix can be used in conjunction with ignore.terms as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.
ignore.terms   A character giving terms from either the random or residual models that are to be ignored in that their contributions to the variance is not to be included in the estimated matrix. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!). This can be used in conjunction with estimateV.asreml as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.
fixed.spline.terms  A character vector giving one or more spline terms in the random model that are regarded as fixed and so are to be ignored because they are not regarded as contributing to the variance. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!).
bound.exclusions

A character specifying one or more bound codes that will result in a variance parameter in the random model being excluded from contributing to the variance. If set to NULL then none will be excluded.

tolerance

The value such that eigenvalues less than it are considered to be zero.

update

If TRUE then the arguments R.param and G.param are set to those in the asreml object supplied in object so that the values from the original model are used as starting values. If FALSE then calls are made to asreml in which the only changes from the previous call are (i) the model is fitted to simulated data and (ii) modifications specified via ... are made, except that changes cannot be made to any of the models.

trace

If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

ncores

A numeric specifying the number of cores to use in doing the simulations.

... Other arguments that are passed down to the function asreml. Changes to the models are not allowed. Other changes are dangerous and generally should be avoided.

Value

A list with the following components:

1. REMLRT: the observed REML ratio statistic.
2. p: the bootstrap p-value for the observed test statistic.
3. DF: the calculated difference in DF for the variance parameters in the two models.
4. totalunconverged: the total number of unconverged analyses over the simulations.
5. REMLRT.sim: a numeric containing the values of the ratio statistics for the simulated data. It has an attribute called na.action that can be retrieved using attr(REMLRT.sim, which = "na.action"); it contains a list of the simulation numbers that were abandoned because max.retries failed to converge for both models.
6. nunconverged: the number of unconverged analyses for each bootstap sample, the maximum being max.retries.

Note

A bootstrap sample is generated using a multivariate normal distribution with expected value as specified by means and variance matrix given by V. Each simulated sample is analysed according to the reduced model and, provided this analysis converges, according to the full.model. If one of these analyses fails to converge, it is abandoned and another sample is generated for this simulation. As many as max.retries attempts are made to generate a data set for which both analyses converge. If data set that converges for both analyses is not generated for a simulation, NA is returned for that bootstrap sample. Hence, the maximum number of data sets that will be generated is nboot * max.retries and less than nboot samples will be generated if a data set that converges for both analyses is not obtained within max.retries attempts.

If a bootstrap sample converges for both analyses, the REML ratio test statistic is calculated as $2(\log(REML)_F - \log(REML)_R)$.

The DF is calculated from the information in full.asreml.obj and reduced.asreml.obj. The degrees of freedom are computed as the difference between the two models in the number of variance parameters whose estimates do not have a code for bound specified in bound.exclusions.
If ASReml-R version 4 is being used then the codes specified in `bound.exclusions` are not restricted to a subset of the default codes, but a warning is issued if a code other than these is specified. For ASReml-R version 3, only a subset of the default codes are allowed: F (Fixed), B (Boundary), C (Constrained) and S (Singular).

Author(s)

Chris Brien

See Also

`REMLRT.asreml`, `infoCriteria.asreml`, `testranfix.asrtests`

Examples

```r
## Not run:
bootREMLRT(ICV.max, ICV.red, ncores = 2)
## End(Not run)
```

changeModelOnIC.asrtests

Uses information criteria to decide whether to change an already fitted model.

Description

Uses information criteria to decide whether to change the fitted model stored in the supplied `asrtests.object` according to the specified modifications. The function `changeTerms` is used to change the model. Thus, the model can be modified using a combination of adding and removing sets of terms from one or both of the fixed or random models, replacing the residual model and changing the bounds and/or initial values of some terms. The model will be unchanged if terms specified in `dropFixed` or `dropRandom` are not in the fitted model.

A row is added to the `test.summary` data.frame of the `asrtests.object` using the supplied `label` and stating whether or not the new model has been swapped for the supplied model. Convergence in fitting the model is checked and a note included in the `action` if there was not. All components of the `asrtests.object` are updated to exhibit the differences between the supplied and new model.

To obtain a list of the information criteria for a set of models use `changeTerms.asrtests` with ICLikelihood set to REML or full, or use `infoCriteria.asreml`.

Usage

```r
## S3 method for class 'asrtests'
changeModelOnIC(asrtests.obj, 
   dropFixed = NULL, addFixed = NULL, 
   dropRandom = NULL, addRandom = NULL, 
   newResidual = NULL, label = "Changed terms", 
   allow.unconverged = TRUE, allow.fixedcorrelation = TRUE, 
   checkboundaryonly = FALSE, 
   trace = FALSE, update = TRUE, denDF = "numeric",
```
arguments

Arguments

**asrtests.obj**
An **asreml.object** containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

**dropFixed**
A single character string in the form of a formula which, after addition of ". ~ . -" and expansion, specifies the sum of a set of terms to be dropped from the fixed formula. The names must match those in the wald.tab component of the asrtests.obj. The fixed terms will be reordered so that single-variable terms come first, followed by two-variable terms and so on. Note that multiple terms specified using a single `asreml::at` function can only be dropped as a whole. If the term was specified using an `asreml::at` function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the `levels` function can be specified.

**addFixed**
A single character string in the form of a formula which, after addition of ". +" and expansion, specifies the sum of a set of terms to be added to the fixed formula. The fixed terms will be reordered so that single-variable terms come first, followed by two-variable terms and so on.

**dropRandom**
A single character string in the form of a formula which, after addition of ". ~ . -" and expansion, specifies the sum of a set of terms to be dropped from the random formula. The names must match those in the vparameters component of the asreml.obj component in the asrtests.obj. Note that multiple terms specified using a single `asreml::at` function can only be dropped as a whole. If the term was specified using an `asreml::at` function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the `levels` function can be specified.

**addRandom**
A single character string in the form of a formula which, after addition of ". +" and expansion, specifies the sum of a set of terms to be added to the random formula.

**newResidual**
A single character string in the form of a formula which, after addition of ". ~ ", specifies the residual (or `rcov`) model. To remove the model, enter "-.(.)".

**label**
A character to use as an entry in the `terms` column in `test.summary` to indicate as far as is possible the terms that are being manipulated.

**allow.unconverged**
A logical indicating whether to accept a new model even when it does not converge. If `FALSE` and the fit of the new model does not converge, the supplied `asrtests.obj` is returned. Also, if `FALSE` and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

**allow.fixedcorrelation**
A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If `FALSE` and the new model contains correlations whose values have not been able to be estimated, the supplied `asrtests.obj` is returned. The fit in the `asreml.obj` component of the supplied `asrtests.obj` will also
be tested and a warning issued if both fixed correlations are found in it and allow.fixed.correlation is FALSE.

checkboundaryonly
If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead.

trace
If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update
If TRUE then update.asreml is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified as specified and (ii) modifications specified via ... are made.

denDF
Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

set.terms
A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting the new model. The names must match those in the vparameters component of the asreml.obj component in the new asrtests.object. The terms in the model do not need to change from those in the model in the supplied asrtests.obj.

ignore.suffices
A logical vector specifying whether the suffices of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in terms.

bounds
A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

which.IC
A character specifying the information criterion to be used in selecting the best model. Possible values are AIC and BIC. The values of the criterion for supplied model must exceed that for changed model for the changed model to be returned.

IClikelihood
A character specifying whether Restricted Maximum Likelihood (REML) or the full likelihood (full) are to be used in calculating the information criteria for family set to asr_gaussian. For family set to asr_binomial or asr_poisson and with dispersion set to 1, the deviance is extracted from object and used to calculate the AIC and BIC (as suggested by Damian Collins); the setting of IClikelihood is ignored and the log-likelihood set to NA. The information criteria are not valid for other settings of family and dispersion.
fixedDF  A numeric giving the number of estimated fixed parameters. If NULL then this is determined from the information in asreml.obj.

varDF  A numeric giving the number of estimated variance parameters. If NULL then this is determined from the information in asreml.obj. It replaces the varDF argument.

initial.values  A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

bound.exclusions  A character specifying the bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in calculating the information criteria. If set to NULL then none will be excluded.

...  Further arguments passed to asreml, wald.asreml and as.asrtests.

Value

An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary. The values of the degrees of freedom and the information criteria are differences between those of the changed model and those of the model supplied to changeModelOnIC.

Author(s)

Chris Brien

See Also

as.asrtests, rmboundary.asrtests, testranfix.asrtests, testresidual.asrtests, newfit.asreml, reparamSigDevn.asrtests, chooseModel.asrtests, changeTerms.asrtests, infoCriteria.asreml

Examples

## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety, 
  random = ~ Row + Column + units, 
  residual = ~ ar1(Row):ar1(Column), 
  data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL, 
  label = "Maximal model")
current.asrt <- rmboundary(current.asrt)

# Drop both Row and Column terms

current.asrt <- changeModelOnIC(current.asr, 
  dropRandom = "Row + Column", 
  checkboundaryonly = TRUE, 
  which.IC = "AIC", ICllikelihood = "full")

current.asrt <- iterate(current.asrt)

# Add and drop both fixed and random terms

current.asrt <- changeModelOnIC(current.asr, 
  addFixed = "vRow", dropFixed = "WithinColPairs",
  ...
changeTerms.asrtests

Add and drops terms from one or both of the fixed or random model, replaces the residual (rcov) model with a new model and changes bounds or initial values of terms.

Description

The specified terms are simply added or dropped, without testing, from either the fixed or random model and/or the residual (rcov) model replaced. Also, the bounds and/or initial values of some terms can be changed. No hypothesis testing is performed, but a check is made for boundary or singular terms.

A row is added to the test.summary data.frame of the asrtests.object using the supplied label and stating which models have been changed. Information criteria can be included in the row of the test.summary. Convergence in fitting the model is checked and a note included in the action if there was not. All components of the asrtests.object are updated.

To only change the terms based on a comparison of information criteria use changeModelOnIC.asrtests.

Usage

## S3 method for class 'asrtests'
changeTerms(asrtests.obj,
dropFixed = NULL, addFixed = NULL,
dropRandom = NULL, addRandom = NULL,
newResidual = NULL, label = "Changed terms",
allow.unconverged = TRUE, allow.fixedcorrelation = TRUE,
checkboundaryonly = FALSE,
trace = FALSE, update = TRUE, denDF = "numeric",
set.terms = NULL, ignore.suffices = TRUE,
bounds = "P", initial.values = NA,
IClikelihood = "none", bound.exclusions = c("F","B","S","C"),
...)

Arguments

asrtests.obj  An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.
dropFixed  A single character string in the form of a formula which, after addition of ". ~ . ~ -" and after expansion, specifies the sum of a set of terms to be dropped from the fixed formula. The names must match those in the wald.tab component of the asrtests.obj. The fixed terms will be reordered so that single-variable terms come first, followed by two-variable terms and so on. Note that multiple terms specified using a single asreml:::at function can only be dropped as a whole. If the term was specified using an asreml:::at function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the levels function can be specified.

addFixed  A single character string in the form of a formula which, after addition of ". ~ . ~ +" and expansion, specifies the sum of a set of terms to be added to the fixed formula. The fixed terms will be reordered so that single-variable terms come first, followed by two-variable terms and so on.

dropRandom  A single character string in the form of a formula which, after addition of ". ~ . ~ -" and expansion, specifies the sum of a set of terms to be dropped from the random formula. The terms must match those in the vparameters component of the asreml.obj component in the asrtests.obj. Note that multiple terms specified using a single asreml:::at function can only be dropped as a whole. If the term was specified using an asreml:::at function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the levels function can be specified.

addRandom  A single character string in the form of a formula which, after addition of ". ~ . ~ +" and expansion, specifies the sum of a set of terms to be added to the random formula.

newResidual  A single character string in the form of a formula which, after addition of ", ~ . ~ -", specifies the residual (or r cov) model. To remove the model, enter "-(.)".

label  A character to use as an entry in the terms column in test.summary to indicate as far as is possible the terms that are being manipulated.

allow.unconverged  A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit does not converge, the supplied asrtests.obj is returned.

allow.fixedcorrelation  A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly  If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead.

trace  If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update  If TRUE then update.asreml is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified as specified and (ii) modifications specified via ... are made.
denDF  Specifies the method to use in computing approximate denominator degrees of freedom when \texttt{wald.asreml} is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

set.terms  A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting the new model. The names must match those in the \texttt{vparameters} component of the \texttt{asreml.obj} component in the new \texttt{asrtests.object}. The terms in the model do not need to change from those in the model in the supplied \texttt{asrtests.obj}.

ignore.suffixes  A logical vector specifying whether the suffixes of the \texttt{asreml}-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffixes are stripped from the \texttt{asreml}-assigned names. If FALSE for an element of terms, the element must exactly match an \texttt{asreml}-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the \texttt{asreml}-assigned suffixes for all the terms in terms.

bounds  A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

initial.values  A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

IClikelihood  A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new \texttt{asrtests.object}. If none, none are included. Otherwise, if REML and family is set to asr_gaussian (the default), then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full and family is set to asr_gaussian, then the AIC and BIC based on the full likelihood are included. if family is asr_binomial or asr_poisson, with dispersion set to 1, the deviance is extracted from object and used to calculate the AIC and BIC. (See also \texttt{infoCriteria.asreml}).

bound.exclusions  A character specifying the bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in calculating the information criteria. If set to \texttt{NULL} then none will be excluded.

...  Further arguments passed to \texttt{asreml}, \texttt{wald.asreml} and \texttt{as.asrtests}.

Value

An \texttt{asrtests.object} containing the components (i) \texttt{asreml.obj}, (ii) \texttt{wald.tab}, and (iii) \texttt{test.summary}.

Author(s)

Chris Brien
ChickpeaEnd.dat

A large data set comprising the end of imaging data from a chick pea experiment conducted in high-throughput greenhouses

Description

The data collected after imaging had been completed on the 1056 plants in the experiment reported by Atieno et al. (2017). The design employed for the experiment was a split-plot design in which two consecutive carts formed a main plot. The split-plot design assigned 245 genotypes to main plots, the genotypes being unequally replicated 2 or 3 times. Treatments (non-saline, saline) were randomized to the two subplots (carts) within each main plot.

The columns in the data.frame are: Smarthouse, Lane, Position, Zone, Mainplot, Subplot, Replicate, xLane, xPosition, Genotypes, Treatments, Biomass, PlantHeight, SenescenceRank, TotalPods, FilledPods, EmptyPods, SeedNo, TotalSeedWt, SeedWt100.

References


See Also

as.asrtests, rmboundary.asrtests, testranfix.asrtests, testresidual.asrtests, newfit.asreml, reparamSigDevn.asrtests, chooseModel.asrtests, changeModelOnIC.asrtests, infoCriteria.asreml

Examples

```r
## Not run:
terms <- "(Date/(Sources * (Type + Species)))"
current.asrt <- changeTerms(current.asrt, addFixed = terms)

current.asrt <- changeTerms(current.asrt, dropFixed = "A + B", denDF = "algebraic")
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                      random = ~ Row + Column + units,
                      residual = ~ ar1(Row):ar1(Column),
                      data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary(current.asrt)
# Add and drop both fixed and random terms
current.asrt <- changeTerms(current.asrt,
                             addFixed = "vRow", dropFixed = "WithinColPairs",
                             addRandom = "spl(vRow)", dropRandom = "units",
                             checkboundaryonly = TRUE)
# Replace residual with model without Row autocorrelation
current.asrt <- changeTerms(current.asrt,
                            newResidual = "Row:ar1(Column)",
                            label="Row autocorrelation")

## End(Not run)
```
The columns Smarthouse, Lane and Position uniquely identify the rows of observations. Zones are groups of 4 Lanes, Mainplots are the 44 pairs of consecutive Subplots within each Zone, and a Subplot is a cart containing a single plant. The columns xLane and xPosition are numeric covariates for location within a Smarthouse. Genotypes and Treatments indicate the genotype and treatment that each plant was allocated. The response variables are Biomass, PlantHeight, SenescenceRank, TotalPods, FilledPods, EmptyPods, SeedNo, TotalSeedWt and SeedWt100.

Usage

\texttt{data(ChickpeaEnd.dat)}

Format

A \texttt{data.frame} with 1056 rows by 20 columns.

References


\begin{verbatim}
chooseModel

\textit{chooseModel} \textit{Determines the set of significant terms using p-values and records the tests performed in a data.frame, taking into account the marginality relations of terms.}

Description

Using p-values from hypothesis tests, determines the set of significant terms, taking into account the hierarchy or marginality of terms. In particular, a term will not be tested if it is marginal to (or nested in) one that is significant. For example, if A:B is significant, then neither A nor B will be tested. The tests conducted in choosing selected model are listed in a summary \texttt{data.frame}.

Usage

\texttt{chooseModel(object, \ldots)}

Arguments

\texttt{object} \hspace{1cm} an object using which p-values can be obtained for use in model selection.

\texttt{\ldots} \hspace{1cm} further arguments passed to or from other methods.

Details

\texttt{chooseModel} is the generic function for the \texttt{chooseModel} method. Use \texttt{methods("chooseModel")} to get all the methods for the \texttt{chooseModel} generic.

\texttt{chooseModel.asrtests} is a method for an \texttt{asrtests.object}. It uses \texttt{testranfix.asrtests} to conduct tests to determine the p-values used in the model selection.

\texttt{chooseModel.data.frame} is a method for a \texttt{data.frame}. It uses the p-values stored in the \texttt{data.frame} in the model selection.
chooseModel.asrtests

Author(s)

Chris Brien

See Also

chooseModel.asrtests, chooseModel.asrtests, changeModelOnIC.asrtests, testranfix.asrtests

chooseModel.asrtests  Determines and records the set of significant terms using an asrtests.object, taking into account the hierarchy or marginality relations of the terms.

Description

Performs a series of hypothesis tests on a set of fixed and/or random terms taking into account the marginality of terms. In particular, a term will not be tested if it is marginal to (or nested in) one that is significant. For example, if A:B is significant, then neither A nor B will be tested. For a random term, the term is removed from the model fit, any boundary terms are removed using rmboundary.asrtests and a REML likelihood ratio test is performed using REMLRT.asreml. If it is not significant and drop.ran.ns is TRUE, the term is permanently removed from the model. Note that if boundary terms are removed, the reduced model may not be nested in the full model in which case the test is not valid. For fixed terms, the Wald tests are performed and the p-value for the term obtained. If it is not significant and drop.fix.ns is TRUE, the term is permanently removed from the model. A row that records the outcome of a test is added to test.summary for each term that is tested.

Usage

## S3 method for class 'asrtests'
chooseModel(object, terms.marginality=NULL, alpha = 0.05, allow.unconverged = TRUE, allow.fixedcorrelation = TRUE, checkboundaryonly = FALSE, drop.ran.ns=TRUE, positive.zero = FALSE, bound.test.parameters = "none", drop.fix.ns=FALSE, denDF = "numeric", dDF.na = "none", dDF.values = NULL, trace = FALSE, update = TRUE, set.terms = NULL, ignore.suffices = TRUE, bounds = "P", initial.values = NA, IClikelihood = "none", ...)

Arguments

object  an asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

terms.marginality  A square matrix of ones and zeros with row and column names being the names of the terms to be tested. The names of fixed terms must match those in the wald.tab component of the object, while the names of random terms must match those in the vparameters component of the asreml.obj component in the object. The diagonal elements of the matrix should be one, indicating that a term is marginal to itself. Elements should be one if the row term is marginal to the column term. All other elements should be zero.
chooseModel.asrtests

alpha
The significance level for the test.

allow.unconverged
A logical indicating whether to accept a new model even when it does not converge. If FALSE and a fit when a term is removed does not converge, the term will not be removed.

allow.fixedcorrelation
A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly
If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead.

drop.ran.ns
A logical indicating whether to drop nonsignificant random terms from the model.

positive.zero
Indicates whether the hypothesized values for the variance components being tested are on the boundary of the parameter space. For example, this is true for positively-constrained variance components that, under the reduced model, are zero. This argument does not need to be set if bound.test.parameters is set.

bound.test.parameters
Indicates whether for the variance components being tested, at least some of the hypothesized values are on the boundary of the parameter space. The possibilities are "none", "onlybound" and "one-and-one". The default is "none", although if it is set to "none" and positive.zero is TRUE then bound.test.parameters is taken to be "onlybound". When bound.test.parameters is set to "one-and-one", it signifies that there are two parameters being tested, one of which is bound and the other is not. For example, the latter is true for testing a covariance and a positively-constrained variance component that, under the reduced model, are zero.

drop.fix.ns
A logical indicating whether to drop a fixed term from the model when it is nonsignificant.

denDF
Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

dDF.na
The method to use to obtain substitute denominator degrees of freedom when the numeric or algebraic methods produce an NA. If dDF.na = "none", no substitute denominator degrees of freedom are employed; if dDF.na = "residual", the residual degrees of freedom from asreml.obj$nedf are used; if dDF.na = "maximum", the maximum of those denDF that are available, excluding that for the Intercept, is used; if all denDF are NA, asreml.obj$nedf is used. If dDF.na = "supplied", a vector of values for the denominator degrees of freedom is to be supplied in dDF.values. Any other setting is ignored and a warning message produced. Generally, substituting these degrees of freedom is anticonservative in that it is likely that the degrees of freedom used will be too large.
dDF.values  A vector of values to be used when dDF.na = "supplied". Its values will be used when denDF in a test for a fixed effect is NA. This vector must be the same length as the number of fixed terms, including (Intercept) whose value could be NA.

trace  If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update  If TRUE then update.asreml is called in testing models. In doing this the arguments R.param and G.param are set to those in the asreml object stored in object so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes to the asreml.obj stored in the supplied object are (i) to the terms in the fixed and random models corresponding to terms in terms.marginality and (ii) those modifications specified via ....

set.terms  A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the vparameters component of the asreml.obj component in the new asrtests.object.

ignore.suffices  A logical vector specifying whether the suffices of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in terms.

bounds  A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

initial.values  A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

IClikelihood  A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new asrtests.object. If none, none are included. Otherwise, if REML, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full, then the AIC and BIC based on the full likelihood are included. (See also infoCriteria.asreml.)

... further arguments passed to asreml, wald.asreml and as.asrtests via testranfix.asrtests.

Value

A list containing:

1. asrtests.obj: an asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary;

2. sig.tests: a character vector whose elements are the the significant terms amongst those tested.
chooseModel.data.frame

Determines the set of significant terms from results stored in a data.frame, taking into account the marginality relations of terms and recording the tests used in a data.frame.

Description

Uses the p.values from a set of hypothesis tests that are stored in the supplied data.frame to choose a model to describe the effects of the terms corresponding to the p-values, taking into account the hierarchy or marginality of terms. In particular, a term will not be tested if it is marginal to (or nested in) one that is significant. For example, if A:B is significant, then neither A nor B will be tested. The tests used in choosing the selected model are listed in the data.frame choose.summary. No change is made to the p.values, the DF and denDF being for information only.
Usage

## S3 method for class 'data.frame'
chooseModel(object, terms=NULL, p.values = "Pr",
    DF = "Df", denDF = "denDF", omit.DF = FALSE,
    terms.marginality=NULL, alpha = 0.05, ...)

Arguments

object

A character giving the name of the column in object containing the terms

terms

A character giving the name of the column in object containing the terms

p.values

A character giving the name of the column in object containing the p-values

terms.marginality

A square matrix of ones and zeros with row and column names being the names

Value

A list containing:
1. `choose.summary`: a `data.frame` summarizing the tests carried out in choosing the significant terms; provided `omit.DF = FALSE`, it has the same columns as a `test.summary` from an `asrtests.object`.

2. `sig.tests`: a character vector whose elements are the significant terms amongst those tested.

**Author(s)**

Chris Brien

**See Also**

chooseModel, chooseModel.asrtests

**Examples**

```r
data("Ladybird.dat")

## Use asreml to get the table of p-values
## Not run:
m1.asr <- asreml(logitP ~ Host*Cadavers*Ladybird,
random = ~ Run,
data = Ladybird.dat)
current.asrt <- as.asrtests(m1.asr)
fixed.tab <- current.asrt$wald.tab
col.p <- "Pr"
df = "Df"
den.df = "denDF"

## End(Not run)

## Use lmeTest to get the table of p-values
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE)) {
m1.lmer <- lmerTest::lmer(logitP ~ Host*Cadavers*Ladybird + (1|Run),
data=Ladybird.dat)
fixed.tab <- anova(m1.lmer, type = "II")
col.p <- "Pr(>F)"
df = "NumDF"
den.df = "DenDF"
}

## Select a model using the table of p-values obtained with either asreml or lmerTest
if (exists("fixed.tab")) {
term.marg <- dae::marginality(dae::pstructure(~ Host*Cadavers*Ladybird,
data = Ladybird.dat))
chosen <- chooseModel(fixed.tab, p.values = col.p, DF = df, denDF = den.df,
terms.marginality = term.marg)
}
```
chooseSpatialModelOnIC.asrtests

Uses information criteria to choose the best fitting spatial model for accounting for local spatial variation.

Description

For a response variable measured on a potentially irregular grid of rows and columns of the units, uses information criteria to decide whether to add to the fitted model stored in the supplied `asrtests.object` either a two-dimensional exponential correlation model, a two-dimensional tensor-product natural cubic smoothing spline model (TPNCS), a two-dimensional tensor-product penalized P-spline model (TPPCS) model, or a two-dimensional tensor-product penalized linear spline model with first-difference penalties (TPP1LS) to account for the local spatial variation. The models from which to select can be reduced to a subset of these four models. The data can be arranged in sections, for each of which there is a grid and for which the model is to be fitted separately. Also, the rows and columns of a grid are not necessarily one observational unit wide. The spatial model is only added if the information criterion of the supplied model is decreased with the addition of the local spatial model.

One or more rows is added to the test.summary data.frame of the `asrtests.object`, for each section and each spatial model, stating whether or not the new model has been swapped for a model in which the spatial model has been added to the supplied model. Convergence in fitting the model is checked and a note included in the action if there was not. All components of the `asrtests.object` are updated to exhibit the differences between the supplied and any new model.

Usage

```r
## S3 method for class 'asrtests'
chooseSpatialModelOnIC(asrtests.obj, trySpatial = "all",
sections = NULL,
row.covar = "cRow", col.covar = "cCol",
row.factor = "Row", col.factor = "Col",
corr.funcs = c("ar1", "ar1"),
row.corrfitfirst = TRUE,
dropRowterm = NULL, dropColterm = NULL,
nsegs = NULL, nestorder = c(1,1),
asreml.option = "mbf",
tpps4mbf.obj = NULL,
allow.unconverged = FALSE,
allow.fixedcorrelation = FALSE,
checkboundaryonly = FALSE, update = FALSE,
IClikelihood = "full", which.IC = "AIC",
return.asrts = "best", ...)
```

Arguments

- `asrtests.obj` An `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary`.
- `trySpatial` A character string nominating the types of spatial model whose fits are to be assessed. Possible values are none, corr, TPNCS, TPPCS, and TPP1LS. If set to none, then just the supplied nonspatial model and the information about its information criteria will be returned.
sections A single character string that specifies the name of the column in the data.frame that contains the factor that identifies different sections of the data to which separate spatial models are to be fitted.

row.covar A single character string nominating a numeric that contains the values of a centred covariate indexing the rows of a grid. The numeric must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

col.covar A single character string nominating a numeric that contains the values of a centred covariate indexing the columns of a grid. The numeric must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

row.factor A single character string nominating a factor that indexes the rows of a grid that are to be one dimension of a spatial correlation model. The factor must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

col.factor A single character string nominating a factor that indexes the columns of a grid that are to be one dimension of a spatial correlation model. The factor must be a column in the data.frame stored in the asreml.obj that is a component of the supplied asrtests.obj.

corr.funcs A single character string of length two that specifies the asreml one-dimensional correlation or variance model function for the row and column dimensions of a two-dimensional separable spatial correlation model; the two-dimensional model is fitted as a random term. If a correlation or variance model is not to be investigated for one of the dimensions, specify "" for that dimension.

row.corrFitfirst If TRUE then the row correlation or variance function is fitted first, followed by the addition of the column correlation or variance function. If FALSE, the order of fitting is reversed.

dropRowterm A single character string nominating a factor in the data.frame that has as many levels as there are unique values in row.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropRowterm and a random row deviations term based on row.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

dropColterm A single character string nominating a factor in the data.frame that has as many levels as there are unique values in col.covar. This argument is required for spatial.model set to TPNCSS or TPPS. It is used to remove a term corresponding to the dropColterm and a random column deviations term based on col.covar will be included in the model. If the argument is NULL, it is assumed that such a term is not included in the fitted model stored in asrtests.obj.

nsegs A pair of numeric values giving the number of segments into which the column and row ranges are to be split, respectively (each value specifies the number of internal knots + 1). If only one number is specified, that value is used in both dimensions. If not specified, then (number of unique values - 1) is used in each dimension; for a grid layout with equal spacing, this gives a knot at each data value.

nestorder A character of length 2. The order of nesting for column and row dimensions, respectively; default=1 (no nesting). A value of 2 generates a spline with half the number of segments in that dimension, etc. The number of segments in each direction must be a multiple of the order of nesting.
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asreml.option A single character string specifying whether the grp or mbf methods are to be used to supply externally formed covariate matrices to asreml. If the mbf methods is to be used, then makeTPPSplineMats.data.frame must be used before calling addSpatialModelOnIC.asrtests. Compared to the mbf method, the grp method creates large asreml objects, but is faster. The grp method adds columns to the data.frame containing the data; the mbf method adds only fixed covariate to data and stores the random covariates externally.

tpps4mbf.obj An object made with makeTPPSplineMats.data.frame and which contains the spline basis information, that is extra to the data.frames created by makeTPPSplineMats.data.frame in the environment in which it is called and that is needed to fit a TPPS model using the mbf method of asreml.

allow.unconverged A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit of the new model does not converge, the supplied asrtests.obj is returned. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

allow.fixedcorrelation A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead. Note that, for correlation models, the fitting of each dimension and the test for a nugget term are performed with checkboundaryonly set to TRUE and its supplied setting only honoured using a call to rmboundary.asrtests immediately prior to returning the final result of the fitting.

update If TRUE then update.asreml is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified as specified and (ii) modifications specified via ... are made.

which.IC A character specifying the information criterion to be used in selecting the best model. Possible values are AIC and BIC. The values of the criterion for supplied model must exceed that for changed model for the changed model to be returned.

IClikelihood A character specifying whether Restricted Maximum Likelihood (REML) or the full likelihood (full) are to be used in calculating the information criteria.

return.asrts A character string specifying whether the asrtests.object for the best fitting model (smallest AIC or BIC) is returned or the asrtests.objects resulting from the attempted fits of all of the models specified using trySpatial are returned.

... Further arguments passed to changeModelOnIC.asrtests, asreml and tpsmmb.

Details

A fitted spatial model is only returned if it improves the fit over an above that achieved with the model fit supplied in the asrtests.obj. If return.asrts is all, then this applies to each spatial
model specified by `trySpatial`. The model fit supplied in the `asrtests.obj` should not include terms that will be included in any local spatial model. All spatial model terms are fitted as fixed or random. Consequently, the residual model does not have to be iid. The improvement in the fit resulting from the addition of a spatial model to the supplied model is evaluated. Note that the data must be in the order that corresponds to the residual argument with a variable to the right of another variable changes levels in the data frame faster than those of the other variable e.g. `Row:Column` implies that all levels for `Column` in consecutive rows of the data frame with a single `Row` level.

For the `corr` spatial model, the default model is an autocorrelation model of order one (`ar1`) for each dimension. However, any of the single dimension correlation/variance models from `asreml` can be specified for each dimension, as can no correlation model for a dimension; the models for the two dimensions can differ. Using a forward selection procedure, a series of models are tried, without removing boundary or singular terms, beginning with the addition of row correlation and followed by the addition of column correlation or, if the `row.corrfitfirst` is set to `FALSE`, the reverse order. If the fitting of the first-fitted correlation did not result in a model change because the fitting did not converge or correlations were fixed, but the fit of the second correlation was successful, then adding the first correlation will be retried. If one of the metric correlation functions is specified (e.g. `exp`), then the `row.covar` or `col.covar` will be used in the spatial model. However, because the correlations are fitted separately for the two dimensions, the `row.factor` and `col.factor` are needed for all models and is used for a dimension that does not involve a correlation/variance function for the fit being performed. Also, the correlation models are fitted as random terms and so the correlation model will include a variance parameter for the grid even when `ar1` is used to specify the correlation model, i.e. the model fitted is a variance model and there is no difference between `ar1` and `ar1v` in fitting the model. The variance parameter for this term represents the spatial variance and the fit necessarily includes a nugget term, this being the residual variance. If any correlation is retained, the need for a nugget term is assessed by fixing the residual variance to one, which will have no effect if heterogeneous residual variances have been specified. Once the fitting of the correlation model has been completed, the `rmboundary` function will be executed with the checkboundaryonly value supplied in the `chooseSpatialModelOnIC.asrtests` call.

The tensor-product natural-cubic-smoothing-spline `TPNCSS` spatial model is as described by Verbyla et al. (2018), the tensor-product penalized-cubic-spline `TPPCS` model is as described by Rodriguez-Alvarez et al. (2018), and the tensor-product, first-difference-penalty, linear spline `TPP1LS` model that is amongst those described by Piepho, Boer and Williams (2022). The fixed terms for these models are `row.covar + col.covar + row.covar:col.covar` and the random terms `spl(row.covar) + spl(col.covar) + dev(row.covar) + dev(col.covar) + spl(row.covar):col.covar + row.covar:spl(col.covar) + spl(row.covar):spl(col.covar)`. The supplied model should not include any of these terms. However, any fixed or random main-effect term for either `dropRowterm` or `dropColterm` will be removed from the fit.

The `TPPCS` and `TPP1LS` models are fitted using functions from the R package `TPSbits` authored by Sue Welham (2022). There are two methods for supplying the spline basis information produced by `tpsmb` to `asreml`. The `grp` method adds the it to the `data.frame` holding the information for the analysis. The `mbf` method requires the spline basis information to be in the same environment as the function that is called to make a fit using `asreml`. To this end, and prior to invoking the calling function, `makeTPPSplineMats.data.frame` must be used produce the `data.frames`.

All models utilize the function `changeModelOnIC.asrtests` to assess the model fit, the information criteria used in assessing the fit being calculated using `infoCriteria`. Arguments from `tpsmb` and `changeModelOnIC.asrtests` can be supplied in calls to `chooseSpatialModelOnIC.asrtests` and will be passed on to the relevant function throught the `ellipses` argument (...).

The data for experiment can be divided into sections and an attempt to fit the same spatial model to each is made. The fit may differ for each of the sections, but the fit over all of the sections is assessed.

Each combination of a `row.coords` and a `col.coords` does not have to specify a single observation;
for example, to fit a local spatial model to the main units of a split-unit design, each combination would correspond to a main unit and all subunits of the main unit would have the same combination.

**Value**

A list containing four components: (i) asrts, (ii) spatial.IC, (iii) best.spatial.mod, and (iv) best.spatial.IC.

The component asrts itself holds a list of one or more asrtests.objects, either the best overall out of the supplied model and the spatial models, or, for each spatial model, the best out of the supplied model and that spatial model. Each asrtests.object contains the components: (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

The spatial.IC component holds a data.frame with summary of the values of the information criteria for the supplied model and those resulting from adding the spatial model to the supplied model. In the case of a spatial correlation model, the information criteria for the selected spatial correlation model is returned. If a spatial model could not be fitted, then all returned values will be NA.

The best.spatial component is a character giving the name of the best spatial model, and best.spatial.AIC gives the value of its AIC.

**Author(s)**

Chris Brien

**References**


Welham, S. J. (2022) TPSbits: Creates Structures to Enable Fitting and Examination of 2D Tensor-Product Splines using ASReml-R. Version 1.0.0 [https://mmade.org/tpsbits/](https://mmade.org/tpsbits/)

**See Also**

as.asrtests, rmboundary.asrtests, addSpatialModelOnIC.asrtests, addSpatialModel.asrtests, testranfix.asrtests, testresidual.asrtests, newfit.asreml, reparamSigDevn.asrtests, chooseModel.asrtests, changeModelOnIC.asrtests, infoCriteria.asreml

**Examples**

```r
## Not run:

data(Wheat.dat)

#Add row and column covariates
Wheat.dat <- within(Wheat.dat,
```
estimateV.asreml

Forms the estimated variance, random or residual matrix for the observations from the variance parameter estimates.

Description

Forms the estimated variance (V), random (G) or (R) matrix for the observations, a square symmetric matrix of order equal to the number of observations. The estimates of the variance parameters and the information about the random and residual models for which they were estimated are obtained from the asreml object. This function is not available in ASReml-R version 3.

Usage

## S3 method for class 'asreml'
estimateV(asreml.obj, which.matrix = "V", extra.matrix = NULL, ignore.terms = NULL, fixed.spline.terms = NULL, bound.exclusions = c("F","B","S","C"), ...)

Arguments

asreml.obj An asreml object from a call to asreml in which the data argument has been set.

which.matrix A character giving the matrix that is to be formed. It must be one of "V", to produce the variance matrix \( V = G + R \), "G" to produce the matrix \( G \), corresponding to the random formula, or "R" to produce the matrix \( R \), corresponding to the residual formula.
extra.matrix  A matrix of order equal to the number of observations that is to be added to the matrix specified by which.matrix, the latter based on the information in asreml.obj. It is assumed that the sigma-parameterized values of the variance parameter estimates, such as is given in the varcomp component of summary.asreml, have been used in calculating extra.matrix; the values in the vparameters component of G.param and R.param may be either gamma- or sigma-parameterized. The argument extra.matrix can be used in conjunction with ignore.terms as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.asreml.

ignore.terms  A character giving terms from either the random or residual models that are to be ignored in that their contributions to the variance is not to be included in the estimated matrix. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!). This can be used in conjunction with estimateV.asreml as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.asreml.

fixed.spline.terms  A character vector giving one or more spline terms in the random model that are regarded as fixed and so are to be ignored because they are not regarded as contributing to the variance. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!).

bound.exclusions  A character specifying one or more bound codes that will result in a variance parameter in the random model being excluded from contributing to the variance. If set to NULL then none will be excluded.

Details

The information about the variance parameters in the fitted mixed model are obtained from the G.param and R.param components of the asreml object. The function can deal with the following variance functions in either the random or residual models: id, diag, us, ar1, ar2, ar3, sar,sar2, ma1, ma2, arma, exp, gau, cor, corb and corg. All of these functions, except us, can be combined with either v or h. It will also cope with the following functions in the random model: at, str, spl, dev, grp, fa and rr. Additionally, it can deal with the function dsum in the residual model. For further information see the ASReml-R User Guide Version 4 (Butler et al., 2018).

Value

A matrix containing the estimated variance matrix.

Author(s)

Chris Brien

References

exploreLSDs.alldiffs

See Also

asreml, simulate.asreml, variofaces.asreml.

Examples

## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
    random = ~ Row + Column + units,
    residual = ~ ar1(Row):ar1(Column),
data=Wheat.dat)
# Form variance matrix based on estimated variance parameters
V <- estimateV(current.asr)

## End(Not run)

exploreLSDs.alldiffs   Explores the computed LSD values for pairwise differences between predictions.

Description

Given an alldiffs.object with an sed component, the LSDs are calculated for all pairwise comparisons of predictions. It then calculates (i) a table of frequencies of the LSD values, (ii) the distinct values of the LSDs after rounding, (iii) various statistics from the LSD values, (iv) a measure of the accuracy of each of the LSD statistics, (v) the numbers of false positives and false negatives for each of the LSD statistics if pairwise comparisons are based on the LSD statistic, (vi) the accuracy of each statistic in representing the LSD values for each prediction and (vii) a matrix containing the LSD values for comparing each pair of predictions. Histograms of the frequencies can also be produced.

Usage

## S3 method for class 'alldiffs'
exploreLSDs(alldiffs.obj, LSDtype = "overall", LSDby = NULL,
    LSDaccuracy = "maxAbsDeviation", alpha = 0.05, digits = 3,
    retain.zeroLSDs = FALSE,
    zero.tolerance = .Machine$double.eps ^ 0.5,
    plotHistogram = FALSE, ...)

Arguments

alldiffs.obj  An alldiffs.object.
LSDtype  A character string that can be overall or factor.combinations. It determines whether the LSD values that are investigated and stored are (i) the overall minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of all pairwise LSDs, or (ii) the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum for the pairwise LSDs for each factor.combination, unless there is only one prediction for a factor.combination, when notional LSDs are calculated. The LSDtype specified here does not have to match that used in the creating the alldiffs.object.
exploreLSDs.alldiffs

See `LSD.frame` for further information on how the LSD statistics are calculated.

**LSDby**
A **character** (vector) of variables names, being the names of the **factors** or **numerals** in the `classify`; for each combination of the values the of the **factors** and **numerals**, the LSD statistics and accuracy are computed, as well histograms plotted, when `LSDtype` is `factor.combinations`. The `LSDby` specified here does not have to match that used in the creating the `alldiffs.object`.

**LSDaccuracy**
A **character** nominating one of `maxAbsDeviation`, `maxDeviation`, `q90Deviation` or `RootMeanSqDeviation` as the statistic to be calculated as a measure of the accuracy of an LSD statistic when its values are used as an approximate LSD. The option `q90Deviation` produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between a set of LSDs and an LSD statistic calculated from those LSDs; the accuracy is expressed as a proportion of the value of the LSD statistic.

**alpha**
A **numeric** specifying the significance level for an LSD to compare a pair of predictions.

**digits**
A **numeric** specifying the number of significant digits to retain in rounding the LSDs before determining the distinct rounded LSDs.

**retain.zeroLSDs**
A **logical** indicating whether to retain or omit LSDs that are zero when calculating the summaries of LSDs.

**zero.tolerance**
A **numeric** specifying the value such that if an LSD is less than it, it will be considered to be zero.

**plotHistogram**
A **logical** indicating whether or not histograms of the LSD values are to be plotted. The `LSDtype` argument controls whether one histogram of all LSD values is plotted or histograms are plotted for each combination of the levels of the factors specified by the `LSDby` argument.

... Provision for passing arguments to functions called internally - not used at present.

**Details**

The false positives and negatives are computed by comparing, for each pair of predictions within each levels-combination of the `LSDby` variables, the significance of the pair difference determined using (i) the true LSD that is computed from the standard error of differences for the pair and (ii) the approximate LSD that is a statistic computed from the true LSDs for all pairwise difference within each levels-combination of the `LSDby` variables. The number of false positives is the number of pairwise differences for which a difference is declared significant using the approximate LSD, but not using the true LSD. The number of false negatives is the number of pairwise differences for which a difference is declared nonsignificant using the approximate LSD, but significant using the true LSD.

The LSD accuracy for a set of LSDs is a function of the deviations of those LSDs and an LSD statistic calculated from them; the accuracy is expressed as a proportion of the value of the LSD statistic.

**Value**

A `list` with components `frequencies`, `distinct.vals`, `statistics`, `accuracy`, `per.pred.accuracy` and `LSD`:

1. `frequencies` is a `data.frame` with the frequency distribution of the LSD values;
2. `distinct.vals` is a list, each component of which contains the distinct values of the LSDs after rounding;

3. `statistics` is a data.frame with the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of the LSD values;

4. `accuracy` is a data.frame with the accuracies of the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of the LSD values with respect to the values from which these statistics are calculated;

5. `false.pos` is a data.frame with the numbers of false positives for the pairwise comparisons within each levels-combination of the LSDby variables when each of the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of the LSD values is used as an approximate LSD in determining the significance of the pairwise differences;

6. `false.neg` is a data.frame with the numbers of false negatives for the pairwise comparisons within each levels-combination of the LSDby variables when each of the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of the LSD values is used as an approximate LSD in determining the significance of the pairwise differences;

7. `per.pred.accuracy` is a data.frame with the accuracies of the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of the LSD values for a set of predictions when these statistics are used to represent the LSDs for the comparisons amongst the set of predictions;

8. `LSD` is a square matrix containing the LSD values for all pairwise comparisons of the predictions contained in the supplied `alldiffs.obj`.

In the statistics, accuracy, false.pos and false.neg data.frames, `c` is the number of pairwise comparisons on which the values in the same row are based. The accuracy measure is specified by the `LSDaccuracy` argument.

**Author(s)**

Chris Brien

**See Also**

`asremlPlus-package`, `plotLSDs.data.frame`, `plotLSDs.alldiffs`, `plotLSDerrors.alldiffs`, `plotLSDerrors.data.frame`, `recalcLSD.alldiffs`, `pickLSStatistics.alldiffs`, `redoErrorIntervals.alldiffs`

**Examples**

```r
data(WaterRunoff.dat)
##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
  random = ~ Benches:MainPlots,
  keep.order=TRUE, data= WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
T5.diffs <- predictPlus(classify = "Sources:Type",
  asreml.obj = current.asr,
  wald.tab = current.asrt$wald.tab,
  present = c("Sources", "Type", "Species"))
```
facCombine.alldiffs

Combines several factors into one in the components of an alldiffs.object

Description

Combines several factors, in the prediction component of object, into one whose levels are the combinations of the used levels of the individual factors. The matching changes are made to the other components and the attributes of the alldiffs.object. If any of the factors to be combined are in LSDby, they are removed from the LSDby, unless the factors to be combined are exactly those in the LSDby. The levels of the factors are combined using fac.combine from the dae package.

Usage

## S3 method for class 'alldiffs'
facCombine(object, factors, order="standard", combine.levels=TRUE, sep="_", level.length = NA, ...)
Arguments

object
An `alldiffs.object`.
factors
A `character` containing the names of factors in the prediction component of object whose levels are to be combined.
order
Either `standard` or `yates`. The order in which the levels combinations of the factors are to be considered as numbered when forming the levels of the combined factor; standard numbers them as if they are arranged in standard order, that is with the levels of the first factor moving slowest and those of the last factor moving fastest; yates numbers them as if they are arranged in Yates order, that is with the levels of the first factor moving fastest and those of the last factor moving slowest.
combine.levels
A logical specifying whether the levels labels of the new factor are to be combined from those of the factors being combined. The default is to use the integers from 1 to the product of the numbers of combinations of used levels of the individual factors, numbering the levels according to order.
sep
A character string to separate the levels when `combine.levels = TRUE`.
level.length
The maximum number of characters from the levels of factors to use in the row and column labels of the tables of pairwise differences and their p-values and standard errors.

Value
A modified `alldiffs.object`.

Author(s)
Chris Brien

See Also
`as.alldiffs`, `alldifferences.data.frame`, `print.alldiffs`, `sort.alldiffs`, `renewClassify.alldiffs`; `fac.combine` in package `dae`.

Examples

data("Ladybird.dat")

## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(logitP ~ Host*Cadavers*Ladybird,
    random = ~ Run,
    data = Ladybird.dat)
current.asrt <- as.asrtests(m1.asr)
HCL.pred <- asreml::predict.asreml(m1.asr, classify="Host:Cadavers:Ladybird",
    sed=TRUE)
HCL.preds <- HCL.pred$pvals
HCL.sed <- HCL.pred$sed
HCL.vcov <- NULL
wald.tab <- current.asrt$wald.tab
den.df <- wald.tab[match("Host:Cadavers:Ladybird", rownames(wald.tab)), "denDF"]
## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
  ml.lmer <- lmerTest::lmer(logitP ~ Host*Cadavers*Ladybird + (1|Run),
    data=Ladybird.dat)
  HCL.emm <- emmeans::emmeans(ml.lmer, specs = ~ Host:Cadavers:Ladybird)
  HCL.preds <- summary(HCL.emm)
  den.df <- min(HCL.preds$df)
  ## Modify HCL.preds to be compatible with a predictions.frame
  HCL.preds <- as.predictions.frame(HCL.preds, predictions = "emmean",
    se = "SE", interval.type = "CI",
    interval.names = c("lower.CL", "upper.CL"))
  HCL.vcov <- vcov(HCL.emm)
  HCL.sed <- NULL
}

## Use the predictions obtained with either asreml or lmerTest
if (exists("HCL.preds"))
{
  ## Form an all.diffs object
  HCL.diffs <- as.alldiffs(predictions = HCL.preds, classify = "Host:Cadavers:Ladybird",
    sed = HCL.sed, vcov = HCL.vcov, tdf = den.df)
  ## Check the class and validity of the alldiffs object
  is.alldiffs(HCL.diffs)
  validAlldiffs(HCL.diffs)
  ## Combine Cadavers and Ladybird
  HCL.diffs <- facCombine(HCL.diffs, factors = c("Cadavers","Ladybird"))
  ## Check the validity of HCL.diffs
  validAlldiffs(HCL.diffs)
}

---

desc

facRecast.alldiffs

Reorders and/or revises the factor levels using the order of old levels in levels.order and the new labels for the levels given in newlabels. The values in levels.order must be unique.

**Description**

Reorders and revises the levels and labels of a factor, in the prediction component of an alldiffs.object. The values in the levels.order vector should be the same as the levels in the existing factor, but the order can be changed. To revise the levels, specify the new levels in the newlabels vector and these will replace the corresponding value in the levels.order vector. The matching changes are made to the other components and attributes of the alldiffs.object.

**Usage**

## S3 method for class 'alldiffs'
facRecast(object, factor, levels.order = NULL, newlabels = NULL, ...)

---

facRecast.alldiffs
Arguments

- **object**: An `alldiffs.object`.
- **factor**: A `character` containing the name of a `factor` in the prediction component of `object` whose levels and labels are to be recast.
- **levels.order**: A `vector` of length `levels(factor)` containing the old levels in the new order for the factor being created. If `levels.order` is `NULL`, then the current levels of `levels(factor)` are used.
- **newlabels**: A `vector` of length `levels(factor)` containing values to use in the revision.
- **...** Further arguments passed to the `factor` call creating the new `factor`.

Value

A modified `alldiffs.object`.

Author(s)

Chris Brien

See Also

- `as.alldiffs`, `allDifferences.data.frame`, `print.alldiffs`, `sort.alldiffs`, `facCombine.alldiffs`, `facRename.alldiffs`, `renewClassify.alldiffs`; fac.recode in package `dae`.

Examples

data("Ladybird.dat")

## Use asreml to get predictions and associated statistics
## Not run:
m1.asr <- asreml(logitP ~ Host*Cadavers*Ladybird, random = ~ Run, data = Ladybird.dat)
current.asrt <- as.asrtests(m1.asr)
HCL.pred <- asreml::predict.asreml(m1.asr, classify="Host:Cadavers:Ladybird", sed=TRUE)
HCL.preds <- HCL.pred$pvals
HCL.sed <- HCL.pred$sed
HCL.vcov <- NULL
wald.tab <- current.asrt$wald.tab
den.df <- wald.tab[match("Host:Cadavers:Ladybird", rownames(wald.tab)), "denDF"]

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE)) {
m1.lmer <- lmerTest::lmer(logitP ~ Host*Cadavers*Ladybird + (1|Run), data=Ladybird.dat)
HCL.emm <- emmmeans::emmmeans(m1.lmer, specs = ~ Host:Cadavers:Ladybird)
HCL.preds <- summary(HCL.emm)
den.df <- min(HCL.preds$df)
## Modify HCL.preds to be compatible with a predictions.frame

```r
code
HCL.preds <- as.predictions.frame(HCL.preds, predictions = "emmean",
    se = "SE", interval.type = "CI",
    interval.names = c("lower.CL", "upper.CL"))
```

HCL.vcov <- vcov(HCL.emm)
HCL.sed <- NULL

## Use the predictions obtained with either asreml or lmerTest

```r
if (exists("HCL.preds")){

  ## Form an all.diffs object

  HCL.diffs <- allDifferences(predictions = HCL.preds, classify = "Host:Cadavers:Ladybird",
    sed = HCL.sed, vcov = HCL.vcov, tdf = den.df)

  ## Check the class and validity of the alldiffs object

  is.alldiffs(HCL.diffs)
  validAlldiffs(HCL.diffs)

  ## Recast the Ladybird and Host factors

  HCL.diffs <- facRecast(HCL.diffs, factor = "Ladybird",
    newlabels = c("none", "present"))
  HCL.diffs <- facRecast(HCL.diffs, factor = "Ladybird",
    levels.order = c("present", "none"),
    newlabels = c("yes","no"))
  HCL.diffs <- facRecast.alldiffs(HCL.diffs, factor = "Host",
    levels.order = c("trefoil", "bean"))

  ## Check the validity of HCL.diffs

  validAlldiffs(HCL.diffs)
}
```

---

**facRename.alldiffs**  
*Renames factors in the prediction component of an alldiffs.object.*

### Description

Renames factors in the prediction component of an alldiffs.object. These changes are propagated to the other components and attributes of the alldiffs.object.

### Usage

```r
## S3 method for class 'alldiffs'
facRename(object, factor.names, newnames, ...)
```

### Arguments

- **object**  
  An alldiffs.object.

- **factor.names**  
  A character containing the names of the factors in the prediction component of object that are to be renamed.

- **newnames**  
  A character containing the new names of the factors in the prediction component of object.
Provision for passing arguments to functions called internally - not used at present.

Value

A modified `alldiffs.object`.

Author(s)

Chris Brien

See Also

`as.alldiffs, allDifferences.data.frame, print.alldiffs, sort.alldiffs, facCombine.alldiffs, facRecast.alldiffs, renewClassify.alldiffs, fac.recode` in package `dae`.

Examples

data("Ladybird.dat")

## Use asreml to get predictions and associated statistics

## Not run:
ml.asr <- asreml(logitP ~ Host*Cadavers*Ladybird,
    random = ~ Run,
    data = Ladybird.dat)
current.asrt <- as.asrtests(ml.asr)
HCL.pred <- asreml::predict.asreml(ml.asr, classify="Host:Cadavers:Ladybird",
    sed=TRUE)
HCL.preds <- HCL.pred$pvals
HCL.sed <- HCL.pred$sed
HCL.vcov <- NULL
wald.tab <- current.asrt$wald.tab
den.df <- wald.tab[match("Host:Cadavers:Ladybird", rownames(wald.tab)), "denDF"]

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmmeans", quietly = TRUE)) {
    m1.lmer <- lmerTest::lmer(logitP ~ Host*Cadavers*Ladybird + (1|Run),
        data=Ladybird.dat)
    HCL.emm <- emmeans::emmeans(m1.lmer, specs = ~ Host:Cadavers:Ladybird)
    HCL.preds <- summary(HCL.emm)
    den.df <- min(HCL.preds$df)
    ## Modify HCL.preds to be compatible with a predictions.frame
    HCL.preds <- as.predictions.frame(HCL.preds, predictions = "emmmean",
        se = "SE", interval.type = "CI",
        interval.names = c("lower.CL", "upper.CL"))
    HCL.vcov <- vcov(HCL.emm)
    HCL.sed <- NULL
}

## Use the predictions obtained with either asreml or lmerTest
if (exists("HCL.preds"))
{
## Form an all.diffs object
HCL.diffs <- allDifferences(predictions = HCL.preds,
                           classify = "Host:Cadavers:Ladybird",
                           sed = HCL.sed, vcov = HCL.vcov, tdf = den.df)

## Check the class and validity of the alldiffs object
is.alldiffs(HCL.diffs)
validAlldiffs(HCL.diffs)

## Rename Cadavers
HCL.diffs <- facRename(HCL.diffs, factor.names = "Cadavers", newnames = "Cadavers.nos")

## Check the validity of HCL.diffs
validAlldiffs(HCL.diffs)
}

getASRemlVersionLoaded

Finds the version of asreml that is loaded and returns the initial characters in version.

Description

Checks that asreml is loaded and, if it is, returns the first nchar characters of the version that is loaded.

Usage

getASRemlVersionLoaded(nchar = NULL, notloaded.fault = FALSE)

Arguments

nchar The number of characters in the asreml version to get.
notloaded.fault A logical indicating whether a fault is to occur if asreml is not loaded.

Value

A character, being the first nchar characters of the version of asreml that is loaded.

Author(s)

Chris Brien

See Also

loadASRemlVersion.

Examples

## Not run:
getASRemlVersionLoaded()
## End(Not run)
getFormulae.asreml  

Gets the formulae from an asreml object.

Description

Gets the formulae nominated in the which argument from the call stored in an asreml object.

Usage

```r
## S3 method for class 'asreml'
getFormulae(asreml.obj, which = c("fixed", "random", "residual"),
            expanded = FALSE, envir = parent.frame(), ...)
```

Arguments

- `asreml.obj`: An asreml object resulting from the fitting of a model using REML.
- `which`: A character listing the formula(e) to be extracted from the call stored in asreml.obj. It should be some combination of fixed, random, residual, sparse and all. If all is included then all formula(e) will be returned, those not having been specified in the call being NULL.
- `expanded`: A logical indicating whether terms are to be expanded to the sum of a set of individual terms.
- `envir`: The environment in which the formula(e) are to be evaluated. May also be NULL, a list, a data.frame, a pairlist or an integer as specified to `sys.call`.
- `...`: Arguments passed on to `update.formula` and ultimately to `terms.formula`.

Value

A list containing a component with each of the extracted formula(e), the name of a component being the formula that it contains.

Author(s)

Chris Brien

See Also

`printFormulae.asreml`

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                      random = ~ Row + Column + units,
                      residual = ~ ar1(Row):ar1(Column),
                      data=Wheat.dat)
getFormulae(current.asr)

## End(Not run)
```
getTestEntry.asrtests  Gets the entry for a test recorded in the test.summary data.frame of an asrtests.object

Description
Matches the label in the term column of the test.summary data.frame in the supplied asrtests.object and extracts the line for it. It only matches the last occurrence of label.

Usage
### S3 method for class 'asrtests'
getTestEntry(asrtests.obj, label, ...)

Arguments
asrtests.obj  An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.
label  A character specifying the label of the test for which the entry is required. If testranfix.asrtests was used for the test of interest, then the label will be the value of the term argument supplied to testranfix.asrtests. For changeModelOnIC.asrtests, the label will be the value of the label argument. Other arguments will be relevant for other test and change functions.
...
provision for passing arguments to functions called internally - not used at present.

Value
A one-line data.frame containing the entry.

Author(s)
Chris Brien

See Also
getTestPvalue.asrtests, as.asrtests, testranfix.asrtests, testswapran.asrtests, testresidual.asrtests, changeModelOnIC.asrtests, changeTerms.asrtests, chooseModel.asrtests

Examples
### Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety, 
random = ~ Row + Column + units, 
residual = ~ ar1(Row):ar1(Column), 
data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary(current.asrt)
# Test nugget term
current.asrt <- testranfix(current.asrt, "units", positive=TRUE)
getTestPvalue(asrtests)

*Gets the p-value for a test recorded in the test.summary data.frame of an asrtests.object*

## Description

Matches the label in the term column of the test.summary data.frame in the supplied `asrtests.object` and extracts its p-value. It only matches the last occurrence of label.

## Usage

```r
## S3 method for class 'asrtests'
getTestPvalue(asrtests.obj, label, ...)
```

## Arguments

- **asrtests.obj** An `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary`.
- **label** A character specifying the label of the test for which the p-value is required. If `testranfix.asrtests` was used for the test of interest, then the label will be the value of the term argument supplied to `testranfix.asrtests`. Other arguments will be relevant for other test functions.
- **...** provision for passing arguments to functions called internally - not used at present.

## Value

An numeric containing the p-value. It can be `NA`, for example when a p-value could not be calculated.

## Author(s)

Chris Brien

## See Also

- `getTestEntry.asrtests`, `as.asrtests`, `testranfix.asrtests`, `testswapran.asrtests`, `testresidual.asrtests`, `changeTerms.asrtests`, `chooseModel.asrtests`
## Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                      random = ~ Row + Column + units,
                      residual = ~ ar1(Row):ar1(Column),
                      data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary(current.asrt)
# Test nugget term
current.asrt <- testranfix(current.asrt, "units", positive=TRUE)
getTestPvalue(current.asrt, label = "units")
## End(Not run)
```

---

### infoCriteria

**Computes AIC and BIC for models.**

### Description

Computes Akaike and Bayesian (Schwarz) Information Criteria for models. Either the Restricted Maximum likelihood (REML) or the full likelihood (full) can be used. The full likelihood is used when it is desired to compare models that differ in their fixed models.

### Usage

```r
## S3 method for class 'asreml'
infoCriteria(object, DF = NULL,
              bound.exclusions = c("F","B","S","C"),
              ICllikelihood = "REML", fixedDF = NULL, varDF = NULL, ...)
## S3 method for class 'list'
infoCriteria(object, bound.exclusions = c("F","B","S","C"),
              ICllikelihood = "REML", fixedDF = NULL, varDF = NULL, ...)
```

### Arguments

- **object**
  An `asreml` object resulting from the fitting of a model using REML or a list of `asreml` objects. If the components of the list are named, then those names will be used as the rownames for the returned data.frame.

- **DF**
  A numeric giving the number of estimated variance parameters. If NULL then this is determined from the information in object. This argument has been replaced by varDF, but is retained for compatibility with legacy code. It is not available with the list method.

- **bound.exclusions**
  A character specifying the bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters. If set to NULL then none will be excluded.

- **IClikelihood**
  A character specifying whether Restricted Maximum Likelihood (REML) or the full likelihood (full) are to be used in calculating the information criteria for family set to asr_gaussian. For family set to asr_binomial or asr_poisson and with dispersion set to 1, the deviance is extracted from object and used
to calculate the AIC and BIC (as suggested by Damian Collins); the setting of IClikelihood is ignored and the log-likelihood set to NA. The information criteria are not valid for other settings of family and dispersion.

**fixedDF**

A numeric giving the number of estimated fixed parameters. If NULL then this is determined from the information in object. For object a list only a single value that is used for all components of the list has been implemented.

**varDF**

A numeric giving the number of estimated variance parameters. If NULL then this is determined from the information in object. It replaces the DF argument. For object a list only a single value that is used for all components of the list has been implemented.

... Provision for passing arguments to functions called internally - not used at present.

**Details**

The variance degrees of freedom (varDF) are the number of number of variance parameters that have been estimated, excluding those whose estimates have a code for bound specified in bound.exclusions. If varDF is not NULL, the supplied value is used. Otherwise varDF is determined from the information in object, i.e. if object is an asreml object then from it, or if object is a list then from each asreml object in the list. Similarly, the fixed degrees of freedom (fixedDF) are the number of number of fixed parameters that have been estimated, any coefficients that have the value NA being excluded. If fixedDF is not NULL, the supplied value is used. Otherwise fixedDF is determined from the information in object.

If ASReml-R version 4 is being used then the codes specified in bound.exclusions are not restricted to a subset of the default codes, but a warning is issued if a code other than these is specified. For ASReml-R version 3, only a subset of the default codes are allowed: F (Fixed), B (Boundary), C (Constrained) and S (Singular).

The calculation of the information criteria is an adaption of the code supplied in File S1 of Verbyla (2019). The log-likelihood is calculated as \( \loglik = \log(REML) - \log(|C|)/2 \), where \( C \) is the inverse coefficient matrix; the term involving \( C \) is omitted for REML. The AIC is calculated as \(-2 \times \loglik + 2 \times (\text{varDF} + \text{fixedDF})\) and the BIC as \(-2 \times \loglik + (\text{fixedDF} + \text{varDF}) \times \log(n - r + \text{fixedDF})\), where \( n \) is the number of observations and \( r \) is the rank of the fixed effects design matrix. For REML, fixedDF = 0.

**Value**

A data.frame containing the numbers of estimated fixed (fixedDF) and variance (varDF) parameters, the number of bound parameters (NBound), AIC, BIC and the value of the log-likelihood (loglik). All elements of the data.frame will be set to NA for the invalid combinations of family and dispersion as noted in the IClikelihood argument. If object is a list and its components are named, then those names will be used to set the rownames of the data.frame.

**Author(s)**

Chris Brien

**References**

is.alldiffs

Tests whether an object is of class alldiffs

Description

A single-line function that tests whether an object is of class alldiffs.

Usage

is.alldiffs(object)
is.asrtests

Tests whether an object is of class asrtests

Description

A single-line function that tests whether an object is of class asrtests.

Usage

is.asrtests(object)
**is.predictions.frame**

Tests whether an object is of class `predictions.frame`

A single-line function that tests whether an object is of class `predictions.frame`.

**Usage**

```r
is.predictions.frame(object)
```

**Arguments**

- `object` An object to be tested.

**Description**

A single-line function that tests whether an object is of class `predictions.frame`.

**Value**

A logical.

**Author(s)**

Chris Brien

**See Also**

`asremlPlus-package`, `asrtests.object`, `is.asrtests`, `as.asrtests`
Value

A logical.

Author(s)

Chris Brien

See Also

asremlPlus-package.predictions.frame, validPredictionsFrame, as.predictions.frame

Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics
## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety,
             random=~Blocks/Wplots,
             data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety",
          sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- as.predictions.frame(Var.pred$pvals, se = "std.error",
          est.status = "status")

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
                      data=Oats.dat)
  Var.emm <- emmeans::emmmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
          se = "SE", interval.type = "CI",
          interval.names = c("lower.CL", "upper.CL"))
}

if (exists("Var.preds"))
{
  ## Check the class and validity of the alldiffs object
  is.predictions.frame(Var.preds)
}
iterate.asrtests

Subject the fitted asreml.obj stored in an asrtests.object to further iterations of the fitting process.

Description

In an effort to improve convergence, subject the fitted asreml.obj stored in an asrtests.object to further iterations of the fitting process; the model specification is not changed. While no change is made to the test.summary, the wald.tab is updated.

Usage

## S3 method for class 'asrtests'
iterate(asrtests.obj, denDF="numeric", trace = FALSE, ...)

Arguments

asrtests.obj an asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

denDF Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

trace If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

...

further arguments passed to update.asreml.

Value

An object of S3-class asrtests.

Author(s)

Chris Brien

References


See Also

as.asrtests, asrtests.object, newfit.asreml
Examples

```r
## Not run:
current.asrt <- iterate(current.asrt)
## End(Not run)
```

**Ladybird.dat**

*Data for an experiment to investigate whether ladybirds transfer aphids*

**Description**

Welham et al. (2015, Example 8.2) describe a three-factor factorial experiment to investigate whether ladybirds transfer fungus to live aphids on plants. The three factors are Host plant (beans, trefoil), infected Cadavers (5, 10, 20), and Ladybird (-, +). A generalized randomized complete-block design is used to assign the three factors to 2 Runs, each of which involves 36 containers with a plant and live aphids. The response to be analyzed is the logit of the proportion of live aphids that were infected.

The columns in the data frame are: ID, Run, Plant, Host, Ladybird, Cadavers, Live, Infected, logitP, Prop. The column ID numbers the observations. Live, Infected, logitP, Prop are response variables.

**Usage**

```r
data(Ladybird.dat)
```

**Format**

A data.frame containing 72 observations of 10 variables.

**Author(s)**

Chris Brien

**Source**


---

**linTransform.alldiffs**

*Calculates a linear transformation of the predictions stored in an alldiffs.object.*
Description

Effects the linear transformation of the predictions in the supplied `alldiffs.object`, the transformation being specified by a matrix or a formula. The values of the transformed values are stored in an `alldiffs.object`. A matrix might be a contrast matrix or a matrix of weights for the levels of a factor used to obtain the weighted average over the levels of that factor. A formula gives rise to a projection matrix that linearly transforms the predictions so that they conform to the model specified by the formula, this model being a submodel of that inherent in the classify.

If pairwise = TRUE, all pairwise differences between the linear transforms of the predictions, their standard errors, p-values and LSD statistics are computed as using `allDifferences.data.frame`. This adds them to the `alldiffs.object` as additional list components named differences, sed, p.differences and LSD.

If a transformation has been applied (any one of transform.power is not one, scale is not one and offset is nonzero), the backtransforms of the transformed values and of the lower and upper limits of their error.intervals are added to a data.frame that is consistent with a predictions.frame. If transform.power is other than one, the standard.error column of the data.frame is set to NA. This data.frame is added to the `alldiffs.object` as a list component called backtransforms.

The printing of the components produced is controlled by the tables argument. The order of plotting the levels of one of the factors indexing the predictions can be modified and is achieved using `sort.alldiffs`.

Usage

```r
## S3 method for class 'alldiffs'
linTransform(alldiffs.obj, classify = NULL, term = NULL,
linear.transformation = NULL, Vmatrix = FALSE,
error.intervals = "Confidence",
avsed.tolerance = 0.25, accuracy.threshold = NA,
LSDtype = "overall", LSDsupplied = NULL,
LSDby = NULL, LSDstatistic = "mean",
LSDaccuracy = "maxAbsDeviation",
zero.tolerance = .Machine$double.eps ^ 0.5,
response = NULL, response.title = NULL,
x.num = NULL, x.fac = NULL,
tables = "all", level.length = NA,
pairwise = TRUE, alpha = 0.05,
inestimable.rm = TRUE, ...)```

Arguments

- `alldiffs.obj` An `alldiffs.object`.
- `classify` A character string giving the variables that define the margins of the multiway table corresponding to the predictions in `alldiffs.obj`. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator.
- `term` A character string giving the variables that define the term that was fitted using `asreml` and that corresponds to classify. It only needs to be specified when it is different to classify; it is stored as an attribute of the `alldiffs.object`. It is likely to be needed when the fitted model includes terms that involve both a numeric covariate and a factor that parallel each other; the classify would include the covariate and the term would include the factor.
linTransform.alldiffs

**linear.transformation**

A *formula* or a *matrix*. If a *formula* is given then it is taken to be a submodel of a model term corresponding to the *classify*. The projection matrix that transforms the predictions so that they conform to the submodel is obtained; the submodel does not have to involve variables in the *classify*, but the variables must be columns in the predictions component of alldiffs.obj and the space for the submodel must be a subspace of the space for the term specified by the *classify*. For example, for *classify* set to "A:B", the submodel ~ A + B will result in the predictions for the combinations of A and B being made additive for the *factors* A and B. The submodel space corresponding to A + B is a subspace of the space A:B. In this case both the submodel and the *classify* involve only the factors A and B. To fit an intercept-only submodel, specify `linear.transformation` to be the formula ~1.

If a *matrix* is provided then it will be used to apply the linear transformation to the predictions. The number of rows in the *matrix* should equal the number of linear combinations of the predictions desired and the number of columns should equal the number of predictions.

In either case, as well as the values of the linear combinations, their standard errors, pairwise differences and associated statistics are returned.

**Vmatrix**

A *logical* indicating whether the variance matrix of the predictions will be stored as a component of the *alldiffs.object* that is returned. If `linear.transformation` is set, it will be stored irrespective of the value of Vmatrix.

**error.intervals**

A *character* string indicating the type of error interval, if any, to calculate in order to indicate uncertainty in the results. Possible values are "none", "StandardError", "Confidence" and "halfLeastSignificant". The default is for confidence limits to be used. The "halfLeastSignificant" option results in half the Least Significant Difference (LSD) being added and subtracted to the predictions, the LSD being calculated using the square root of the mean of the variances of all or a subset of pairwise differences between the predictions. If the LSD is zero, as can happen when predictions are constrained to be equal, then the limits of the error intervals are set to NA. If LSDtype is set to overall, the avsed.tolerance is not NA and the range of the SEDs divided by the average of the SEDs exceeds avsed.tolerance then the error.intervals calculations and the plotting will revert to confidence intervals.

**avsed.tolerance**

A *numeric* giving the value of the SED range, the range of the SEDs divided by the square root of the mean of the variances of all or a subset of the pairwise differences, that is considered reasonable in calculating error.intervals. To have it ignored, set it to NA. It should be a value between 0 and 1. The following rules apply:

1. If avsed.tolerance is NA then mean LSDs of the type specified by LSDtype are calculated and used in error.intervals and plots.
2. Irrespective of the setting of LSDtype, if avsed.tolerance is not exceeded then the mean LSDs are used in error.intervals and plots.
3. If LSDtype is set to overall, avsed.tolerance is not NA, and avsed.tolerance is exceeded then error.intervals and plotting revert to confidence intervals.
4. If LSDtype is set to factor.combinations and avsed.tolerance is not exceeded for any factor combination then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.
linTransform.alldiffs

5. If LSDtype is set to per.prediction and avsed.tolerance is not exceeded for any prediction then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

accuracy.threshold

A numeric specifying the value of the LSD accuracy measure, which measure is specified by LSDaccuracy, as a threshold value in determining whether the halfLeastSignificant error.interval for a predicted value is a reasonable approximation; this will be the case if the LSDs across all pairwise comparisons for which the interval’s LSD was computed, as specified by LSDtype and LSDby, are similar enough to the interval’s LSD, as measured by LSDaccuracy. If it is NA, it will be ignored. If it is not NA, a column of logicals named LSDwarning will be added to the predictions component of the alldiffs.object. The value of LSDwarning for a predicted.value will be TRUE if the value of the LSDaccuracy measure computed from the LSDs for differences between this predicted.value and the other predicted.values as compared to its assignedLSD exceeds the value of accuracy.threshold. Otherwise, the value of LSDwarning for a predicted.value will be FALSE.

LSDtype

A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparison2, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations.

See LSD.frame for further information on the values in a row of this data.frame and how they are calculated.

LSDsupplied

A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function dae::fac.combine to the predictions component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into assignedLSD column of the LSD.frame stored as the LSD component of the alldiffs.object.

LSDby

A character (vector) of variables names, being the names of the factors or numerics in the classify; for each combination of their levels and values, there will be or is a row in the LSD.frame stored in the LSD component of the alldiffs.object when LSDtype is factor.combinations.

LSDstatistic

A character nominating one or more of minimum, q10, q25, mean, median, q75, q90 or maximum as the value(s) to be stored in the assignedLSD column in an LSD.frame; the values in the assignedLSD column are used in computing
halfLeastSignificant error.intervals. Here q10, q25, q75 and q90 indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function quantile is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median is calculated using the median function. Multiple values are only produced for LSDtype set to factor.combination, in which case LSDby must not be NULL and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is NULL, it is reset to mean.

LSDaccuracy A character nominating one of maxAbsDeviation, maxDeviation, q90Deviation or RootMeanSqDeviation as the statistic to be calculated as a measure of the accuracy of assignedLSD. The option q90Deviation produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named accuracyLSD in an LSD.frame.

zero.tolerance A numeric specifying the value such that if a predicted.value, its variance-covariance, or an LSD is less than it, it will be considered to be zero.

response A character specifying the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

response.title A character specifying the title for the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

x.num A character string giving the name of the numeric covariate that (i) corresponds to x.fac, (ii) is potentially included in terms in the fitted model, and (iii) which corresponds to the x-axis variable. It should have the same number of unique values as the number of levels in x.fac.

x.fac A character string giving the name of the factor that (i) corresponds to x.num, (ii) is potentially included in terms in the fitted model, and (iii) which corresponds to the x-axis variable. It should have the same number of levels as the number of unique values in x.num. The levels of x.fac must be in the order in which they are to be plotted - if they are dates, then they should be in the form yyyyymmdd, which can be achieved using as.Date. However, the levels can be non-numeric in nature, provided that x.num is also set.

tables A character vector containing a combination of none, predictions, vcov, backtransforms, differences, p.differences, sed, LSD and all. These nominate which components of the alldiffs.object to print.

level.length The maximum number of characters from the levels of factors to use in the row and column labels of the tables of pairwise differences and their p-values and standard errors.

pairwise A logical indicating whether all pairwise differences of the predictions and their standard errors and p-values are to be computed and stored. If tables is equal to "differences" or "all" or error.intervals is equal to "halfLeastSignificant", they will be stored irrespective of the value of pairwise.

alpha A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

inestimable.rm A logical indicating whether rows for predictions that are not estimable are to be removed from the components of the alldiffs.object.

... further arguments passed to redoErrorIntervals.alldiffs.
Details

For a matrix \( L \), vector of predictions \( p \) and variance matrix of the predictions \( V_p \), the linear transformed predictions are given by \( Lp \) with variance matrix \( LV_pL^T \). The last matrix is used to compute the variance of pairwise differences between the transformed values.

The matrix \( L \) is directly specified by setting `linear.transformation` to it. If `linear.transformation` is a `formula` then \( L \) is formed as the sum of the orthogonal projection matrices obtained using `pstructure.formula` from the package `dae`; `grandMean` is set to `TRUE` and `orthogonalize` to "eigenmethods".

Value

A `alldiffs.object` with the linear transformation of the predictions and their standard errors and all pairwise differences between the linear transforms of their predictions, their standard errors and p-values and LSD statistics.

If the supplied `alldiffs.object` contained a `backtransforms` component, then the returned `alldiffs.object` will contain a `backtransforms` component with the backtransformed linear transformation of the predictions. The backtransformation will, after backtransforming for any power transformation, subtract the offset and then divide by the scale.

If `error.intervals` is not "none", then the predictions component and, if present, the backtransforms component will contain columns for the lower and upper values of the limits for the interval. The names of these columns will consist of three parts separated by full stops: 1) the first part will be `lower` or `upper`; 2) the second part will be one of `Confidence`, `StandardError` or `halfLeastSignificant`; 3) the third component will be `limits`.

The name of the response, the `response.title`, the term, the `classify`, `tdf`, `alpha`, `sortFactor` and the `sortOrder` will be set as attributes to the object. Also, if `error.intervals` is "halfLeastSignificant", then those of `LSDtype`, `LSDby` and `LSDstatistic` that are not `NULL` will be added as attributes of the object and of the predictions frame; additionally, `LSDvalues` will be added as attribute of the predictions frame, `LSDvalues` being the LSD values used in calculating the `error.intervals`.

Author(s)

Chris Brien

See Also

`linTransform`, `predictPlus.asreml`, `as.alldiffs`, `print.alldiffs`, `sort.alldiffs`, `subset.alldiffs`, `alldifferences.data.frame`, `redoErrorIntervals.alldiffs`, `recalcLSD.alldiffs`, `pickLSDstatistics.alldiffs`, `predictPresent.asreml`, `plotPredictions.data.frame`, `as.Date`, `predict.asreml`

Examples

data(WaterRunoff.dat)

## Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) # required for asreml-R4 only

current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                      random = ~ Benches:MainPlots,
                      keep.order=TRUE, data= WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
# Get additive predictions directly using predictPlus

diffs.sub <- predictPlus(asreml(classify = "Sources:Species", Vmatrix = TRUE,
    linear.transformation = ~ Sources + Species,
    asreml.obj = current.asr, tables = "none",
    wald.tab = current.asrt$wald.tab,
    present = c("Type","Species","Sources"))

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmmeans", quietly = TRUE))
{
    m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * Species) +
                                (1|Benches:MainPlots),
                                data=na.omit(WaterRunoff.dat))
    SS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Species)
    SS.preds <- summary(SS.emm)
    den.df <- min(SS.preds$df, na.rm = TRUE)
    ## Modify SS.preds to be compatible with a predictions.frame
    SS.preds <- as.predictions.frame(SS.preds, predictions = "emmean",
        se = "SE", interval.type = "CI",
        interval.names = c("lower.CL", "upper.CL"))

    ## Form an all.diffs object and check its validity
    SS.vcov <- vcov(SS.emm)
    SS.diffs <- allDifferences(predictions = SS.preds, classify = "Sources:Species",
                                vcov = SS.vcov, tdf = den.df)
    validAlldiffs(SS.diffs)

    ## Get additive predictions
    diffs.sub <- linTransform(SS.diffs, classify = "Sources:Species",
                                linear.transformation = ~ Sources + Species,
                                Vmatrix = TRUE, tables = "none")
}

## Calculate contrasts from prediction obtained using asreml or lmerTest
if (exists("diffs.sub"))
{
    # Contrast matrix for differences between each species and non-planted for the last source
    L <- cbind(matrix(rep(0,7*32), nrow = 7, ncol = 32),
               diag(1, nrow = 7),
               matrix(rep(-1, 7), ncol = 1))
    rownames(L) <- as.character(diffs.sub$predictions$Species[33:39])
    diffs.L <- linTransform(diffs.sub,
                            classify = "Sources:Species",
                            linear.transformation = L,
                            tables = "predictions")
}

---

loadASRemlVersion | Ensures that a specific version of asreml is loaded.
**LSD.frame**

**Description**

Loads the specified version of asreml, provided that it is not already loaded. If the version of asreml is not the required version, then the loaded version is unloaded first.

**Usage**

```r
loadASRemlVersion(version = 4, ...)
```

**Arguments**

- `version`: The version that is to be loaded, the version consisting of just the initial characters that are significant in the version that should be loaded. For example, the default value of 4 implies that any version that begins with "4" is acceptable. It is used to check that the required version is loaded.

- `...`: Other library/require arguments that are needed to load the specified version of asreml.

**Value**

A character, being all characters in the version of asreml that is loaded on exit from the function.

**Author(s)**

Chris Brien

**See Also**

`getASRemlVersionLoaded`.

**Examples**

```r
## Not run:
loadASRemlVersion(3, lib.loc = "D:\Analyses\R asreml3")
## End(Not run)
```

---

**LSD.frame**

*Description of an LSD frame*

**Description**

A data.frame that stores Least Significant differences (LSDs) for predictions for a fitted model.

**Value**

A data.frame that can be a component of an `alldiffs.object` and that contains LSD values and statistics to be used in determining the significance of the pairwise differences. In particular, they are used in calculating halfLeastSignificant limits to be included in a predictions.frame.

Exactly what an LSD.frame contains is determined by the following arguments to functions that return an `alldiffs.object`: LSDtype, LSDby, LSDstatistic, LSDaccuracy and LSDsupplied. The rownames of the LSD.frame indicate, for each of its rows, for what group of predictions the entries in the row were calculated, this being controlled by the LSDtype and LSDby arguments.
The values for all of the LSD arguments are stored as attributes to the `alldiffs.object` and the predictions and, if present backtransforms, components of the `alldiffs.object`.

An `LSD.frame` always has the eight columns `c`, `minimumLSD`, `meanLSD`, `maximumLSD`, `assignedLSD`, `accuracyLSD`, `falsePos` and `falseNeg`.

1. `c`: This gives the number of pairwise coomparison of predictions for the combinations of the factor levels given by the row name. If the row name is `overall` then it is for all predictions.

2. `minimumLSD`, `meanLSD`, `maximumLSD`: These are computed for either `overall`, `factor.combinations`, `per.prediction` or supplied LSD values, as specified by the `LSDtype` argument. The `meanLSD` is calculated using the square root of the mean of the variances of set of pairwise differences appropriate to the specific `LSDtype` argument.

   For `overall`, the mean, minimum and maximum of the LSDs for all pairwise comparisons are computed.

   If `factor.combinations` was specified for `LSDtype` when the LSDs were being calculated, then the `LSD.frame` contains a row for each combination of the values of the `factors` and `numeric` specified by `LSDby`. The values in a row are calculated from the LSD values for the pairwise differences for each combination of the `factors` and `numeric` values, unless there is only one prediction for a combination, when notional LSDs are calculated that are based on the standard error of the prediction multiplied by the square root of two.

   For `per.prediction`, the minimum, mean and maximum LSD, based, for each prediction, on the LSD values for all pairwise differences involving that prediction are computed.

   For supplied, the `LSD.frame` is set up based on the setting of `LSDby`: a single row with name `overall` if `LSDby` is `NULL` or, if `LSDby` is a vector of `factor` and `numeric` names, rows for each observed combinations of the values of the named `factors` and `numeric`. The `LSDsupplied` argument is used to provide the values to be stored in the column `assignedLSD`.

3. `assignedLSD`: The `assignedLSD` column contains the values that are assigned for use in calculating `halfLeastSignificant error.intervals`. Its contents are determined by `LSDstatistic` and `LSDsupplied` arguments. The `LSDsupplied` argument allows the direct specification of values to be placed in the `assignedLSD` column of the `LSD.frame`. The default is to use the values in the `meanLSD` column.

4. `LSDaccuracy`: The `LSDaccuracy` gives an indication of the proportion that the correct LSD for a single `predicted.value` might deviate from its `assignedLSD` value. The contents of the `accuracyLSD` column is controlled by the `LSDaccuracy` argument.

5. `falsePos` and `falseNeg`: These columns contain the number of false positives and negatives if the `assignedLSD` value(s) is(are) used to determine the significance of the pairwise predictions differences. Each LSD value in the `assignedLSD` column is used to determine the significance of pairwise differences that involve predictions for the combination of values given by the row name for the LSD value.

See `recalcLSD.alldiffs` for more information.

**Author(s)**

Chris Brien

**See Also**

`recalcLSD.alldiffs`, `redoErrorIntervals.alldiffs`, `predictPresent.asreml`, `predictPlus.asreml`
Examples

```r
data(Oats.dat)

## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety,
                 random=~Blocks/Wplots,
                 data=Oats.dat)
current.asr <- as.asrtests(m1.asr)
Var.diffs <- predictPlus(m1.asr, classify="Nitrogen:Variety",
                         wald.tab = current.asr$wald.tab,
                         tables = "none")

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
   requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
                            data=Oats.dat)
  #Get predictions
  Var.emm <- emmeans::emmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                     se = "SE", interval.type = "CI",
                                     interval.names = c("lower.CL", "upper.CL"))
  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
  #Set up an alldiffs object, which includes overall LSDs
  Var.diffs <- allDifferences(predictions = Var.preds, classify = "Variety:Nitrogen",
                              sed = Var.sed, vcov = Var.vcov, tdf = 45)
}

if (exists("Var.diffs"))
{
  ## Use recalcLSD to get LSDs for within Variety differences
  Var.LSD.diffs <- recalcLSD(Var.diffs,
                              LSDtype = "factor.combinations", LSDby = "Variety")
  print(Var.LSD.diffs$LSD)
}
```

Description

Prepares the fixed and random sP-spline basis matrices, and associated information, that are needed for fitting of Tensor Product P-splines (TPPS) as described by Rodriguez-Alvarez et al. (2018).
When \texttt{asreml.option} is set to \texttt{mbf}, \texttt{makeTPPSplineMats.data.frame} must be run prior to fitting TPPS models for local spatial variation using \texttt{addSpatialModelOnIC.asrtests} and \texttt{chooseSpatialModelOnIC.asrtests}. The spline basis matrices are created in the parent environment of \texttt{makeTPPSplineMats.data.frame} when it is called. If the \texttt{grp} is to be used to supply the basis functions to asreml-R, then this function need not be called; the spatial spline fitting functions will set up the basis functions.

### Usage

```r
define makeTPPSplineMats(data, sections = NULL, row.covar, col.covar, nsegs = NULL, nestorder = c(1,1), degree = c(3,3), difforder = c(2,2), asreml.option = "mbf", ...)
```

### Arguments

- **data**: An \texttt{data.frame} that holds the spline bases for a section. It is indexed by columns named \texttt{col} and \texttt{row}.
- **sections**: A single character string that species the name of the column in the \texttt{data.frame} that contains the \texttt{factor} that identifies different sections of the data to which separate spatial models are to be fitted.
- **row.covar**: A single character string nominating a \texttt{numeric} column in the \texttt{data.frame} that contains the values of a covariate indexing the rows of the grid.
- **col.covar**: A single character string nominating a \texttt{numeric} column in the \texttt{data.frame} that contains the values of a covariate indexing the columns of the grid.
- **nsegs**: A pair of \texttt{numeric} values giving the number of segments into which the column and row ranges are to be split, respectively (each value specifies the number of internal knots + 1). If only one number is specified, that value is used in both dimensions. If not specified, then (number of unique values - 1) is used in each dimension; for a grid layout with equal spacing, this gives a knot at each data value.
- **nestorder**: A character of length 2. The order of nesting for column and row dimensions, respectively; default=1 (no nesting). A value of 2 generates a spline with half the number of segments in that dimension, etc. The number of segments in each direction must be a multiple of the order of nesting.
- **degree**: A character of length 2. The degree of polynomial spline to be used for column and row dimensions respectively; default=3.
- **difforder**: A character of length 2. The order of differencing for column and row dimensions, respectively; default=2.
- **asreml.option**: A single character string whose value may be \texttt{mbf} or \texttt{grp}, indicating the method is to be used to supply externally formed covariate matrices to asreml.
- **...**: Further arguments passed to \texttt{tpsmmb} from package TPSbits.

### Details

The objects are formed using the function \texttt{tpsmmb} from the R package TPSbits authored by Sue Welham (2022).

Each combination of a \texttt{row.covar} and a \texttt{col.covar} does not have to specify a single observation; for example, to fit a local spatial variation model to the main units of a split-unit design, each
combination would correspond to a main unit and all subunits of the main unit would have the same combination.

The data for experiment can be divided sections and the spline bases and associated data will be produced for each section. If there is more than one sections, then a list is returned that has a component for each section. The component for a section contains:

**Value**

A list of length equal to the number of sections is produced. Each of these components is a list with 8 or 9 components named data.plus, being the input data.frame to which has been added the columns required to fit the TPPS model (the data.frame stored in the data component holds only the covariates from data).

List of length 8 or 9 (according to the asrem1 option).

1. data = the input data frame augmented with structures required to fit tensor product splines in asreml-R. This data frame can be used to fit the TPS model.
   
   Added columns:
   
   - TP.col, TP.row = column and row coordinates
   - TP.CxR = combined index for use with smooth x smooth term
   - TP.C.n for n=1:diff.c = X parts of column spline for use in random model (where diff.c is the order of column differencing)
   - TP.R.n for n=1:diff.r = X parts of row spline for use in random model (where diff.r is the order of row differencing)
   - TP.CR.n for n=1:(diff.c*diff.r) = interaction between the two X parts for use in fixed model. The first variate is a constant term which should be omitted from the model when the constant (1) is present. If all elements are included in the model then the constant term should be omitted, eg. y ~ -1 + TP.CR.1 + TP.CR.2 + TP.CR.3 + TP.CR.4 + other terms...
   - when asrem1="grp" or "sepgrp", the spline basis functions are also added into the data frame. Column numbers for each term are given in the grp list structure.

2. mbflist = list that can be used in call to asreml (so long as Z matrix data frames extracted with right names, eg BcZ<stub>.df)

3. BcZ.df = mbf data frame mapping onto smooth part of column spline, last column (labelled TP.col) gives column index

4. BrZ.df = mbf data frame mapping onto smooth part of row spline, last column (labelled TP.row) gives row index

5. BcrZ.df = mbf data frame mapping onto smooth x smooth term, last column (labelled TP.CxR) maps onto col x row combined index

6. dim = list structure, holding dimension values relating to the model:
   
   - "diff.c" = order of differencing used in column dimension
   - "nbc" = number of random basis functions in column dimension
   - "nbcn" = number of nested random basis functions in column dimension used in smooth x smooth term
   - "diff.r" = order of differencing used in column dimension
   - "nbr" = number of random basis functions in column dimension
   - "nbrn" = number of nested random basis functions in column dimension used in smooth x smooth term
7. trace = list of trace values for \(Z G Z'\) for the random TP spline terms, where \(Z\) is the design matrix and \(G\) is the known diagonal variance matrix derived from eigenvalues. This can be used to rescale the spline design matrix (or equivalently variance components).

8. grp = list structure, only added for setting \(asreml="grp\). For \(asreml="grp\), provides column indexes for each of the 5 random components of the 2D splines in data.plus. Dimensions of the components can be derived from the values in the dim item.

9. data.plus = the input data.frame to which has been added the columns required to fit tensor product splines in asreml-R. This data.frame can be used to fit the TPS model. For multiple sections, this data.frame will occur in the component for each section.

Author(s)
Chris Brien

References

Welham, S. J. (2022) TPSbits: Creates Structures to Enable Fitting and Examination of 2D Tensor-ProductSplines using ASReml-R. Version 1.0.0 [https://mmade.org/tpsbits/]

See Also
addSpatialModelOnIC.asrtests, chooseSpatialModelOnIC.asrtests, tpsmmb

Examples

```r
## Not run:

data(Wheat.dat)

#Add row and column covariates
Wheat.dat <- within(Wheat.dat, {
  cColumn <- dae::as.numfac(Column)
  cColumn <- cColumn - mean(unique(cColumn))
  cRow <- dae::as.numfac(Row)
  cRow <- cRow - mean(unique(cRow))
})

#Set up the matrices
tps.Xmat <- makeTPPSplineMats(wheat.dat,
  row.covar = "cRow", col.covar = "cColumn")

## End(Not run)
```

newfit.asreml

Refits an asreml model with modified model formula using either a call to update.asreml or a direct call to asreml.
Description

Extracts the call from the asreml.obj and evaluates that call, replacing any arguments with changed values. If update is TRUE and set.terms is not set, the call is evaluated using update.asreml; otherwise, it is evaluated using a direct call to asreml. The principal difference is that the latter does not enforce the use of previous values of the variance parameters as initial values; it sets G.param and R.param to NULL or to values as specified for set.terms. The ... argument can be used to pass G.param and/or R.param, provided update is FALSE and set.terms is not set.

Usage

```r
## S3 method for class 'asreml'
newfit(asreml.obj, fixed., random., sparse., residual., rcov., update = TRUE, trace = FALSE,
       allow.unconverged = TRUE, allow.fixedcorrelation = TRUE,
       keep.order = TRUE,
       set.terms = NULL, ignore.suffices = TRUE,
       bounds = "P", initial.values = NA, ...)
```

Arguments

- `asreml.obj` A valid asreml object with with a component named call (from a previous call to either asreml or update.asreml).
- `fixed.` A character or formula specifying changes to the fixed formula. This is a two-sided formula where "." is substituted for existing components in the fixed component of asreml.obj$call. If changes are specified, the fixed terms will be reordered so that single-variable terms come first, followed by two-variable terms and so on.
- `random.` A character or formula specifying changes to the random formula. This is a one-sided formula where "." is substituted for existing components in the random component of asreml.obj$call.
- `sparse.` A character or formula specifying changes to the sparse formula. This is a one-sided formula where "." is substituted for existing components in the sparse component of asreml.obj$call.
- `residual.` A character or formula specifying changes to the error formula, used when version 4 or later of ASReml-R is loaded. This is a one-sided formula where "." is substituted for existing components in the residual component of asreml.obj$call.
- `rcov.` A character or formula specifying changes to the error formula, used when version 3 of ASReml-R is loaded. This is a one-sided formula where "." is substituted for existing components in the residual component of asreml.obj$call.
- `update` A logical indicated whether to use update.asreml or asreml to evaluate the modified call. If TRUE, use update.asreml to evaluate the modified call. In doing this the arguments R.param and G.param are set to those in the asreml.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml itself, in which the only changes from the previous call are those specified in the arguments to newfit.asreml.
- `trace` A logical that control output from ASReml-R. If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.
allow.unconverged
A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit does not converge, the supplied asreml.obj is returned.

allow.fixedcorrelation
A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asreml.obj is returned. The fit in the supplied the asreml.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

keep.order
A logical value indicating whether the terms should keep their positions. If FALSE the terms are reordered so that main effects come first, followed by the interactions, all second-order, all third-order and so on. Effects of a given order are kept in the order specified.

set.terms
A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the vparameters component of the new asreml.obj.

ignore.suffixes
A logical vector specifying whether the suffixes of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffixes are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffixes for all the terms in terms.

bounds
A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

initial.values
A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

... additional arguments to the call, or arguments with changed values.

Value
An asreml object.

Author(s)
Chris Brien

References
num.recode

Recodes the unique values of a vector using the values in a new vector.

Description

Recodes the unique values of a variate using the value in position i of the new.values vector to replace the ith sorted unique values of x. The new levels do not have to be unique.

Usage

num.recode(x, new.values)

Arguments

x The vector to be recoded.
new.values A vector of length unique(x) containing values to use in the recoding.

Value

A vector.

Author(s)

Chris Brien

See Also

dae::fac.recode.

Examples

## set up a factor with labels
x <- rep(c(-42, -14, 14, 42), 4)

## recode x
b <- num.recode(x, c(0, 28, 56, 84))
pairdiffsTransform.alldiffs

Data for an experiment to investigate nitrogen response of 3 oats varieties

Description

Yates (1937) describes a split-plot experiment that investigates the effects of three varieties of oats and four levels of Nitrogen fertilizer. The varieties are assigned to the main plots using a randomized complete block design with 6 blocks and the nitrogen levels are randomly assigned to the subplots in each main plot.

The columns in the data frame are: Blocks, Wplots, Subplots, Variety, Nitrogen, xNitrogen, Yield. The column xNitrogen is a numeric version of the factor Nitrogen. The response variable is Yield.

Usage

data(Oats.dat)

Format

A data.frame containing 72 observations of 7 variables.

Author(s)

Chris Brien

Source


pairdiffsTransform.alldiffs

Calculates the differences between nominated pairs of predictions stored in an alldiffs.object.

Description

Predictions of differences and their error intervals are formed for two levels of a factor, the pairs.factor in numerator.levels with a level in denominator.levels, an alldiffs.object is formed that contains the differences between predictions with this pair of levels for all of the combinations of the levels of the other factors in the classify of the alldiffs.object. These prediction differences are obtained using linTransform by forming a suitable contrast matrix to specify the linear.transformation. This function has the advantage that the factors indexing the differences are included in the components of the alldiffs.objects. If pairwise = TRUE, all pairwise differences between the linear transforms of the predictions, their standard errors, p-values and LSD statistics are computed as using allDifferences.data.frame. This adds them to the alldiffs.object as additional list components named differences, sed, p.differences and LSD.

The printing of the components produced is controlled by the tables argument. The order of plotting the levels of one of the factors indexing the predictions can be modified and is achieved using sort.alldiffs.
Usage

```r
## S3 method for class 'alldiffs'
pairdiffsTransform(alldiffs.obj, pairs.factor, first.levels, second.levels,
                   Vmatrix = FALSE, error.intervals = "Confidence",
                   avsed.tolerance = 0.25, accuracy.threshold = NA,
                   LSDtype = "overall", LSDsupplied = NULL, LSDby = NULL,
                   LSDstatistic = "mean", LSDaccuracy = "maxAbsDeviation",
                   response = NULL, response.title = NULL, tables = "all",
                   pairwise = TRUE, alpha = 0.05, ...)```

Arguments

- `alldiffs.obj` An `alldiffs.object`
- `pairs.factor` A character string giving the name of the factor for whose levels the differences are to be calculated.
- `first.levels` A character string containing the levels of the `pairs.factor` whose predictions are those subtracted from.
- `second.levels` A character string containing the levels of the `pairs.factor` whose predictions are those that are subtracted.
- `Vmatrix` A logical indicating whether the variance matrix of the predictions will be stored as a component of the `alldiffs.object` that is returned.
- `error.intervals` A character string indicating the type of error interval, if any, to calculate in order to indicate uncertainty in the results. Possible values are "none", "StandardError", "Confidence" and "halfLeastSignificant". The default is for confidence limits to be used. The "halfLeastSignificant" option results in half the Least Significant Difference (LSD) being added and subtracted to the predictions, the LSD being calculated using the square root of the mean of the variances of all or a subset of pairwise differences between the predictions. If the LSD is zero, as can happen when predictions are constrained to be equal, then the limits of the error intervals are set to `NA`. If `LSDtype` is set to `overall`, the `avsed.tolerance` is not `NA` and the range of the SEDs divided by the average of the SEDs exceeds `avsed.tolerance` then the `error.intervals` calculations and the plotting will revert to confidence intervals.
- `avsed.tolerance` A numeric giving the value of the SED range, the range of the SEDs divided by the square root of the mean of the variances of all or a subset of the pairwise differences, that is considered reasonable in calculating `error.intervals`. To have it ignored, set it to `NA`. It should be a value between 0 and 1. The following rules apply:
  1. If `avsed.tolerance` is `NA` then mean LSDs of the type specified by `LSDtype` are calculated and used in `error.intervals` and plots.
  2. Irrespective of the setting of `LSDtype`, if `avsed.tolerance` is not exceeded then the mean LSDs are used in `error.intervals` and plots.
  3. If `LSDtype` is set to `overall`, `avsed.tolerance` is not `NA`, and `avsed.tolerance` is exceeded then `error.intervals` and plotting revert to confidence intervals.
  4. If `LSDtype` is set to `factor.combinations` and `avsed.tolerance` is not exceeded for any factor combination then the half LSDs are used in `error.intervals` and plots; otherwise, `error.intervals` and plotting revert to confidence intervals.
5. If LSDtype is set to per.prediction and avsed.tolerance is not exceeded for any prediction then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

accuracy.threshold
A numeric specifying the value of the LSD accuracy measure, which measure is specified by LSDaccuracy, as a threshold value in determining whether the halfLeastSignificant error.interval for a predicted value is a reasonable approximation; this will be the case if the LSDs across all pairwise comparisons for which the interval’s LSD was computed, as specified by LSDtype and LSDby, are similar enough to the interval’s LSD, as measured by LSDaccuracy. If it is NA, it will be ignored. If it is not NA, a column of logicals named LSDwarning will be added to the predictions component of the alldiffs.object. The value of LSDwarning for a predicted.value will be TRUE if the value of the LSDaccuracy measure computed from the LSDs for differences between this predicted.value and the other predicted.values as compared to its assignedLSD exceeds the value of accuracy.threshold. Otherwise, the value of LSDwarning for a predicted.value will be FALSE.

LSDtype
A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparison2, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations. See LSD.frame for further information on the values in a row of this data.frame and how they are calculated.

LSDsupplied
A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function dae::fac.combine to the predictions component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into assignedLSD column of the LSD.frame stored as the LSD component of the alldiffs.object.

LSDby
A character (vector) of variables names, being the names of the factors or numerics in the classify; for each combination of their levels and values, there will be or is a row in the LSD.frame stored in the LSD component of the alldiffs.object when LSDtype is factor.combinations.

LSDstatistic
A character nominating one or more of minimum, q10, q25, mean, median, q75, q90 or maximum as the value(s) to be stored in the assignedLSD column in an LSD.frame; the values in the assignedLSD column are used in computing
halfLeastSignificant error.intervals. Here q10, q25, q75 and q90 indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function quantile is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median is calculated using the median function. Multiple values are only produced for LSDtype set to factor.combination, in which case LSDby must not be NULL and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is NULL, it is reset to mean.

LSDaccuracy A character nominating one of maxAbsDeviation, maxDeviation, q90Deviation or RootMeanSqDeviation as the statistic to be calculated as a measure of the accuracy of assignedLSD. The option q90Deviation produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named accuracyLSD in an LSD.frame.

response A character specifying the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

response.title A character specifying the title for the response variable for the predictions. It is stored as an attribute to the alldiffs.object.

tables A character vector containing a combination of none, predictions, vcov, backtransforms, differences, p.differences, sed, LSD and all. These nominate which components of the alldiffs.object to print.

pairwise A logical indicating whether all pairwise differences of the predictions and their standard errors and p-values are to be computed and stored. If tables is equal to "differences" or "all" or error.intervals is equal to "halfLeastSignificant", they will be stored irrespective of the value of pairwise.

alpha A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

... further arguments passed to linTransform.alldiffs.

Value

A list of alldiffs.objects with a component for each combination of a first.levels with a second.levels. The name of a component will be a level from first.levels combined with a level from second.levels, separated by a comma. If the predictions in the supplied alldiffs.object are based on a response that was transformed, each alldiffs.object in the list will include a backtransforms component that contains a column labelled backtransformed.predictions, along with the backtransforms of the nominated error.intervals. The predictions and backtransforms components in an alldiffs.object will be indexed by the variables in the classify of alldiffs.obj, except that the pairs.factor is omitted. If the transformation was the logarithmic transformation, these backtransformed.predictions are predicted ratios of the untransformed response.

If sortFactor attribute is set and is not the ratio.factor, the predictions and, if present, their backtransforms will be sorted using the sortOrder attribute of the alldiffs.object, and both sortFactor and sortOrder will be set as attributes to the object.

Author(s)

Chris Brien
See Also

pairdiffsTransform, alldiffs, print.alldiffs, sort.alldiffs, subset.alldiffs, alldifferences.data.frame, redoErrorIntervals.alldiffs, recalcLSD.alldiffs, pickLSDstatistics.alldiffs, predictPresent.asreml, plotPredictions.data.frame, as.Date, predict.asreml

Examples

#### Form the differences for log(RGR) for Salinity

```r
load(system.file("extdata", "testDiffs.rda", package = "asremlPlus", mustWork = TRUE))
#### For the ratios for Cl per WU Temperature - use backtransforms of log-predictions
Preds.ratio.ClUp <- pairdiffsTransform(diffs.ClUp, pairs.factor = "Temperature", first.levels = "Hot", second.levels = "Cool", error.intervals = "halfLeast", tables = "backtransforms")  # Backtransforms are ratios

#### Form the differences for Nitrogen compared to no Nitrogen

data("Oats.dat")
## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety, random=~Blocks/Wplots, data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
wald.tab <- current.asrt$wald.tab
Var.diffs <- predictPlus(m1.asr, classify="Nitrogen:Variety", pairwise = TRUE, Vmatrix = TRUE, error.intervals = "halfLeast", LSDtype = "factor", LSDby = "Variety", wald.tab = wald.tab)

## End(Not run)

## Use lme4 and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE)) {
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots), data=Oats.dat)
  ## Set up a wald.tab
  int <- as.data.frame(rbind(rep(NA,4)))
  rownames(int) <- "(Intercept)"
  wald.tab <- anova(m1.lmer, ddf = "Kenward", type = 1)[3:6]
  names(wald.tab) <- c("Df", "denDF", "F.inc", "Pr")
  wald.tab <- rbind(int, wald.tab)
  # Get predictions
  Var.emm <- emmeans::emmmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean", se = "SE", interval.type = "CI", interval.names = c("lower.CL", "upper.CL"))
  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
den.df <- wald.tab[match("Variety", rownames(wald.tab)), "denDF"]
}
permute.square

Permutes the rows and columns of a square matrix.

Description
Permutes the rows and columns of a square matrix.

Usage
permute.square(x, permutation)

Arguments
x
A square matrix.

permutation
A vector specifying the new order of rows and columns.

Value
A square matrix.

Author(s)
Chris Brien

See Also
permute.to.zero.lowertri

Examples
terms.marginality <- matrix(c(1,0,0,0, 0,1,0,0, 0,1,1,0, 0,1,1,0, 1,1,1,1,1), nrow=5)
permtn <- c(3,2,4,5)
terms.marginality <- permute.square(terms.marginality, permtn)
permute.to.zero.lowertri

Permutes a square matrix until all the lower triangular elements are zero.

Description
Permutes a square matrix until all the lower triangular elements are zero.

Usage
permute.to.zero.lowertri(x)

Arguments
x  A square matrix of order n with at least n*(n-1)/2 zero elements.

Value
A square matrix.

Author(s)
Chris Brien

See Also
permute.square

Examples

```r
terms.marginality <- matrix(c(1,0,0,0, 0,1,0,0, 0,1,1,0, 0,1,1,1, 1,1,1,1,1), nrow=5)
terms.marginality <- permute.to.zero.lowertri(terms.marginality)
```

pickLSDstatistics.alldiffs

Pick LSDstatistics whose values minimize the number of errors in pairwise comparisons of predictions.

Description
Given an alldiffs.object with an sed component, exploreLSDs.alldiffs is used to calculate the LSD values for each set of prediction comparisons specified by LSDtype and LSDby using each of the statistics minimum, q10, q25, mean, median, q75, q90 and maximum. Then the numbers of false positives and false negatives resulting from employing each of the calculated LSDs is obtained. For each set of comparisons, the LSD value(s) with the lowest number of false positives are identified and, from these, the smallest value with the lowest number of false negatives. That is, a conservative approach is taken to picking LSD values by putting the priority on avoiding false positives. Before using the LSDstatistics that this function suggests, the number of false positives and negatives
generated by them should be checked. For example, it may be that there are too many false negatives
and a better balance between the numbers of false positives and negatives can be identified using
`exploreLSDs.alldiffs`.

Usage

```r
## S3 method for class 'alldiffs'
pickLSDstatistics(alldiffs.obj,
   LSDtype = "overall", LSDby = NULL,
   alpha = 0.05, digits = 3,
   retain.zeroLSDs = FALSE,
   zero.tolerance = .Machine$double.eps ^ 0.5,
   ...)
```

Arguments

- **alldiffs.obj**: An `alldiffs.object`.
- **LSDtype**: A character string that can be overall or factor.combinations. It
determines whether the LSD values that are investigated and stored are (i) the
overall minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum of all pairwise LSDs, or (ii) the minimum, quantile10, quantile25, mean, median, quantile75, quantile90, and maximum for the pairwise LSDs for each combination of the values of the factors and numerics
named in LSDby, unless there is only one prediction for a combination, when
notional LSDs are calculated. The LSDtype specified here does not have to
match that used in the creating the `alldiffs.object`.
See `LSD.frame` for further information on how the LSD statistics are calculated.
- **LSDby**: A character (vector) of variables names, being the names of the
factors or numerics in the classify; for each combination of the values the of the
factors and numerics, the LSD errors are to be computed when LSDtype is
factor.combinations. The LSDby specified here does not have to match that
used in the creating the `alldiffs.object`.
- **alpha**: A numeric specifying the significance level for an LSD to compare a pair of
predictions.
- **digits**: A numeric specifying the number of significant digits to retain in rounding the
LSDs before determining the distinct rounded LSDs.
- **retain.zeroLSDs**: A logical indicating whether to retain or omit LSDs that are zero when calculating the summaries of LSDs.
- **zero.tolerance**: A numeric specifying the value such that if an LSD is less than it, it will be
considered to be zero.
- **...**: Provision for passsing arguments to functions called internally - not used at present.

Value

A character of length one for LSDby set to overall or of length equal to the number of observed
combinations of the values of the factors and numerics in LSDby. Each element of the returned
character is one of minimum, q10, q25, mean, median, q75, q90 or maximum, reflecting the value(s)
of the LSD from amongst those calculated that minimizes the number of false positives; if there is
more than one such value, then the element will be correspond to the value of the LSD from amongst
those with the minimum number of false positives that minimizes the number of false negatives.
Author(s)

Chris Brien

See Also

asremlPlus-package, exploreLSDs.alldiffs plotLSDs.data.frame, plotLSDs.alldiffs, plotLSDerrors.alldiffs, plotLSDerrors.data.frame, recalcLSD.alldiffs, redoErrorIntervals.alldiffs

Examples

data(WaterRunoff.dat)

## Use asreml to get predictions and associated statistics

### Not run:

```r
asreml.options(keep.order = TRUE) # required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                      random = ~ Benches:MainPlots,
                      keep.order = TRUE, data = WaterRunoff.dat)

current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus(classify = "Sources:Type",
                        asreml.obj = current.asr,
                        wald.tab = current.asrt$wald.tab,
                        present = c("Sources", "Type", "Species"))
```

### End(Not run)

### Use lmerTest and emmmeans to get predictions and associated statistics

```r
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmmeans", quietly = TRUE)) {

  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                           (1|Benches:MainPlots),
                           data = na.omit(WaterRunoff.dat))

  TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
  TS.preds <- summary(TS.emm)
  den.df <- min(TS.preds$df, na.rm = TRUE)

  ## Modify TS.preds to be compatible with a predictions.frame
  TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
                                   se = "SE", interval.type = "CI",
                                   interval.names = c("lower.CL", "upper.CL"))

  ## Form an all.diffs object and check its validity
  els <- as.numeric(rownames(TS.preds))
  TS.vcov <- vcov(TS.emm)[els, els]
  TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
                             vcov = TS.vcov, tdf = den.df)

  validAlldiffs(TS.diffs)
}
```

### Plot p-values for predictions obtained using asreml or lmerTest

```r
if (exists("TS.diffs")) {
  ## Pick the LSD values for predictions obtained using asreml or lmerTest
```
plotLSDerrors.alldiffs

Plots a map of the errors that occur in using the computed LSD values for pairwise differences between predictions.

Description

Produces a plot of the errors that occur in using the computed LSD values for pairwise differences predictions by comparing the result obtained from using the LSDs stored in the assignedLSD column of the LSD component of the alldiffs.object with those computed from the sed component using the t-value for the df stored in the tdf attribute of the alldiffs.object. The sed component is generally a matrix whose rows and columns are labelled by the levels of one or more factors, the set of labels being the same for rows and columns. The sections argument allows multiple plots to be produced, one for each combination of the levels of the factors listed in sections. Otherwise, a single plot is produced for all observed combinations of the levels of the factors in the classify attribute for the alldiffs.object. The plots are produced using plotLSDerrors.data.frame. The order of plotting the levels of one of the factors indexing the predictions can be modified using sort.alldiffs.

Usage

plotLSDerrors(object, ...)

## S3 method for class 'alldiffs'
plotLSDerrors(object, alpha = 0.05, useIntervals = FALSE, sections = NULL, gridspacing = 0, factors.per.grid = 0,
triangles = "both", title = NULL,
axis.labels = TRUE, axis.text.size = 12,
sep="", colours = c("white","blue","red","grey"),
ggplotFuncs = NULL, printPlot = TRUE,
sortFactor = NULL, sortParallelToCombo = NULL,
sortNestingFactor = NULL, sortOrder = NULL,
decreasing = FALSE, ...)

Arguments

object An alldiffs.object with both LSD and sed components that are not NULL.
alpha A numeric giving the significance level for the LSD.
useIntervals A logical indicating whether to use the interval limits stored in the predictions component of object, instead of the LSDs stored in the LSD component, for determining whether pairs of predictions are significantly different. It allows a
check of how the error.intervals in the predictions component will perform if they are used for all pairwise predictions comparisons, whereas the comparisons to which the LSDs apply may be restricted by the setting of the LSDby attribute of object. There is no restriction on the error.intervals that can be used. However, the limits for them must be in columns in the predictions component of object and their names must end with .limits and begin with lower. and upper.

sections A character listing the names of the factors that are to be used to break the plot into sections. A separate plot will be produced for each observed combination of the levels of these factors.

gridspacing A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. An alternative is to specify the factors.per.grid argument to have the grid spacings automatically calculated. Grids are most useful when two or more factors index the rows and columns. If a single, nonzero number, k say, is given then a grid line is placed after every kth row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.

factors.per.grid A numeric specifying the number of factors to include within each grid of differences. The gridspacing will then be computed based on the numbers of combinations observed within the levels of the remaining factors in a single plot. The gridspacing argument to this function will be ignored if factors.per.grid is greater than zero. Grids are most useful when two or more factors index the rows and columns of each plot.

triangles A character indicating whether the plot should include the lower, upper or both triangle(s).

title A character string giving the main title for the plot and to which is appended the levels combination of the sectioning factors, if any, for each plot.

axis.labels A logical indicating whether a label is to be added to the x- and y-axes. If TRUE, the label is the comma-separated list of factors whose levels combinations are involved in the prediction differences for which the LSD values are calculated.

axis.text.size A numeric giving the size of the labels on the axes of the heatmap.

sep A character giving the characters separating the levels of different factors in the row and column names of the sed component.

colours A vector of of colours to be passed to the ggplot function scale\_colour\_gradientn.

ggplotFuncs A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. It is passed to ggplot via plotLSDerrors.data.frame to be applied in creating the ggplot object.

printPlot A logical indicating whether or not the a plot is to be printed. This would be used when just the returned data.frame is required.

sortFactor A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the
sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.

sortParallelToCombo
A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the supplied list is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.

sortNestingFactor
A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

sortOrder
A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It speci-

Value
A list with components named LSDresults and plots. The LSDresults component contains the data.frame with the columns Rows, Columns, LSDresults, sections1 and sections2. This data.frame is formed using the LSD and sed components of object and is used by plotLSDerrors.data.frame in producing the plots. The plots component contains a list of ggplot objects, one for each plot produced. Multiple plots are stored in the plots component if the sections argument is set and the plots are are named for the levels combinations of the sections.

Author(s)
Chris Brien

See Also
plotLSDerrors.alldiffs, plotLSDerrors.data.frame, plotLSDs.data.frame, exploreLSDs, sort.alldiffs, subset.alldiffs, ggplot
Examples

## Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18" & Benches != "3")

## Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
    random = ~ Benches:MainPlots,
    keep.order=TRUE, data=tmp)
current.asr <- as.asreml(current.asr, NULL, NULL)
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
    asreml.obj = current.asr, tables = "none",
    wald.tab = current.asr$wald.tab,
    present = c("Type","Species","Sources"))

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE)) {
    m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
        (1|Benches:MainPlots),
        data=na.omit(WaterRunoff.dat))
    TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
    TS.preds <- summary(TS.emm)
    den.df <- min(TS.preds$df, na.rm = TRUE)
    ## Modify TS.preds to be compatible with a predictions.frame
    TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
        se = "SE", interval.type = "CI",
        interval.names = c("lower.CL", "upper.CL"))

    ## Form an all.diffs object and check its validity
    els <- as.numeric(rownames(TS.preds))
    TS.vcov <- vcov(TS.emm)[els,els]
    TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
        vcov = TS.vcov, tdf = den.df)
    validAlldiffernts(TS.diffs)
}

## Plot LSD values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs")) {
    plotLSDerrors(TS.diffs, gridspacing = rep(c(3,4), c(4,2)))
    plotLSDerrors(TS.diffs, sections = "Sources", axis.labels = TRUE)
}
plotLSDerrors.data.frame

Plots a map of the supplied errors that occur in using the computed LSD values for pairwise differences between predictions.

Description

Produces a plot of the errors that have been supplied in a data.frame. The data.frame includes two factors whose levels specify, for each LSD result, which combinations of factor levels are being compared. The function plotLSDerrors.alldiffs produces such data.frames.

Usage

```r
## S3 method for class 'data.frame'
plotLSDerrors(object, LSDresults = "LSDresults", x, y,
               alpha = 0.05, triangles = "both",
               gridspacing = 0, title = NULL,
               axis.labels = NULL, axis.text.size = 12,
               colours = c("white","blue","red","grey"),
               ggplotFuncs = NULL, printPlot = TRUE, ...)
```

Arguments

- `object`: A data.frame containing the three columns specified by LSDresults, x and y.
- `LSDresults`: A character giving the name of the column in object that contains the LSD results values to be plotted. The column should be a character or factor with values or levels that are a subset of Ok, FN, FP and na.
- `x`: A character giving the name of the column in object that contains the factor whose levels index the LSD values that are to be plotted in the same column.
- `y`: A character giving the name of the column in object that contains the labels of the LSD values that are to be plotted as the rows.
- `alpha`: A numeric giving the significance level for the LSD.
- `triangles`: A character indicating whether the plot should include the lower, upper or both triangle(s). Here it is only used to adjust gridlines for the omission of the diagonal.
- `gridspacing`: A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. This is most useful when two or more factors index the rows and columns. If a single, nonzero number, k say, is given then a grid line is placed after every kth row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.
- `title`: A character string giving the main title for the plot.
- `axis.labels`: A character string giving the label to use for both the x- and y-axis.
- `axis.text.size`: A numeric giving the size of the labels on the axes of the heatmap.
- `colours`: A vector of of colours to be passed to the ggplot function scale_colour_gradientn.
- `ggplotFuncs`: A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. These functions are applied in creating the ggplot object.
plotLSDerrors.data.frame

printPlot

A logical indicating whether or not the a plot is to be printed. This would be used when just the returned ggplot object is required.

... 

Provision for passing arguments to functions called internally - not used at present.

Value

An object of class "ggplot", which can be plotted using print or otherwise manipulated.

Author(s)

Chris Brien

See Also

plotLSDs.data.frame, plotLSDs.all diffs, exploreLSDs, ggplot

Examples

##Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18")

##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
random = ~ Benches:MainPlots,
keep.order=TRUE, data= tmp))
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
asreml.obj = current.asr, tables = "none",
wald.tab = current.asrt$wald.tab,
present = c("Type","Species","Sources"))

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
requireNamespace("emmmeans", quietly = TRUE))
{
m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
(1|Benches:MainPlots),
data=na.omit(WaterRunoff.dat))
TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
se = "SE", interval.type = "CI",
interval.names = c("lower.CL", "upper.CL"))

## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
plotLSDs.alldiffs

```
TS.vcov <- vcov(TS.emm)[els,els]
TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
                         vcov = TS.vcov, tdf = den.df)
validAlldiffs(TS.diffs)
```

## Plot LSD values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs")) {
  LSDresults <- within(reshape2::melt(TS.diffs$p.differences),
                        {
                          Var1 <- factor(Var1, levels=dimnames(TS.diffs$p.differences)[[1]])
                          Var2 <- factor(Var2, levels=levels(Var1))
                        })
  names(LSDresults) <- c("Rows","Columns","LSDresults")
  plotLSDerrors(LSDresults, x = "Rows", y = "Columns", gridspacing = rep(c(3,4), c(4,2)))
}

---

**plotLSDs.alldiffs**  
Plots a heat map of computed LSD values for pairwise differences between predictions.

**Description**

Produces a heat-map plot of the computed LSD values for pairwise differences between predictions by multiplying the values stored in the `sed` component of an `all.diffs` object by the t-value for the `df` stored in the `tdf` attribute of the object. This component is generally a matrix whose rows and columns are labelled by the levels of one or more factors, the set of labels being the same for rows and columns. The `sections` argument allows multiple plots to be produced, one for each combination of the levels of the factors listed in `sections`. Otherwise, a single plot is produced for all observed combinations of the levels of the factors in the `classify` attribute for the `alldiffs.object`. The plots are produced using `plotLSDs.data.frame`. The order of plotting the levels of one of the factors indexing the predictions can be modified using `sort.alldiffs`.

**Usage**

```
plotLSDs(object, ...)  
## S3 method for class 'alldiffs'
plotLSDs(object, alpha = 0.05,
         sections = NULL, gridspacing = 0, factors.per.grid = 0,
         triangles = "both",
         title = NULL, axis.labels = TRUE, axis.text.size = 12,
         sep="", colours = RColorBrewer::brewer.pal(3, "Set2"),
         ggplotFuncs = NULL, printPlot = TRUE,
         sortFactor = NULL, sortParallelToCombo = NULL,
         sortNestingFactor = NULL, sortOrder = NULL,
         decreasing = FALSE, ...)
```

**Arguments**

- **object**  
  An `alldiffs.object` with an `sed` component that is not `NULL`.

- **alpha**  
  A numeric giving the significance level for the LSD.
sections  A character listing the names of the factors that are to be used to break the plot into sections. A separate plot will be produced for each observed combination of the levels of these factors.

gridspacing  A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. An alternative is to specify the factors.per.grid argument to have the grid spacings automatically calculated. Grids are most useful when two or more factors index the rows and columns. If a single, nonzero number, $k$ say, is given then a grid line is placed after every $k$th row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.

factors.per.grid  A numeric specifying the number of factors to include within each grid of differences. The gridspacing will then be computed based on the numbers of combinations observed within the levels of the remaining factors in a single plot. The gridspacing argument to this function will be ignored if factors.per.grid is greater than zero. Grids are most useful when two or more factors index the rows and columns of each plot.

triangles  A character indicating whether the plot should include the lower, upper or both triangles.

title  A character string giving the main title for the plot and to which is appended the levels combination of the sectioning factors, if any, for each plot.

axis.labels  A logical indicating whether a label is to be added to the x- and y-axes. If TRUE, the label is the comma-separated list of factors whose levels combinations are involved in the prediction differences for which the LSD values are calculated.

axis.text.size  A numeric giving the size of the labels on the axes of the heatmap.

sep  A character giving the characters separating the levels of different factors in the row and column names of the sed component.

colours  A vector of of colours to be passed to the ggplot function scale_color_gradientn.

ggplotFuncs  A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. It is passed to ggplot via plotLSDs.data.frame to be applied in creating the ggplot object.

printPlot  A logical indicating whether or not the a plot is to be printed. This would be used when just the returned data.frame is required.

sortFactor  A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.

sortParallelToCombo  A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the
supplied **list** is named for a **classify** variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of **sortFactor**. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If **sortParallelToCombo** is **NULL** then the first value of each **classify** variable, except for the **sortFactor** factor, in the predictions component is used to define **sortParallelToCombo**. If there is only one variable in the **classify** then **sortParallelToCombo** is ignored.

**sortNestingFactor**

A [character](https://en.wikipedia.org/wiki/Character_(computer)) containing the name of the factor that defines groups of the **sortFactor** within which the predicted values are to be ordered. If there is only one variable in the **classify** then **sortNestingFactor** is ignored.

**sortOrder**

A character vector whose length is the same as the number of levels for **sortFactor** in the predictions component of the **alldiffs.object**. It specifies the desired order of the levels in the reordered components of the **alldiffs.object**. The argument **sortParallelToCombo** is ignored.

The following creates a **sortOrder** vector **levs** for factor **f** based on the values in **x**:  

```r
levs <- levels(f)[order(x)]
```

**decreasing**

A logical passed to **order** that determines whether the order for sorting the **alldiffs.object** components is for increasing or decreasing magnitude of the predicted values.

... Provision for passing arguments to functions called internally - not used at present.

**Value**

A **list** with components named LSDs and plots. The LSDs component contains the data.frame with the columns **Rows**, **Columns**, **LSDs**, **sections1** and **sections2**. This data.frame is formed using the **sed** component of object and is used by **plotLSDs.data.frame** in producing the plot. The plots component contains a list of ggplot objects, one for each plot produced. Multiple plots are stored in the plots component if the **sections** argument is set and the plots are are named for the levels combinations of the sections.

**Author(s)**

Chris Brien

**See Also**

`plotLSDs.data.frame`, `exploreLSDs`, `sort.alldiffs`, `subset.alldiffs`, `ggplot`

**Examples**

```r
# Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18" & Benches != "3")

# Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) # required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
            random = ~ Benches:MainPlots,
```
keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
asreml.obj = current.asr, tables = "none",
wald.tab = current.asr$wald.tab,
present = c("Type","Species","Sources"))

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE)) {
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                           (1|Benches:MainPlots),
data=na.omit(WaterRunoff.dat))
  TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
  TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)

  ## Modify TS.preds to be compatible with a predictions.frame
  TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
                                   se = "SE", interval.type = "CI",
                                   interval.names = c("lower.CL", "upper.CL"))
}

## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els,els]
TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
                           vcov = TS.vcov, tdf = den.df)
validAlldiffs(TS.diffs)

## Plot LSD values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs")) {
  plotLSDs(TS.diffs, gridspacing = rep(c(3,4), c(4,2)))
  plotLSDs(TS.diffs, sections = "Sources", axis.labels = TRUE)
}

---

**plotLSDs.data.frame** *Plots a heat map of computed LSD-values for pairwise differences between predictions.*

**Description**

Produces a heat-map plot of the computed LSD values for pairwise differences between predictions that are stored in in a data.frame. The data.frame includes two factors whose levels specify, for each LSD value, which combinations of factor levels are being compared.
Usage

## S3 method for class 'data.frame'
plotLSDs(object, LSD = "LSDs", x, y, alpha = 0.05, triangles = "both", gridspacing = 0, title = NULL, axis.labels = NULL, axis.text.size = 12, colours = RColorBrewer::brewer.pal(3, "Set2"), ggplotFunucs = NULL, printPlot = TRUE, ...)

Arguments

object A data.frame containing the three columns specified by LSD, x and y.
LSD A character giving the name of the column in object that contains the LSD values to be plotted.
x A character giving the name of the column in object that contains the factor whose levels index the LSD values that are to be plotted in the same column.
y A character giving the name of the column in object that contains the labels of the LSD values that are to be plotted as the rows.
alpha A numeric giving the significance level for the LSD.
triangles A character indicating whether the plot should include the lower, upper or both triangle(s). Here it is only used to adjust gridlines for the omission of the diagonal.
gridspacing A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. This is most useful when two or more factors index the rows and columns. If a single, nonzero number, k say, is given then a grid line is placed after every kth row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.
title A character string giving the main title for the plot.
axis.labels A character string giving the label to use for both the x- and y-axis.
axis.text.size A numeric giving the size of the labels on the axes of the heatmap.
colours A vector of of colours to be passed to the ggplot function scale_colour_gradientn.
ggplotFunucs A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. These functions are applied in creating the ggplot object.
printPlot A logical indicating whether or not the a plot is to be printed. This would be used when just the returned ggplot object is required.
...

Value

An object of class "ggplot", which can be plotted using print or otherwise manipulated.

Author(s)

Chris Brien
See Also

plotLSDs.alldiffs, plotLSDerrors.alldiffs, plotLSDerrors.data.frame, exploreLSDs, gplot

Examples

##Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18")

##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                     random = ~ Benches:MainPlots,
                     keep.order = TRUE, data = tmp))
current.asr <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
                               asreml.obj = current.asr, tables = "none",
                               wald.tab = current.asr$wald.tab,
                               present = c("Type", "Species", "Sources"))

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE)) {
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                           (1|Benches:MainPlots),
                           data = na.omit(WaterRunoff.dat))
  TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
  TS.preds <- summary(TS.emm)
  den.df <- min(TS.preds$df, na.rm = TRUE)

  ## Modify TS.preds to be compatible with a predictions.frame
  TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
                                  se = "SE", interval.type = "CI",
                                  interval.names = c("lower.CL", "upper.CL"))

  ## Form an all.diffs object and check its validity
  els <- as.numeric(rownames(TS.preds))
  TS.vcov <- vcov(TS.emm)[els, els]
  TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
                              vcov = TS.vcov, tdf = den.df)
  validAlldiffs(TS.diffs)
}

## Plot LSD values for predictions obtained using asreml or lmerTest

if (exists("TS.diffs")) {
  LSD <- within(reshape2::melt(TS.diffs$p.differences),
                {Var1 <- factor(Var1, levels = dimnames(TS.diffs$p.differences)[[1]])
                 Var2 <- factor(Var2, levels = levels(Var1))})
plotPredictions.data.frame

```r
)}
names(LSD) <- c("Rows","Columns","LSDs")
plotLSDs(LSD, x = "Rows", y = "Columns", gridspacing = rep(c(3,4), c(4,2)))
```

---

**plotPredictions.data.frame**

*Plots the predictions for a term, possibly with error bars.*

### Description

This function plots the predictions \( y \) that are based on \texttt{classify} and stored in the \texttt{data.frame} data. The package \texttt{ggplot2} is used to produce the plots. Line plots are produced when variables involving \texttt{x.num} or \texttt{x.fac} are involved in \texttt{classify} for the predictions; otherwise, bar charts are produced. Further, for line charts, the argument \texttt{panels} determines whether a single plot or multiple plots in a single window are produced; for bar charts, the argument \texttt{panels} is ignored.

### Usage

```r
## S3 method for class 'data.frame'
plotPredictions(data, classify, y,
    x.num = NULL, x.fac = NULL, nonx.fac.order = NULL,
    colour.scheme = "colour", panels = "multiple",
    graphics.device = NULL,
    error.intervals = "Confidence", interval.annotate = TRUE,
    titles = NULL, y.title = NULL,
    filestem = NULL, ggplotFuncs = NULL, 
    ...)
```

### Arguments

- **data**: A \texttt{predictions.frame}, or \texttt{data.frame}, containing the values of the variables to be plotted. Generally, it should contain the variables classifying the predictions and include a column with the name specified in the \texttt{y} argument, usually \texttt{predicted.value} or \texttt{backtransformed.predictions}; each row contains a single predicted value. It should also include columns for the \texttt{standard.error} and \texttt{est.status}. The number of rows should equal the number of unique combinations of the classifying variables. While such a \texttt{data.frame} can be constructed from the beginning, the \texttt{pvals} component of the value produced by \texttt{predict.asreml} is a suitable value to supply for this argument. Note that the names \texttt{standard.error} and \texttt{est.status} have been changed to \texttt{std.error} and \texttt{status} in the \texttt{pvals} component produced by \texttt{asreml-R4}; if the new names are in the \texttt{data.frame} supplied to \texttt{predictions}, they will be returned to the previous names.

- **If** \texttt{error.intervals} is not "none", then the predictions component and, if present, the backtransforms component should contain columns for the lower and upper values of the limits for the interval with names that begin with \texttt{lower} and \texttt{upper}, respectively. The second part of the name must be one of \texttt{Confidence}, \texttt{StandardError} or \texttt{halfLeastSignificant}. The last part needs to be consistent between the lower and upper limits.
classify  A character string giving the combinations of the independent variables on which the predictions are based. It is an interaction type term formed from the independent variables, that is, separating the variable names with the `:` operator. To predict the overall mean, set the `classify` to "(Intercept)".

y  A character string giving the name of the variable that is to be plotted on the Y axis.

x.num  A character string giving the name of the numeric covariate that corresponds to `x.fac`, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of unique values as the number of levels in `x.fac`.

x.fac  A character string giving the name of the factor that corresponds to `x.num`, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of levels as the number of unique values in `x.num`. The levels of `x.fac` must be in the order in which they are to be plotted - if they are dates, then they should be in the form yyyymmdd, which can be achieved using `as.Date`. However, the levels can be non-numeric in nature, provided that `x.num` is also set.

nonx.fac.order  A character vector giving the order in which factors other than `x.fac` are to be plotted in facetted plots (i.e. where the number of non x factors is greater than 1). The first factor in the vector will be plotted on the X axis (if there is no `x.num` or `x.fac`). Otherwise, the order of plotting the factors is in columns (X facets) and then rows (Y facets). By default the order is in decreasing order for the numbers of levels of the non x factors.

colour.scheme  A character string specifying the colour scheme for the plots. The default is "colour" which produces coloured lines and bars, a grey background and white gridlines. A value of "black" results in black lines, grey bars and gridlines and a white background.

panels  Possible values are "single" and "multiple". When line plots are to be produced, because variables involving `x.num` or `x.fac` are involved in `classify` for the predictions, `panels` determines whether or not a single panel or multiple panels in a single window are produced. The `panels` argument is ignored for bar charts.

graphics.device  A character specifying a graphics device for plotting. The default is `graphics.device = NULL`, which will result in plots being produced on the current graphics device. Setting it to "windows", for example, will result in a windows graphics device being opened.

error.intervals  A character string indicating the type of error interval, if any, to plot in order to indicate uncertainty in the results. Possible values are "none", "StandardError", "Confidence" and "halfLeastSignificant". Here, any option other than "none" will result in the interval limits contained in `data` being plotted.

interval.annotate  A logical indicating whether the plot annotation indicating the type of `error.interval` is to be included in the plot.

titles  A list, each component of which is named for a column in the `data.frame` for the `asreml.obj` used in making the predictions and contains a character string giving a title to use in output (e.g. tables and graphs). Here they will be used for axis labels for nonresponse variables. For response variable labels see `y.title`.
filestem  A character string giving the beginning of the name of the file in which to save the plot. If filestem = NULL, the plot is not saved. The remainder of the file name will be generated automatically and consists of the following elements separated by full stops: the classify term, Bar or Line and, if error.intervals is not "none", one of SE, CI or LSI. The file will be saved as a 'png' file in the current work directory.

y.title  The title to be displayed on the y axis of any plot.

ggplotFuncs  A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. These functions are applied in creating the ggplot object for plotting.

...  further arguments passed to ggplot.

Value  no values are returned.

Author(s)  Chris Brien

See Also  
allDifferences.data.frame, predictPresent.asreml, redoErrorIntervals.alldiffs, recalcLSD.alldiffs, ggplot, Devices

Examples

## Not run:
data(WaterRunoff.dat)
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = log.Turbidity ~ Benches + Sources + Type + Species + 
Sources:Type + Sources:Species + 
Sources:xDay + Species:xDay + Species:Date, 
data = WaterRunoff.dat, keep.order = TRUE)
current.asrt <- as.asrtests(current.asr, NULL, NULL)

#### Get the observed combinations of the factors and variables in classify
class.facs <- c("Species","Date","xDay")
levs <- as.data.frame(table(WaterRunoff.dat[class.facs]))
levs <- as.list(levs[levs$Freq != 0, class.facs])
levs$xDay <- as.numfac(levs$xDay)
predictions <- predict(current.asr, classify="Species:Date:xDay", 
parallel = TRUE, levels = levs, 
present = c("Type","Species","Sources"))

#### for asreml-R3
predictions <- predictions$predictions$pvals
predictions <- predictions[predictions$est.status == "Estimable",]

#### for asreml-R4
predictions <- predictions$pvals
predictions <- predictions[predictions$status == "Estimable",]

#### end
plotPredictions(classify="Species:Date:xDay", y = "predicted.value", 
data = predictions,
plotPvalues.alldiffs

Plots a heat map of p-values for pairwise differences between predictions.

Description

Produces a heat-map plot of the p-values for pairwise differences between predictions that is stored in the p.differences component of an all.diffs object. This is generally a matrix whose rows and columns are labelled by the levels of one or more factors, the set of labels being the same for rows and columns. The sections argument allows multiple plots to be produced, one for each combination of the levels of the factors listed in sections. Otherwise, a single plot is produced for all observed combinations of the levels of the factors in the classify for the alldiffs.object.
The plots are produced using `plotPvalues.data.frame`. The order of plotting the levels of one of the factors indexing the predictions can be modified using `sort.alldiffs`.

**Usage**

```r
plotPvalues(object, ...)  
## S3 method for class 'alldiffs'
plotPvalues(object, sections = NULL,  
gridspacing = 0, factors.per.grid = 0,  
show.sig = FALSE, alpha = 0.10,  
sig.size = 3, sig.colour = "black",  
sig.face = "plain", sig.family = "",  
triangles = "both",  
title = NULL, axis.labels = TRUE, axis.text.size = 12,  
seps="", colours = RColorBrewer::brewer.pal(3, "Set2"),  
ggplotFuncs = NULL, printPlot = TRUE,  
sortFactor = NULL, sortParallelToCombo = NULL,  
sortNestingFactor = NULL, sortOrder = NULL,  
decreasing = FALSE, ...)
```

**Arguments**

- **object**: An `alldiffs.object` with a `p.differences` component that is not `NULL`.
- **sections**: A character listing the names of the factors that are to be used to break the plot into sections. A separate plot will be produced for each observed combination of the levels of these factors.
- **gridspacing**: A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. An alternative is to specify the `factors.per.grid` argument to have the grid spacings automatically calculated. Grids are most useful when two or more factors index the rows and columns. If a single, nonzero number, \( k \) say, is given then a grid line is placed after every \( k \)th row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.
- **factors.per.grid**: A numeric specifying the number of factors to include within each grid of differences. The gridspacing will then be computed based on the numbers of combinations observed within the levels of the remaining factors in a single plot. The gridspacing argument to this function will be ignored if `factors.per.grid` is greater than zero. Grids are most useful when two or more factors index the rows and columns of each plot.
- **show.sig**: A logical that specifies whether asterisks indicating the level of significance are to be added to the plot. If they are then `"***"` indicates that \( p \leq 0.001 \), `"**"` that \( 0.001 < p \leq 0.01 \), `"*"` that \( 0.01 < p \leq 0.05 \) and `"."` that \( 0.05 < p \leq 0.10 \). The last is only included for \( \alpha = 0.10 \).
- **alpha**: A numeric giving the significance level for testing pairwise differences; must be 0.05 or 0.10.
- **sig.size**: A numeric specifying the size, in pts, of the significance asterisks.
- **sig.colour**: A character specifying the colour to use for the significance asterisks.
- **sig.face**: A character specifying the font face for the significance asterisks ("plain", "italic", "bold", "bold.italic").
**sig.family** A character specifying the font family for the significance asterisks. The font families that are available depends on the system. For font families other than the basic Postscript fonts, see the extrafont package.

**triangles** A character indicating whether the plot should include the lower, upper or both triangle(s).

**title** A character string giving the main title for the plot and to which is appended the levels combination of the sectioning factors, if any, for each plot.

**axis.labels** A logical indicating whether a label is to be added to the x- and y-axes. If TRUE, the label is the comma-separated list of factors whose levels combinations are involved in the prediction differences for which the p-values are calculated.

**axis.text.size** A numeric giving the size of the labels on the axes of the heatmap.

**sep** A character giving the characters separating the levels of different factors in the row and column names of the p.differences component.

**colours** A vector of of colours to be passed to the ggplot function scale\_colour\_gradientn.

**ggplotFuncs** A list, each element of which contains the results of evaluating a ggplot function. It is created by calling the list function with a ggplot function call for each element. It is passed to ggplot via plotPvalues.data.frame to be applied in creating the ggplot object.

**printPlot** A logical indicating whether or not the a plot is to be printed. This would be used when just the returned data.frame is required.

**sortFactor** A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.

**sortParallelToCombo** A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the supplied list is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.

**sortNestingFactor** A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

**sortOrder** A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the desired order of the levels in the reordered components of the alldiffs.object. The argument sortParallelToCombo is ignored.
The following creates a `sortOrder` vector `levs` for factor `f` based on the values in `x`: `levs <- levels(f)[order(x)]`.

A logical passed to `order` that determines whether the order for sorting the `alldiffs.object` components is for increasing or decreasing magnitude of the predicted values.

Provision for passing arguments to functions called internally - not used at present.

Value

A `list` with components named `pvalues` and `plots`. The `pvalues` component contains the `data.frame` with the columns `Rows`, `Columns`, `p`, `sections1` and `sections2`. This data.frame is formed using the `sed` component of `object` and is used by `plotPvalues.data.frame` in producing the plot. The `plots` component contains a list of `ggplot` objects, one for each plot produced. Multiple plots are stored in the `plots` component if the `sections` argument is set and the plots are are named for the levels combinations of the sections.

Author(s)

Chris Brien

See Also

`plotPvalues.data.frame`, `allDifferences.data.frame`, `sort.alldiffs`, `subset.alldiffs`, `ggplot`

Examples

```r
##Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18" & Benches != "3")

##Use asreml to get predictions and associated statistics
## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                   random = ~ Benches:MainPlots,
                   keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
                                asreml.obj = current.asr, tables = "none",
                                wald.tab = current.asrt$wald.tab,
                                present = c("Type","Species","Sources"))

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                            (1|Benches:MainPlots),
                            data=na.omit(WaterRunoff.dat))
```
TS.emm <- emmeans::emmeans(ml1.lmer, specs = ~ Sources:Type)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
    se = "SE", interval.type = "CI",
    interval.names = c("lower.CL", "upper.CL"))

## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els, els]
TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
    vcov = TS.vcov, tdf = den.df)
validAll diffs(TS.diffs)
}

## Plot p-values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs")) {
    plotPvalues(TS.diffs, gridspacing = rep(c(3,4), c(4,2)), show.sig = TRUE)
    plotPvalues(TS.diffs, sections = "Sources", show.sig = TRUE, axis.labels = TRUE)
}
```

---

**plotPvalues.data.frame**

*Plots a heat map of p-values for pairwise differences between predictions.*

### Description

Produces a heat-map plot of the p-values for pairwise differences between predictions that is in a data.frame. The data.frame includes two factors whose levels specify, for each p-value, which factor levels are being compared.

### Usage

```
## S3 method for class 'data.frame'
plotPvalues(object, p = "p", x, y,
    gridspacing = 0, show.sig = FALSE, alpha = 0.10,
    sig.size = 3, sig.colour = "black",
    sig.face = "plain", sig.family = "",
    triangles = "both",
    title = NULL, axis.labels = NULL, axis.text.size = 12,
    colours = RColorBrewer::brewer.pal(3, "Set2"),
    ggplotFuncs = NULL, printPlot = TRUE, ...)
```

### Arguments

- **object**: A data.frame containing the three columns specified by p, x and y.
- **p**: A character giving the name of the column in object that contains the p-values to be plotted.
A character giving the name of the column in `object` that contains the factor whose levels index the p-values that are to be plotted in the same column.

A character giving the name of the column in `object` that contains the labels of the p-values that are to be plotted as the rows.

A numeric specifying the number(s) of rows and columns that form groups in the grid of differences. This is most useful when two or more factors index the rows and columns. If a single, nonzero number, \( k \) say, is given then a grid line is placed after every \( k \)th row and column. If a vector of values is given then the number of grid lines is the length of the vector and the spacing between each is specified by the elements of the vector.

A logical that specifies whether asterisks indicating the level of significance are to be added to the plot. If they are then '***' indicates that \( p \leq 0.001 \), '**' that \( 0.001 < p \leq 0.01 \), '*' that \( 0.01 < p \leq 0.05 \) '.' that \( 0.05 < p \leq 0.10 \). The last is only included for \( \alpha = 0.10 \).

A numeric giving the significance level for testing pairwise differences; must be 0.05 or 0.10.

A numeric specifying the size, in pts, of the significance asterisks.

A character specifying the colour to use for the significance asterisks.

A character specifying the font face for the significance asterisks ("plain", "italic", "bold", "bold.italic").

A character specifying the font family for the significance asterisks. The font families that are available depends on the system. For font families other than the basic Postscript fonts, see the `extrafont` package.

A character indicating whether the plot should include the lower, upper or both triangle(s). Here it is only used to adjust gridlines for the omission of the diagonal.

A character string giving the main title for the plot.

A character string giving the label to use for both the x- and y-axis.

A numeric giving the size of the labels on the axes of the heatmap.

A vector of of colours to be passed to the `ggplot` function `scale_colour_gradientn`.

A list, each element of which contains the results of evaluating a `ggplot` function. It is created by calling the `list` function with a `ggplot` function call for each element. These functions are applied in creating the `ggplot` object.

A logical indicating whether or not the a plot is to be printed. This would be used when just the returned `ggplot` object is required.

Provision for passing arguments to functions called internally - not used at present.

An object of class "`ggplot`", which can be plotted using `print` or otherwise manipulated.

Chris Brien

`plotPvalues.alldiffs, allDifferences.data.frame, ggplot`
Examples

```r
##Subset WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18")

##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
random = ~ Benches:MainPlots,
keep.order=TRUE, data= tmp))
current.asrt <- as.asrtests(current.asr, NULL, NULL)
SS.diffs <- predictPlus.asreml(classify = "Sources:Type",
asreml.obj = current.asr, tables = "none",
wald.tab = current.asrt$wald.tab,
present = c("Type","Species","Sources"))

## Use lmerTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
(1|Benches:MainPlots),
data=na.omit(WaterRunoff.dat))
  TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
  TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)

## Modify TS.preds to be compatible with a predictions.frame
  TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
  se = "SE", interval.type = "CI",
  interval.names = c("lower.CL", "upper.CL"))

## Form an all.diffs object and check its validity
  els <- as.numeric(rownames(TS.preds))
  TS.vcov <- vcov(TS.emm)[els,els]
  TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Type",
  vcov = TS.vcov, tdf = den.df)
  validAllDiffs(TS.diffs)
}

## Plot p-values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs"))
{
  p <- within(reshape2::melt(TS.diffs$p.differences),
  {Var1 <- factor(Var1, levels=dimnames(TS.diffs$p.differences)[[1]])
  Var2 <- factor(Var2, levels=levels(Var1))
  })
  names(p) <- c("Rows","Columns","p")
  plotPvalues(p, x = "Rows", y = "Columns",
  gridspacing = rep(c(3,4), c(4,2)), show.sig = TRUE)
}
```

Plots empirical variogram faces, including envelopes, from supplied residuals as described by Stefanova, Smith & Cullis (2009).

Usage

```r
## S3 method for class 'data.frame'
plotVariofaces(data, residuals, restype="Residuals", ...)
```

Arguments

- **data**: A `data.frame` with either 3 or 4 columns. Only if there are 4 columns, the first should be a factor indexing sections for which separate variogram plots are to be produced. In either case, the other 3 columns should be, in order, (i) a factor indexing the x-direction, (ii) a factor indexing the y-direction, and (iii) the residuals for the observed response.
- **residuals**: A `data.frame`, with either 2 or 3 initial columns followed by columns, each of which are the residuals from a simulated data set.
- **restype**: A character describing the type of residuals that have been supplied. It will be used in the plot titles.
- **...**: Other arguments that are passed down to the function `asreml.variogram`.

Details

For each set of residuals, `asreml.variogram` is used to obtain the empirical variogram, from which the values for its faces are obtained. Plots are produced for each face and include the observed residuals and the 2.5%, 50% & 97.5% quantiles.

Value

A list with the following components:

1. **face1**: A `data.frame` containing the variogram values on which the plot for the first dimension is based.
2. **face2**: A `data.frame` containing the variogram values on which the plot for the second dimension is based.
powerTransform

Performs a combination of a linear and a power transformation on a variable. The transformed variable is stored in the data.frame data.

Author(s)

Chris Brien

References


See Also

asremlPlus-package, asreml, asreml.variogram, variofaces.asreml, simulate.asreml.

Examples

## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
  random = ~ Row + Column + units,
  residual = ~ ar1(Row):ar1(Column),
  data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary.asrtests(current.asrt)
# Form variance matrix based on estimated variance parameters
s2 <- current.asr$sigma2
gamma.Row <- current.asr$gammas[1]
gamma.unit <- current.asr$gammas[2]
rho.r <- current.asr$gammas[4]
rho.c <- current.asr$gammas[5]
row.ar1 <- mat.ar1(order=10, rho=rho.r)
col.ar1 <- mat.ar1(order=15, rho=rho.c)
V <- gamma.Row * fac.sumop(Wheat.dat$Row) +
gamma.unit * diag(1, nrow=150, ncol=150) +
mat.dirprod(col.ar1, row.ar1)
V <- s2*V

#Produce variogram faces plot (Stefanova et al, 2009)
resid <- simulate(current.asr, V=V, which="residuals")
resid$residuals <- cbind(resid$observed[c("Row","Column")],
  resid$residuals)
plotVariofaces(data=resid$observed[C("Row","Column","residuals")],
  residuals=resid$residuals,
  retype="Standardized conditional residuals")

## End(Not run)
**powerTransform**

**Description**

Perform a combination of a linear and a power transformation on a variable whose name is given as a character string in `var.name`. The transformed variable is stored in the `data.frame` data. The name of the transformed variable is made by prepending to the original `var.name` a combination of (i) offset, if offset is nonzero, (ii) `neg.` if scale is -1, or scaled., if abs(scale) is other than one, and (iii) either `log.`, `sqrt.`, `recip.` or `power.`, if power is other than one. No action is taken if there is no transformation (i.e. offset = 0, scale = 1 and power = 1). Also, the titles list is extended to include a component with a generated title for the transformed variable with text indicating the transformation prepended to the title for the `var.name` obtained from the titles list. For nonzero offset, ‘Offset ’ is prepended. For scaled not equal to one, the possible prepends are ‘Negative of ’ and ‘Scaled’. The possible prepended texts for power not equal to one are ‘Logarithm of’, ‘Square root of’, ‘Reciprocal of ’ and ‘Power nnnn of ’, where nnn is the power used.

**Usage**

```r
powerTransform(var.name, power = 1, offset = 0, scale = 1, titles = NULL, data)
```

**Arguments**

- `var.name`: A character string specifying the name of the variable in the `data.frame` data that is to be transformed.
- `power`: A number specifying the power to be used in the transformation. If equal to 1, the default, no power transformation is applied. Otherwise, the variable is raised to the specified power, after scaling and applying any nonzero offset. If power = 0, the natural logarithm is used to transform the response; however, if the smallest value to be log-transformed is less than 1e-04, an error is generated. A log-transformation in this situation may be possible if a nonzero offset and/or a scale not equal to one is used.
- `offset`: A number to be added to each value of the variable, after any scaling and before applying any power transformation.
- `scale`: A number to multiply each value of the variable, before adding any offset and applying any power transformation.
- `titles`: A character vector, each element of which is named for a variable in `data` and is a character string giving a title to use in output (e.g. tables and graphs) involving the variable. If titles are not supplied, the column name of the variable in `data` is used.
- `data`: A `data.frame` containing the variable to be transformed and to which the transformed variable is to be appended.

**Value**

A list with a component named `data` that is the `data.frame` containing the transformed variable, a component named `tvar.name` that is a character string that is the name of the transformed variable in `data`, and a component named `titles` that extends the list supplied in the `titles` argument to include a generated title for the transformed title, the name of the new component being `tvar.name`.

**Author(s)**

Chris Brien
See Also

angular, angular.mod.

Examples

```r
## set up a factor with labels
x.dat <- data.frame(y = c(14, 42, 120, 150))

## transform y to logarithms
trans <- powerTransform("y", power = 0, titles=list(y = "Length (cm)"), data = x.dat)
x.dat <- trans$data
tvar.name <- trans$tvar.name

## transform y to logarithms after multiplying by -1 and adding 1.
z.dat <- data.frame(y = c(-5.25, -4.29, -1.22, 0.05))
trans <- powerTransform("y", power = 0, scale = -1, offset = 1,
titles=list(y = "Potential"), data = z.dat)
z.dat <- trans$data
tvar.name <- trans$tvar.name
```

predictions.frame

Description of a predictions object

Description

A data.frame of S3-class predictions.frame that stores the predictions for a fitted model.

as.predictions.frame is function that converts a data.frame to an object of this class.

is.predictions.frame is the membership function for this class; it tests that an object has class predictions.frame.

validPredictionsFrame can be used to test the validity of a predictions.frame.

Value

A data.frame that begins with the variables classifying the predictions, in the same order as in the classify, followed by a column of predictions that is named either predicted.value or backtransformed.predictions; it also contains columns named standard.error and est.status. The number of rows should equal the number of unique combinations of the classifying variables. While such a data.frame can be constructed from the beginning, the pvals component of the value produced by predict.asreml is a suitable value to supply for this argument. Note that the names standard.error and est.status have been changed to std.error and status in the pvals component produced by asreml-R4; if the new names are in the data.frame supplied to predictions, they will be returned to the previous names.

The data.frame may also include columns for the lower and upper values of error intervals, either standard error, confidence or half-LSD intervals. The names of these columns will consist of three parts separated by full stops: 1) the first part will be lower or upper; 2) the second part will be one of Confidence, StandardError or halfLeastSignificant; 3) the third component will be limits.

IF accuracy.threshold is set to a numeric value at the time the prediction.frame is formed, it will also include a column logical values named LSDwarning.
When halfLeastSignificant limits have been included in a `predictions.frame`, its attributes will include those that are not NULL of `LSDtype`, `LSDby`, `LSDstatistic`, `LSDaccuracy` and `LSDvalues`. `LSDvalues` are the LSD values used to calculate the halfLeastSignificant error.intervals and are an expanded version of the values stored in the assigned.LSD column of the `LSD.frame`. See `predictPlus.asreml` for more information.

Author(s)

Chris Brien

See Also

`predictPlus.asreml`, `is.predictions.frame`, `as.predictions.frame`, `validPredictionsFrame`

Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics

## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety, random=~Blocks/Wplots, data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety", sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- as.predictions.frame(Var.pred$pvals, se = "std.error", est.status = "status")

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots), data=Oats.dat)
  Var.emm <- emmeans::emmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean", se = "SE", interval.type = "CI", interval.names = c("lower.CL", "upper.CL"))
}
if (exists("Var.preds"))
{
  ## Check the class and validity of the alldiffs object
  is.predictions.frame(Var.preds)
  validPredictionsFrame(Var.preds)
}
predictPlus.asreml Forms the predictions for a term, their pairwise differences and associated statistics. A factor having parallel values may occur in the model and a linear transformation of the predictions can be specified. It results in an object of class alldiffs.

Description

This function forms the predictions for `term` using `classify` and the supplied `asreml` object and stores them in an `alldiffs.object`. If `x.num` is supplied, the predictions will be obtained for the values supplied in `x.pred.values` and, if supplied, `x.plot.values` will replace them in the `alldiffs.object` that is returned. If `x.fac`, but not `x.num`, is specified, predictions will involve it and, if supplied, `x.plot.values` will replace the levels of `x.fac` in the `alldiffs.object` that is returned. In order to get the correct predictions you may need to supply additional arguments to `predict.asreml` through ... e.g. present, parallel, levels. Any aliased predictions will be removed, as will any standard error of pairwise differences involving them.

Also calculated are the approximate degrees of freedom of the standard errors of the predictions. If the denominator degrees of freedom for `term` are available in `wald.tab`, they are used. Otherwise the residual degrees of freedom or the maximum of the denominator degrees in `wald.tab`, excluding the Intercept, are used. Which is used depends on the setting of `dDF.na`. These degrees of freedom are used for the t-distribution on which p-values and confidence intervals are based. It is stored as an attribute to the `alldiffs.object`. The degrees of freedom are also used in calculating the minimum, mean and maximum LSD for comparing pairs of predictions, which are also stored in the `alldiffs.object`.

If `pairwise = TRUE`, all pairwise differences between the predictions, their standard errors, p-values and LSD statistics are computed using `allDifferences.data.frame`. This adds them to the `alldiffs.object` as additional list components named `differences`, `sed`, `p.differences` and `LSD`.

If a linear transformation of the predictions is specified then the values of this linear transformation are returned, instead of the original predictions, along with their standard errors and the pairwise differences and associated statistics.

If a transformation has been applied in the analysis (any one of `transform.power` is not one, `scale` is not one and `offset` is nonzero), the backtransforms of the transformed values and their lower and upper error intervals are added to a `data.frame` that is consistent with the predictions `data.frame`. If `transform.power` is other than one, the `standard.error` column of the `data.frame` is set to NA. This `data.frame` is added to the `alldiffs.object` as a list component called backtransforms.

The printing of the components produced is controlled by the `tables` argument. The order of plotting the levels of one of the factors indexing the predictions can be modified and is achieved using `sort.alldiffs`.

Usage

```r
## S3 method for class 'asreml'
predictPlus(asreml.obj, classify, term = NULL,
            inestimable.rm = TRUE, linear.transformation = NULL,
            error.intervals = "Confidence", alpha = 0.05,
            wald.tab = NULL, dDF.na = "residual", dDF.values = NULL,
            pairwise = TRUE, Vmatrix = FALSE,
```

predictPlus.asreml

Forms the predictions for a term, their pairwise differences and associated statistics. A factor having parallel values may occur in the model and a linear transformation of the predictions can be specified. It results in an object of class alldiffs.
asreml.obj asreml object for a fitted model.

classify A character string giving the variables that define the margins of the multi-way table to be predicted. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator. To predict the overall mean, set the classify to "(Intercept)".

term A character string giving the variables that define the term that was fitted using asreml and that corresponds to classify. It only needs to be specified when it is different to classify; it is stored as an attribute of the alldiffs.object. It is likely to be needed when the fitted model includes terms that involve both a numeric covariate and a factor that parallel each other; the classify would include the covariate and the term would include the factor.

inestimable.rm A logical indicating whether rows for predictions that are not estimable are to be removed from the components of the alldiffs.object.

linear.transformation A formula or a matrix. If a formula is given then it is taken to be a submodel of a model term corresponding to the classify. The projection matrix that transforms the predictions so that they conform to the submodel is obtained; the submodel does not have to involve variables in the classify, but the variables must be columns in the predictions component of alldiffs.obj and the space for the submodel must be a subspace of the space for the term specified by the classify. For example, for classify set to "A:B" the submodel ~ A + B will result in the predictions for the combinations of A and B being made additive for the factors A and B. The submodel space corresponding to A + B is a subspace of the space A:B. In this case both the submodel and the the classify involve only the factors A and B. To fit an intercept-only submodel, specify linear.transformation to be the formula ~1.
If a matrix is provided then it will be used to apply the linear transformation to the predictions. It might be a contrast matrix or a matrix of weights for a factor used to obtain the weighted average over that factor. The number of rows in the matrix should equal the number of linear combinations of the predictions desired and the number of columns should equal the number of predictions.
In either case, as well as the values of the linear combinations, their standard errors, pairwise differences and associated statistics are returned.

error.intervals A character string indicating the type of error interval, if any, to calculate in order to indicate uncertainty in the results. Possible values are "none", "none".
"StandardError", "Confidence" and "halfLeastSignificant". The default is for confidence limits to be used. The "halfLeastSignificant" option results in half the Least Significant Difference (LSD) being added and subtracted to the predictions, the LSD being calculated using the square root of the mean of the variances of all or a subset of pairwise differences between the predictions. If the LSD is zero, as can happen when predictions are constrained to be equal, then the limits of the error intervals are set to NA. If LSDtype is set to overall, the avsed.tolerance is not NA and the range of the SEDs divided by the average of the SEDs exceeds avsed.tolerance then the error.intervals calculations and the plotting will revert to confidence intervals.

alpha
A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

wald.tab
A data.frame containing the pseudo-anova table for the fixed terms produced by a call to wald.asreml. The main use of it here is in determining the degrees of freedom for calculating confidence or half-LSD error.intervals and p-values, the latter to be stored in the p.differences component of the alldiffs.object that is created.

dDF.na
A character specifying the method to use to obtain approximate denominator degrees of freedom. when the numeric or algebraic methods produce an NA. Consistent with when no denDF are available, the default is "residual" and so the residual degrees of freedom from asreml.obj$nedf are used. If dDF.na = "none", no substitute denominator degrees of freedom are employed; if dDF.na = "maximum", the maximum of those denDF that are available, excluding that for the Intercept, is used; if all denDF are NA, asreml.obj$nedf is used. If dDF.na = "supplied", a vector of values for the denominator degrees of freedom is to be supplied in dDF.values. Any other setting is ignored and a warning message produced. Generally, substituting these degrees of freedom is anticonservative in that it is likely that the degrees of freedom used will be too large.

dDF.values
A vector of values to be used when dDF.na = "supplied". Its values will be used when denDF in a test for a fixed effect is NA. This vector must be the same length as the number of fixed terms, including (Intercept) whose value could be NA.

pairwise
A logical indicating whether all pairwise differences of the predictions and their standard errors and p-values are to be computed and stored. If tables is equal to "differences" or "all" or error.intervals is equal to "halfLeastSignificant", they will be stored irrespective of the value of pairwise.

Vmatrix
A logical indicating whether the variance matrix of the predictions will be stored as a component of the alldiffs.object that is returned. If linear.transformation is set, it will be stored irrespective of the value of Vmatrix.

avsed.tolerance
A numeric giving the value of the SED range, the range of the SEDs divided by the square root of the mean of the variances of all or a subset of the pairwise differences, that is considered reasonable in calculating error.intervals. To have it ignored, set it to NA. It should be a value between 0 and 1. The following rules apply:

1. If avsed.tolerance is NA then mean LSDs of the type specified by LSDtype are calculated and used in error.intervals and plots.
2. Irrespective of the setting of LSDtype, if avsed.tolerance is not exceeded then the mean LSDs are used in error.intervals and plots.
3. If LSDtype is set to overall, avsed.tolerance is not NA, and avsed.tolerance is exceeded then error.intervals and plotting revert to confidence intervals.

4. If LSDtype is set to factor.combinations and avsed.tolerance is not exceeded for any factor combination then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

5. If LSDtype is set to per.prediction and avsed.tolerance is not exceeded for any prediction then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

accuracy.threshold
A numeric specifying the value of the LSD accuracy measure, which measure is specified by LSDaccuracy, as a threshold value in determining whether the halfLeastSignificant error.interval for a predicted value is a reasonable approximation; this will be the case if the LSDs across all pairwise comparisons for which the interval's LSD was computed, as specified by LSDtype and LSDby, are similar enough to the interval's LSD, as measured by LSDaccuracy. If it is NA, it will be ignored. If it is not NA, a column of logicals named LSDwarning will be added to the predictions component of the alldiffs.object. The value of LSDwarning for a predicted.value will be TRUE if the value of the LSDaccuracy measure computed from the LSDs for differences between this predicted.value and the other predicted.values as compared to its assignedLSD exceeds the value of accuracy.threshold. Otherwise, the value of LSDwarning for a predicted.value will be FALSE.

LSDtype
A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparisons, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations. See LSD.frame for further information on the values in a row of this data.frame and how they are calculated.

LSDsupplied
A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function dae::fac.combine to the predictions component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into assignedLSD column of the LSD.frame stored as the LSD component of the alldiffs.object.
LSDbys A character (vector) of variables names, being the names of the factors or numerics in the classify; for each combination of their levels and values, there will be or is a row in the LSD.frame stored in the LSD component of the alldiffs.object when LSDtype is factor.combinatons.

LSDstatistic A character nominating one or more of minmum, q10, q25, mean, median, q75, q90 or maximum as the value(s) to be stored in the assignedLSD column in an LSD.frame; the values in the assignedLSD column are used in computing halfLeastSignificant error.intervals. Here q10, q25, q75 and q90 indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function quantile is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median is calculated using the median function. Multiple values are only produced for LSDtype set to factor.combination, in which case LSDby must not be NULL and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is NULL, it is reset to mean.

LSDaccuracy A character nominating one of maxAbsDeviation, maxDeviation, q90Deviation or RootMeanSqDeviation as the statistic to be calculated as a measure of the accuracy of assignedLSD. The option q90Deviation produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named accuracyLSD in an LSD.frame.

titles A list, each component of which is named for a column in the data.frame for asreml.obj and contains a character string giving a title to use in output (e.g. tables and graphs). Here they will be used for table headings.

tables A character vector containing a combination of none, predictions, vcov, backtransforms, differences, p.differences, sed, LSD and all. These nominate which components of the alldiffs.object to print.

x.num A character string giving the name of the numeric covariate that (i) corresponds to x.fac, (ii) is potentially included in terms in the fitted model, and (iii) which corresponds to the x-axis variable. It should have the same number of unique values as the number of levels in x.fac.

x.fac A character string giving the name of the factor that (i) corresponds to x.num, (ii) is potentially included in terms in the fitted model, and (iii) which corresponds to the x-axis variable. It should have the same number of levels as the number of unique values in x.num. The levels of x.fac must be in the order in which they are to be plotted - if they are dates, then they should be in the form yyyymmd, which can be achieved using as.Date. However, the levels can be non-numeric in nature, provided that x.num is also set.

x.pred.values The values of x.num for which predicted values are required. If levels is set for passing to predict.asreml, x.pred.values is ignored. Note that while levels is an alternative to x.pred.values, it allows more general setting of the levels to be predicted.

x.plot.values The actual values to be plotted on the x axis. They are needed when values different to those in x.num are to be plotted or x.fac is to be plotted because there is no x.num term corresponding to the same term with x.fac.

level.length The maximum number of characters from the levels of factors to use in the row and column labels of the tables of pairwise differences and their p-values and standard errors.
transform.power

A **numeric** specifying the power of a transformation, if one has been applied to the response variable. Unless it is equal to 1, the default, back-transforms of the predictions will be obtained and stored in the backtransorms component of the `alldiffs.object`. The back-transformation raises the predictions to the power equal to the reciprocal of `transform.power`, unless it equals 0 in which case the exponential of the predictions is taken.

offset

A **numeric** that has been added to each value of the response after any scaling and before applying any power transformation.

scale

A **numeric** by which each value of the response has been multiplied before adding any offset and applying any power transformation.

transform.function

A **character** giving the name of a function that specifies the scale on which the predicted values are defined. This may be the result of a transformation of the data using the function or the use of the function as a link function in the fitting of a generalized linear (mixed) model (GLM(M)). The possible `transform.functions` are `identity`, `log`, `inverse`, `sqrt`, `logit`, `probit`, and `cloglog`. The predicted values and error.intervals, if not StandardError intervals, will be back-transformed using the inverse function of the `transform.function`. The standard.error column will be set to NA, unless (i) `asreml` returns columns named `transformed.value` and `approx.se`, as well as those called `predicted.values` and `standard.error` (such as when a GLM is fitted) and (ii) the values in `transformed.value` are equal to those obtained by backtransforming the `predicted.values` using the inverse function of the `transform.function`. Then, the `approx.se` values will be saved in the `standard.error` column of the `backtransforms` component of the returned `alldiffs.obj`. Also, the `transformed.value` and `approx.se` columns are removed from both the predictions and `backtransforms` components of the `alldiffs.obj`. Note that the values that end up in the `standard.errors` column are approximate for the backtransformed values and are not used in calculating `error.intervals`.

sortFactor

A **character** containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the `classify` term then `sortFactor` can be `NULL` and the order is defined by the complete set of predicted values. If there is more than one variable in the `classify` term then `sortFactor` must be set. In this case the `sortFactor` is sorted in the same order within each combination of the values of the `sortParallelToCombo` variables: the `classify` variables, excluding the `sortFactor`. There should be only one predicted value for each unique value of `sortFactor` within each set defined by a combination of the values of the `classify` variables, excluding the `sortFactor` factor. The order to use is determined by either `sortParallelToCombo` or `sortOrder`.

sortParallelToCombo

A **list** that specifies a combination of the values of the factors and numerics, excluding `sortFactor`, that are in `classify`. Each of the components of the supplied list is named for a `classify` variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of `sortFactor`. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If `sortParallelToCombo` is `NULL` then the first value of each `classify` variable, except for the `sortFactor` factor, in the predictions component is used to define `sortParallelToCombo`. If there is only one variable in the `classify` then `sortParallelToCombo` is ignored.
sortNestingFactor
A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

sortOrder
A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the desired order of the levels in the reordered components of the alldiffs.object. The argument sortParallelToCombo is ignored.

The following creates a sortOrder vector levs for factor f based on the values in x: levs <- levels(f)[order(x)].

decreasing
A logical passed to order that determines whether the order for sorting the components of the alldiffs.object is for increasing or decreasing magnitude of the predicted values.

trace
A logical that control output from ASReml-R. If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

... further arguments passed to predict.asreml.

Value
For linear.transformations set to NULL, an S3-class alldiffs.object with predictions and their standard errors and, depending on the settings of the arguments, all pairwise differences between predictions, their standard errors and p-values and LSD statistics. Also, unless the sortFactor or sortOrder arguments are invoked, the rows of predictions component are ordered so that they are in standard order for the variables in the classify. That is, the values of the last variable change with every row, those of the second-last variable only change after all the values of the last variable have been traversed; in general, the values of a variable are the same for all the combinations of the values to the variables to its right in the classify. In addition, if necessary, the order of the columns of the variables in the predictions component are changed to match their order in the classify.

If transform.power or scale is not one or offset is not zero, it will contain a data.frame with the backtransformed linear transformation of the predictions. The backtransformation will, after backtransforming for any power transformation, subtract the offset and then divide by the scale.

If error.intervals is not "none", then the predictions component and, if present, the backtransforms component will contain columns for the lower and upper values of the limits for the interval.

The name of the response, the response.title, the term, the classify, tdf, sortFactor and the sortOrder will be set as attributes to the object. Also, if error.intervals is "halfLeastSignificant", then those of LSDtype, LSDby and LSDstatistic that are not NULL will be added as attributes of the object and of the predictions frame; additionally, LSDvalues will be added as attribute of the predictions frame, LSDvalues being the LSD values used in calculating the error.intervals. Note that the classify in an alldiffs.object is based on the variables indexing the predictions, which may differ from the classify used to obtain the original predictions (for example, when the alldiffs.objects stores a linear transformation of predictions.

For linear.transformations set to other than NULL, an alldiffs.object with the linear.transformation applied to the predictions and their standard errors and, depending on the settings of the arguments, all pairwise differences between the linearly transformed predictions, their standard errors and p-values and LSD statistics. (See also linTransform.alldiffs.)

Author(s)
Chris Brien
See Also

alldiffs.object, as.alldiffs, print.alldiffs, linTransform.alldiffs, sort.alldiffs, subset.alldiffs, allDifferences.data.frame, redoErrorIntervals.alldiffs, recalclSD.alldiffs, exploreLSDs.alldiffs, pickLSDstatistics.alldiffs, predictPresent.asreml, plotPredictions.data.frame, as.Date, predict.asreml

Examples

## Not run:
data(WaterRunoff.dat)
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
random = ~ Benches:MainPlots,
keep.order = TRUE, data = WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
diffs <- predictPlus(classify = "Sources:Type",
asreml.obj = current.asr,
wald.tab = current.asrt$wald.tab,
present = c("Sources", "Type", "Species"))
## End(Not run)

predictPresent.asreml  Forms the predictions for each of one or more terms and presents them in tables and/or graphs.

Description

This function forms the predictions for each term in terms using a supplied asreml object and predictPlus.asreml. Tables are produced using predictPlus.asreml, in conjunction with allDifferences.data.frame, with the argument tables specifying which tables are printed. The argument plots, along with transform.power, controls which plots are produced. The plots are produced using plotPredictions.data.frame, with line plots produced when variables involving x.num or x.fac are involved in classify for the predictions and bar charts otherwise. In order to get the correct predictions you may need to supply additional arguments to predict.asreml through ... e.g. present, parallel, levels. The order of plotting the levels of one of the factors indexing the predictions can be modified and is achieved using sort.alldiffs.

Usage

## S3 method for class 'asreml'
predictPresent(asreml.obj, terms, inestimable.rm = TRUE,
linear.transformation = NULL,
error.intervals = "Confidence", alpha = 0.05,
wald.tab = NULL, dDF.na = "residual", dDF.values = NULL,
pairwise = TRUE, Vmatrix = FALSE,
avsed.tolerance = 0.25, accuracy.threshold = NA,
LSDtype = "overall", LSDsupplied = NULL, LSDby = NULL,
LSDstatistic = "mean", LSDaccuracy = "maxAbsDeviation",
x.num = NULL, x.fac = NULL, nonx.fac.order = NULL,
predictPresent.asreml

Arguments

asreml.obj asreml object for a fitted model.
terms A character vector giving the terms for which predictions are required.
inestimable.rm A logical indicating whether rows for predictions that are not estimable are to be removed from the components of the alldiffs.object.
linear.transformation

A formula or a matrix. If a formula is given then it is taken to be a submodel of a model term corresponding to the classify. The projection matrix that transforms the predictions so that they conform to the submodel is obtained; the submodel does not have to involve variables in the classify, but the variables must be columns in the predictions component of alldiffs.obj and the space for the submodel must be a subspace of the space for the term specified by the classify. For example, for classify set to "A:B", the submodel ~ A + B will result in the predictions for the combinations of A and B being made additive for the factors A and B. The submodel space corresponding to A + B is a subspace of the space A:B. In this case both the submodel and the the classify involve only the factors A and B. To fit an intercept-only submodel, specify linear.transformation to be the formula ~1.

If a matrix is provided then it will be used to apply the linear transformation to the predictions. It might be a contrast matrix or a matrix of weights for a factor used to obtain the weighted average over that factor. The number of rows in the matrix should equal the number of linear combinations of the predictions desired and the number of columns should equal the number of predictions.

In either case, as well as the values of the linear combinations, their standard errors, pairwise differences and associated statistics are returned in the alldiffs.object.

tables A character string indicating the type of error interval, if any, to calculate in order to indicate uncertainty in the results. Possible values are "none", "StandardError", "Confidence" and "halfLeastSignificant". The default is for confidence limits to be used. The "halfLeastSignificant" option results in half the Least Significant Difference (LSD) being added and subtracted to the predictions, the LSD being calculated using the square root of the mean of the variances of all or a subset of pairwise differences between the predictions. If the LSD is zero, as can happen when predictions are constrained to be equal, then the limits of the error intervals are set to NA. If LSDtype is set to overall, the avsed.tolerance is not NA and the range of the SEDs divided by the average of the SEDs exceeds avsed.tolerance then the error.intervals calculations and the plotting will revert to confidence intervals.
predictPresent.asreml

alpha
A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

wald.tab
A data.frame containing the pseudo-anova table for the fixed terms produced by a call to wald.asreml. The main use of it here is in determining the degrees of freedom for calculating confidence or half-LSD error.intervals and p-values, the latter to be stored in the p.differences component of the alldiffs.object that is created.

dDF.na
The method to use to obtain approximate denominator degrees of freedom. When the numeric or algebraic methods produce an NA. Consistent with when no denDF are available, the default is "residual" and so the residual degrees of freedom from asreml.obj$nedf are used. If dDF.na = "none", no substitute denominator degrees of freedom are employed; if dDF.na = "maximum", the maximum of those denDF that are available, excluding that for the Intercept, is used; if all denDF are NA, asreml.obj$nedf is used. If dDF.na = "supplied", a vector of values for the denominator degrees of freedom is to be supplied in dDF.values. Any other setting is ignored and a warning message produced. Generally, substituting these degrees of freedom is anticonservative in that it is likely that the degrees of freedom used will be too large.

dDF.values
A vector of values to be used when dDF.na = "supplied". Its values will be used when denDF in a test for a fixed effect is NA. This vector must be the same length as the number of fixed terms, including (Intercept) whose value could be NA.

pairwise
A logical indicating whether all pairwise differences of the predictions and their standard errors and p-values are to be computed and stored. If tables is equal to "differences" or "all" or error.intervals is equal to "halfLeastSignificant", they will be stored irrespective of the value of pairwise.

Vmatrix
A logical indicating whether the variance matrix of the predictions will be stored as a component of the alldiffs.object that is returned. If linear.transformation is set, it will be stored irrespective of the value of Vmatrix.

avsed.tolerance
A numeric giving the value of the SED range, the range of the SEDs divided by the square root of the mean of the variances of all or a subset of the pairwise differences, that is considered reasonable in calculating error.intervals. It should be a value between 0 and 1. The following rules apply:

1. If avsed.tolerance is NA then mean LSDs of the type specified by LSDtype are calculated and used in error.intervals and plots.
2. Irrespective of the setting of LSDtype, if avsed.tolerance is not exceeded then the mean LSDs are used in error.intervals and plots.
3. If LSDtype is set to overall, avsed.tolerance is not NA, and avsed.tolerance is exceeded then error.intervals and plotting revert to confidence intervals.
4. If LSDtype is set to factor.combinations and avsed.tolerance is not exceeded for any factor combination then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.
5. If LSDtype is set to per.prediction and avsed.tolerance is not exceeded for any prediction then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.
predictPresent.asreml

**accuracy.threshold**

A numeric specifying the value of the LSD accuracy measure, which measure is specified by LSDaccuracy, as a threshold value in determining whether the halfLeastSignificant error.interval for a predicted value is a reasonable approximation; this will be the case if the LSDs across all pairwise comparisons for which the interval’s LSD was computed, as specified by LSDtype and LSDby, are similar enough to the interval’s LSD, as measured by LSDaccuracy. If it is NA, it will be ignored. If it is not NA, a column of logicals named LSDwarning will be added to the predictions component of the alldiffs.object. The value of LSDwarning for a predicted.value will be TRUE if the value of the LSDaccuracy measure computed from the LSDs for differences between this predicted.value and the other predicted.values as compared to its assignedLSD exceeds the value of accuracy.threshold. Otherwise, the value of LSDwarning for a predicted.value will be FALSE.

**LSDtype**

A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparison2, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations. See LSD.frame for further information on the values in a row of this data.frame and how they are calculated.

**LSDsupplied**

A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function dae::fac.combine to the predictions component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into assignedLSD column of the LSD.frame stored as the LSD component of the alldiffs.object.

**LSDby**

A character (vector) of variables names, being the names of the factors or numerics in the classify; for each combination of their levels and values, there will be or is a row in the LSD.frame stored in the LSD component of the alldiffs.object when LSDtype is factor.combinations.

**LSDstatistic**

A character nominating one or more of minmum, q10, q25, mean, median, q75, q90 or maximum as the value(s) to be stored in the assignedLSD column in an LSD.frame; the values in the assignedLSD column are used in computing halfLeastSignificant error.intervals. Here q10, q25, q75 and q90 indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function quantile is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median
is calculated using the \texttt{median} function. Multiple values are only produced for LSDtype set to \texttt{factor.combination}, in which case LSDby must not be \texttt{NULL} and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is \texttt{NULL}, it is reset to mean.

\textbf{LSDaccuracy} A character nominating one of \texttt{maxAbsDeviation}, \texttt{maxDeviation}, \texttt{q90Deviation} or \texttt{RootMeanSqDeviation} as the statistic to be calculated as a measure of the accuracy of assignedLSD. The option \texttt{q90Deviation} produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named \texttt{accuracyLSD} in an \texttt{LSD.frame}.

\textbf{x.num} A character string giving the name of the numeric covariate that corresponds to \texttt{x.fac}, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of unique values as the number of levels in \texttt{x.fac}.

\textbf{x.fac} A character string giving the name of the factor that corresponds to \texttt{x.num}, is potentially included in terms in the fitted model and which corresponds to the x-axis variable. It should have the same number of levels as the number of unique values in \texttt{x.num}. The levels of \texttt{x.fac} must be in the order in which they are to be plotted - if they are dates, then they should be in the form yyyymmd, which can be achieved using \texttt{as.Date}. However, the levels can be non-numeric in nature, provided that \texttt{x.num} is also set.

\textbf{nonx.fac.order} A character vector giving the order in which factors other than \texttt{x.fac} are to be plotted in plots with multiple panels (i.e. where the number of non-x factors is greater than 1). The first factor in the vector will be plotted on the X axis (if there is no \texttt{x.num} or \texttt{x.fac}). Otherwise, the order of plotting the factors is in columns (X facets) and then rows (Y facets). By default the order is in decreasing order for the numbers of levels of the non x factors.

\textbf{x.pred.values} The values of \texttt{x.num} for which predicted values are required.

\textbf{x.plot.values} The actual values to be plotted on the x axis or in the labels of tables. They are needed when values different to those in \texttt{x.num} are to be plotted or \texttt{x.fac} is to be plotted because there is no \texttt{x.num} term corresponding to the same term with \texttt{x.fac}.

\textbf{plots} Possible values are "none", "predictions", "backtransforms" and "both". Plots are not produced if the value is "none". If data are not transformed for analysis (\texttt{transform.power = 1}), a plot of the predictions is produced provided plots is not "none". If the data are transformed, the value of plots determines what is produced.

\textbf{panels} Possible values are "single" and "multiple". When line plots are to be produced, because variables involving \texttt{x.num} or \texttt{x.fac} are involved in \texttt{classify} for the predictions, panels determines whether or not a single panel or multiple panels in a single window are produced. The panels argument is ignored for bar charts.

\textbf{graphics.device} A character specifying a graphics device for plotting. The default is \texttt{graphics.device = NULL}, which will result in plots being produced on the current graphics device. Setting it to "windows", for example, will result in a windows graphics device being opened.
interval.annotate
A logical indicating whether the plot annotation indicating the type of error.interval is to be included in the plot.

titles
A list, each component of which is named for a column in the data.frame for asreml.obj and contains a character string giving a title to use in output (e.g. tables and graphs). Here they will be used for axis labels.

colour.scheme
A character string specifying the colour scheme for the plots. The default is "colour" which produces coloured lines and bars, a grey background and white gridlines. A value of "black" results in black lines, grey bars and gridlines and a white background.

save.plots
A logical that determines whether any plots will be saved. If they are to be saved, a file name will be generated that consists of the following elements separated by full stops: the response variable name with .back if backtransformed values are being plotted, the classify term, Bar or Line and, if error.intervals is not "none", one of SE, CI or LSI. The file will be saved as a 'png' file in the current work directory.

transform.power
A numeric specifying the power of a transformation, if one has been applied to the response variable. Unless it is equal to 1, the default, back-transforms of the predictions will be obtained and stored in the backtransorms component of the alldiffs.object. The plots and tables arguments control the plotting and output of the predictions and backtransforms. The back-transformation raises the predictions to the power equal to the reciprocal of transform.power, unless it equals 0 in which case the exponential of the predictions is taken.

offset
A number that has been added to each value of the response after any scaling and before applying any power transformation. Unless it is equal to 0, the default, back-transforms of the predictions will be obtained and stored in the backtransorms component of the alldiffs.object. The plots and tables arguments control the plotting and output of the predictions and backtransforms. The backtransformation will, after backtransforming for any power transformation, subtract the offset.

scale
A number by which each value of the response has been multiply before adding any offset and applying any power transformation. Unless it is equal to 1, the default, back-transforms of the predictions will be obtained and stored in the backtransorms component of the alldiffs.object. The plots and tables arguments control the plotting and output of the predictions and backtransforms. The backtransformation will, after backtransforming for any power transformation and then subtracting the offset, divide by the scale.

transform.function
A character giving the name of a function that specifies the scale on which the predicted values are defined. This may be the result of a transformation of the data using the function or the use of the function as a link function in the fitting of a generalized linear (mixed) model (GL(M)M). The possible transform.functions are identity, log, inverse, sqrt, logit, probit, and cloglog. The predicted.values and error.intervals, if not StandardError.intervals, will be back-transformed using the inverse function of the transform.function. The standard.error column will be set to NA, unless (i) asreml returns columns named transformed.value and approx.se, as well as those called predicted.values and standard.error (such as when a GLM is fitted) and (ii) the values in transformed.value are equal to those obtained by backtransforming the predicted.values using the inverse function of the transform.function. Then, the approx.se
values will be saved in the `standard.error` column of the backtransforms component of the returned `alldiffs.obj`. Also, the `transformed.value` and `approx.se` columns are removed from both the predictions and backtransforms components of the `alldiffs.obj`. Note that the values that end up in the `standard.errors` column are approximate for the backtransformed values and are not used in calculating `error.intervals`.

**tables**

A character vector containing a combination of predictions, `vcov`, backtransforms, differences, `p.differences`, `sed`, `LSD` and `all`. These nominate which components of the `alldiffs.object` to print.

**level.length**

The maximum number of characters from the levels of factors to use in the row and column labels of the tables produced by `allDifferences.data.frame`.

**sortFactor**

A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the `classify` term then `sortFactor` can be `NULL` and the order is defined by the complete set of predicted values. If there is more than one variable in the `classify` term then `sortFactor` must be set. In this case the `sortFactor` is sorted in the same order within each combination of the values of the `sortParallelToCombo` variables: the `classify` variables, excluding the `sortFactor`. There should be only one predicted value for each unique value of `sortFactor` within each set defined by a combination of the values of the `classify` variables, excluding the `sortFactor` factor. The order to use is determined by either `sortParallelToCombo` or `sortOrder`.

**sortParallelToCombo**

A list that specifies a combination of the values of the factors and numerics, excluding `sortFactor`, that are in `classify`. Each of the components of the supplied list is named for a `classify` variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of `sortFactor`. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If `sortParallelToCombo` is `NULL` then the first value of each `classify` variable, except for the `sortFactor` factor, in the predictions component is used to define `sortParallelToCombo`. If there is only one variable in the `classify` then `sortParallelToCombo` is ignored.

**sortNestingFactor**

A character containing the name of the factor that defines groups of the `sortFactor` within which the predicted values are to be ordered. If there is only one variable in the `classify` then `sortNestingFactor` is ignored.

**sortOrder**

A character vector whose length is the same as the number of levels for `sortFactor` in the predictions component of the `alldiffs.object`. It specifies the desired order of the levels in the reordered components of the `alldiffs.object`. The argument `sortParallelToCombo` is ignored. The following creates a `sortOrder` vector `levs` for factor `f` based on the values in `x`:

```r
levs <- levels(f)[order(x)]
```

**decreasing**

A logical passed to order that determines whether the order for sorting the components of the `alldiffs.object` is for increasing or decreasing magnitude of the predicted values.

**trace**

If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

**ggplotFuncs**

A list, each element of which contains the results of evaluating a `ggplot` function. It is created by calling the `list` function with a `ggplot` function call for each element. It is passed to `plotPredictions.data.frame`.
... further arguments passed to predict.asreml via predictPlus.asreml and to ggplot via plotPredictions.data.frame.

Value

A list containing an alldiffs.object for each term for which tables are produced. The names of the components of this list are the terms with full-stops (.) replacing colons (:). Plots are also produced depending on the setting of the plot argument.

Author(s)

Chris Brien

See Also

predictPlus.asreml, allDifferences.data.frame, sort.alldiffs, subset.alldiffs, redoErrorIntervals.alldiffs, recalclSD.alldiffs, pickLSDstatistics.alldiffs, plotPredictions.data.frame, print.alldiffs, as.Date, Devices

Examples

## Not run:
data(WaterRunoff.dat)
titles <- list("Days since first observation", "Days since first observation", "pH", "Turbidity (NTU)"

names(titles) <- names(WaterRunoff.dat)[c(5,7,11:12)]
asreml.options(keep.order = TRUE)  #required for asreml-R4 only
current.asr <- asreml(fixed = log.Turbidity ~ Benches + Sources + Type + Species + Sources:Type + Sources:Species + Sources:Species:xDay + Sources:Species:Date,

  data = WaterRunoff.dat, keep.order = TRUE)
current.asrt <- as.asrtests(current.asr, NULL, NULL)

#### Get the observed combinations of the factors and variables in classify
class.facs <- c("Sources","Species","Date","xDay")
levs <- as.data.frame(table(WaterRunoff.dat[class.facs]))
levs <- levs[do.call(order, levs), ]
levs <- as.list(levs[levs$Freq != 0, class.facs])
levs$xDay <- as.numfac(levs$xDay)

#### parallel and levels are arguments from predict.asreml
diff.list <- predictPresent.asreml(asreml.obj = current.asrt$asreml.obj,

terms = "Date:Sources:Species:xDay",

  x.num = "xDay", x.fac = "Date",

  parallel = TRUE, levels = levs,

  wald.tab = current.asrt$wald.tab,

  plots = "predictions",

  error.intervals = "StandardError",

  titles = titles,

  transform.power = 0,

  present = c("Type","Species","Sources"),

  tables = "none",

  level.length = 6)

## End(Not run)
print.alldiffs  Prints the values in an alldiffs.object in a nice format.

Description

Prints the predictions and standard errors from a fitted model, including the attributes of the predictions.frame. Also prints all pairwise differences between the predictions to 2 significant figures, along with their p-values and standard errors to 4 decimal places. If LSDs are requested the mean, minimum and maximum LSDs will be printed.

Usage

## S3 method for class 'alldiffs'
print(x, which = "all", colourise = FALSE, ...)

Arguments

x  
An alldiffs.object.

which  
A character vector containing a combination of predictions, vcov, backtransforms, differences, p.differences, sed, LSD and all. These nominate which components of the alldiffs.object to print.

colourise  
A logical which, if TRUE, results in the header text produced by predict.asreml being displayed in a different colour, if supported by the output terminal device. It overrides the TRUE setting of the colourise argument of asreml::asreml.options.

...  
further arguments passed to print.predictions.frame.

Value

No value is returned, but the components of x are printed.

Author(s)

Chris Brien

See Also

print.predictions.frame, as.alldiffs, allDifferences.data.frame

Examples

## Not run:
print.alldiffs(diffs, which = "predictions")

## End(Not run)
print.asrtests  **Prints the values in an asrtests.object**

Description

Prints a summary of the asreml object, the pseudoanova and the test.summary data.frame that are stored in the asrtests.object.

Usage

```r
## S3 method for class 'asrtests'
print(x, which = "key", colourise = FALSE, ...)  
```

Arguments

- **x**: An **asrtests.object**.
- **which**: Which elements of the asrtests.object to print. Possible values are some combination of asremlsummary, vparametersummary, pseudoanova, wald.tab, testsummary and key or all. The option wald.tab is a synonym for pseudoanova. The options key and all are mutually exclusive; key includes vparametersummary, but not the rest of asremlsummar, while all includes the full asremlsummary that includes the vparametersummary.
- **colourise**: A logical which, if TRUE, results in the header text produced by wald.asreml being displayed in a different colour, if supported by the output terminal device. It overrides the TRUE setting of the colourise argument of asreml::asreml.options.
- **...**: further arguments passed to print and print.wald.tab.

Value

No value is returned, but the elements of the list in x are printed.

Author(s)

Chris Brien

See Also

print.wald.tab, as.asrtests, asremlPlus-package

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,  
  random = ~ Row + Column + units,  
  residual = ~ ar1(Row):ar1(Column),  
  data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary.asrtests(current.asrt)
# Test Row autocorrelation
current.asrt <- testresidual(current.asrt, , "~ Row:ar1(Column)",  
  label="Row autocorrelation", simpler=TRUE)
```
print.LSDdata

print(current.asrt)
## End(Not run)

print.LSDdata

Prints the components of a list containing data on the LSDs for all pairwise differences of predictions.

Description
Prints the components of an LSDdata list created by `exploreLSDs`, that contains data on the LSDs for all pairwise differences of predictions stored in an `alldiffs.object`.

Usage

## S3 method for class 'LSDdata'
print(x, which.print = c("statistics", "false.pos", "false.neg"), ...)

Arguments

x
An object that, ideally, is of class LSDdata.

which.print
Which components of the LSDdata list to print. Possible values are any combination of frequencies, distinct.vals, statistics, accuracy, false.pos, false.neg, per.pred.accuracy, LSD, summary and all, except that summary and all cannot occur together. For a description of the components, see `alldiffs.object`. The default is to print statistics, false.pos, false.neg. The option summary results in the printing of distinct.vals, statistics, false.pos, false.neg.

... further arguments passed to print.

Value
No value is returned, but components of x are printed as specified in which.print.

Author(s)
Chris Brien

See Also

`exploreLSDs.alldiffs`, `alldiffs.object`

Examples

## Not run:
data(WaterRunoff.dat)
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
    random = ~ Benches:MainPlots,
    keep.order=TRUE, data= WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
diffs <- predictPlus(classify = "Sources:Type",
    asreml.obj = current.asr,
wald.tab = current.asrt$wald.tab,
present = c("Sources", "Type", "Species"))
LSDdata <- exploreLSDs(diffs, LSDtype = "factor.combinations", LSDby = "Sources")
print(LSDdata)

## End(Not run)

print.predictions.frame

Prints the values in a predictions.frame, with or without title and heading.

Description
Prints the predictions from a fitted model, along with their standard errors and, if present, their error
intervals, with or without title and headings.

Usage

## S3 method for class 'predictions.frame'
print(x, title = NULL,
      which.predictions = c("title", "heading", "table"),
      colourise = FALSE, ...)

Arguments

x An object that, ideally, is of class predictions.frame.
title A character giving a title to be printed out before the heading and table for
the predictions.frame.
which.predictions
what Which elements of the predictions.frame to print. Possible values are
some combination of title, heading, table and all. The heading is an attribute of x.
colourise A logical which, if TRUE, results in the header text produced by predict.asreml
being displayed in a different colour, if supported by the output terminal device.
It overrides the TRUE setting of the colourise argument of asreml::asreml.options,
but is only operational when the table is also printed.

Value
No value is returned, but the components of x are printed.

Author(s)
Chris Brien

See Also

print.alldiffs, as.alldiffs, allDifferences.data.frame
print.test.summary

Examples

```r
## Not run:
print.predictions.frame(diffs$predictions, which = "all")

## End(Not run)
```

print.test.summary  

Prints a data.frame containing a test.summary.

Description

Prints a test.summary (also a choose.summary) with or without a title and with p-values limited to 4-digits.

Usage

```r
## S3 method for class 'test.summary'
print(x, which.print = c("title", "table"), omit.columns = NULL, ...)
```

Arguments

- `x`  
  A object that, ideally, is of class test.summary.

- `which.print`  
  A character specifying the aspects of the test.summary to print. Possible values are some combination of title, table and all.

- `omit.columns`  
  A character specifying the columns of the test.summary table to be omitted from the print. If NULL, none are omitted.

- `...`  
  Further arguments passed to print, but is only operational when the table is also printed.

Value

No value is returned, but x is printed, possibly with a title.

Author(s)

Chris Brien

See Also

print.wald.tab, print.asrtests, as.asrtests, asremlPlus-package

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety, 
  random = ~ Row + Column + units, 
  residual = ~ ar1(Row):ar1(Column), 
  data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary.asrtests(current.asrt)
# Test Row autocorrelation
```
current.asrt <- testresidual(current.asrt, "~ Row:ar1(Column)",
    label="Row autocorrelation", simpler=TRUE)
print(current.asrt$test.summary)

### End(Not run)

print.wald.tab Prints a data.frame containing a Wald or pseudoanova table.

Description

Prints a wald.tab with or without title and/or heading. The printing of the p-values is limited to 4 digits.

Usage

### S3 method for class 'wald.tab'
print(x, which.wald = c("title", "heading", "table"),
    colourise = FALSE, ...)

Arguments

- **x**: An object that, ideally, is of class wald.tab.
- **which.wald**: Which elements of the wald.tab to print. Possible values are some combination of title, heading, table and all. The heading is an attribute of x.
- **colourise**: A logical which, if TRUE, results in the header text produced by wald.asreml being displayed in a different colour, if supported by the output terminal device. It overrides the TRUE setting of the colourise argument of asreml::asreml.options.
- **...**: further arguments passed to print and print.wald.tab, but is only operational when the table is also printed.

Value

No value is returned, but x is printed as specified in which.wald.

Author(s)

Chris Brien

See Also

print.test.summary, print.asrtests, as.asrtests, asremlPlus-package

Examples

### Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
    random = ~ Row + Column + units,
    residual = ~ ar1(Row):ar1(Column),
    data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
current.asrt <- rmboundary.asrtests(current.asrt)  # Test Row autocorrelation
current.asrt <- testresidual(current.asrt, "~ Row:ar1(Column)",
              label="Row autocorrelation", simpler=TRUE)
print(current.asrt$wald.tab)

## End(Not run)

printFormulae(asreml)

# Prints the formulae from an asreml object.

Description

Prints the formulae nominated in the which argument from the call stored in an asreml object.

Usage

## S3 method for class 'asreml'
printFormulae(asreml.obj, which = c("fixed", "random", "residual"),
              expanded = FALSE, envir = parent.frame(), ...)

Arguments

- `asreml.obj`: An asreml object resulting from the fitting of a model using REML.
- `which`: A character listing the formula(e) to be printed from the call stored in asreml.obj. It should be some combination of fixed, random, residual, sparse and all. If all is included then all formula(e) will be printed.
- `expanded`: A logical indicating whether terms are to be expanded to the sum of a set of individual terms.
- `envir`: The environment in which the formula(e) are to be evaluated. May also be NULL, a list, a data.frame, a pairlist or an integer as specified to sys.call.
- `...`: Arguments passed on to getFormulae.asreml and ultimately to update.formula and terms.formula.

Value

Invisibly returns a character, each element of which contains one of the extracted formulae.

Author(s)

Chris Brien

See Also

printFormulae.asreml
Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                   random = ~ Row + Column + units,
                   residual = ~ ar1(Row):ar1(Column),
                   data=Wheat.dat)
printFormulae(current.asr)
## End(Not run)
```

ratioTransform.alldiffs

*Calculates the ratios of nominated pairs of predictions stored in an alldiffs.object.*

Description

Ratio predictions and error intervals are formed for two levels of a factor, the `ratio.factor`. For each pair of levels of the `ratio.factor` in `numerator.levels` with a level in `denominator.levels`, the ratio predictions are formed from all combinations of the other factors as the ratio of the two predictions for each combination, along with confidence intervals for the ratio predictions computed using the Fieller (1954) method.

The printing of the components produced is controlled by the `tables` argument.

Usage

```r
## S3 method for class 'alldiffs'

ratioTransform(alldiffs.obj, ratio.factor, numerator.levels, denominator.levels, method = "Fieller", alpha = 0.05, response = NULL, response.title = NULL, tables = "predictions", ...)
```

Arguments

- `alldiffs.obj` An *alldiffs.object*.
- `ratio.factor` A character string giving the name of the factor for whose levels the ratios are to be calculated.
- `numerator.levels` A character string containing the levels of `ratio.factor` to be used as numerators of the ratio.
- `denominator.levels` A character string containing the levels of `ratio.factor` to be used as denominators of the ratio.
- `method` A character string specifying the method to use in calculating the ratios and their error.intervals. At present only Fieller is available. For the Fieller method, ratios of predictions are formed and confidence intervals formed for them using Fieller’s (1954) theorem.
alpha       A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals.
response    A character specifying the response variable for the predictions. It is stored as an attribute to the alldiffs.object.
response.title A character specifying the title for the response variable for the predictions. It is stored as an attribute to the alldiffs.object.
tables      A character vector containing either none or predictions
...         further arguments passed to linTransform.alldiffs.

Value
A list of predictions.frame, each containing the ratio predictions and their confidence limits for a combination of the numerator.levels with the denominator.levels. It will also contain the values of the variables in the classify of alldiffs.obj that index the ratio predictions, except that the ratio.factor is omitted.

If sortFactor attribute of the alldiffs.object is set and is not the ratio.factor, the predictions and their backtransforms will be sorted using the sortOrder attribute of the alldiffs.object.

Author(s)
Chris Brien

References

See Also
pairdiffsTransform, linTransform, predictPlus.asreml, as.alldiffs, print.alldiffs, sort.alldiffs, subset.alldiffs, allDifferences.data.frame, redoErrorIntervals.alldiffs, recalcLSD.alldiffs, predictPresent.asreml, plotPredictions.data.frame, as.Date, predict.asreml

Examples
#### Form the ratios and Fieller CIs for RGR Salinity
load(system.file("extdata", "testDiffs.rda", package = "asremlPlus", mustWork = TRUE))
Preds.ratio.RGR <- ratioTransform(diffs.RGR,
                                   ratio.factor = "Salinity",
                                   numerator.levels = "Salt",
                                   denominator.levels = "Control")

#### Form the ratios and Fieller CIs for Nitrogen compared to no Nitrogen
data("Oats.dat")
## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety,
                 random=~Blocks/Wplots,
                 data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
wald.tab <- current.asrt$wald.tab
Var.diffs <- predictPlus(m1.asr, classify="Nitrogen:Variety", pairwise = TRUE,
Vmatrix = TRUE, error.intervals = "halfLeast",
LSDtope = "factor", LSDby = "Variety",
wald.tab = wald.tab)

## End(Not run)

## Use lme4 and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
data=Oats.dat)
  ## Set up a wald.tab
  int <- as.data.frame(rbind(rep(NA,4)))
  rownames(int) <- "(Intercept)"
  wald.tab <- anova(m1.lmer, ddf = "Kenward", type = 1)[,3:6]
  names(wald.tab) <- names(int) <- c("Df", "denDF", "F.inc", "Pr")
  wald.tab <- rbind(int, wald.tab)
  #Get predictions
  Var.emm <- emmeans::emmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                   se = "SE", interval.type = "CI",
                                   interval.names = c("lower.CL", "upper.CL"))
  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
  den.df <- wald.tab[match("Variety", rownames(wald.tab)), "denDF"]
  #Create alldiffs object
  Var.diffs <- as.alldiffs(predictions = Var.preds,
                           sed = Var.sed, vcov = Var.vcov,
                           classify = "Nitrogen:Variety", response = "Yield", tdf = den.df)
}

if (exists("Var.diffs"))
  Preds.ratio.OatsN <- ratioTransform(alldiffs.obj = Var.diffs,
                                      ratio.factor = "Nitrogen",
                                      numerator.levels = c("0.2","0.4","0.6"),
                                      denominator.levels = "0.2")

recalcLSD.alldiffs

Add or recalculates the LSD.frame that is a component of an alldiffs.object.

Description

Given an alldiffs.object, adds or recalculate its LSD.frame. N.B. No changes are made to the error.intervals — use redoErrorIntervals.alldiffs to modify both the error.intervals and the LSD.frame.

Usage

## S3 method for class 'alldiffs'

### Description

Given an alldiffs.object, adds or recalculate its LSD.frame. N.B. No changes are made to the error.intervals — use redoErrorIntervals.alldiffs to modify both the error.intervals and the LSD.frame.

### Usage

## S3 method for class 'alldiffs'

recalcLSD(alldiffs.obj, LSDtype = "overall", LSDsupplied = NULL, 
LSDby = NULL, LSDstatistic = "mean", LSDaccuracy = "maxAbsDeviation", 
alpha = 0.05, ...
)

**Arguments**

**alldiffs.obj**  
An **alldiffs.object**.

**LSDtype**  
A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a **LSD.frame** are the values calculated (i) overall from the LSD values for all pairwise comparison2, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the **LSD.frame** stored in an **alldiffs.object** so that they can be used in LSD calculations.

See **LSD.frame** for further information on the values in a row of this data.frame and how they are calculated.

**LSDsupplied**  
A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds. (Applying the function dae::fac.combine to the predictions component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into assignedLSD column of the **LSD.frame** stored as the LSD component of the **alldiffs.object**.

**LSDby**  
A character (vector) of variables names, being the names of the factors or numerics in the classify; for each combination of their levels and values, there will be or is a row in the **LSD.frame** stored in the LSD component of the **alldiffs.object** when LSDtype is factor.combinators.

**LSDstatistic**  
A character nominating one or more of minimum, q10, q25, mean, median, q75, q90 or maximum as the value(s) to be stored in the assignedLSD column in an **LSD.frame**; the values in the assignedLSD column are used in computing halfLeastSignificant error.intervals. Here q10, q25, q75 and q90 indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function quantile is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median is calculated using the median function. Multiple values are only produced for LSDtype set to factor.combination, in which case LSDby must not be NULL and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is NULL, it is reset to mean.
LSDaccuracy  
A character nominating one of maxAbsDeviation, maxDeviation, q90Deviation or RootMeanSqDeviation as the statistic to be calculated as a measure of the accuracy of assignedLSD. The option q90Deviation produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named accuracyLSD in an LSD.frame.

alpha  
The significance level for an LSD to compare a pair of predictions. It is stored as an attribute to the alldiffs.object.

Value  
An alldiffs.object with components predictions, vcov, differences, p.differences sed, LSD and, if present in alldiffs.obj, backtransforms.

Author(s)  
Chris Brien

See Also  
asremlPlus-package, as.alldiffs, sort.alldiffs, subset.alldiffs, print.alldiffs, renewClassify.alldiffs, exploreLSDs.alldiffs, pickLSDtoStatistics.alldiffs, redoErrorIntervals.alldiffs, plotPredictions.data.frame, predictPlus.asreml, predictPresent.asreml

Examples  
data(WaterRunoff.dat)

## Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) # required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                     random = ~ Benches:MainPlots,
                     keep.order=TRUE, data= WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus(classify = "Sources:Type",
                        asreml.obj = current.asr,
                        wald.tab = current.asrt$wald.tab,
                        present = c("Sources", "Type", "Species"))

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                           (1|Benches:MainPlots),
                           data=na.omit(WaterRunoff.dat))
TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Species)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
                                    se = "SE", interval.type = "CI",
                                    interval.names = c("lower.CL", "upper.CL"))

## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els, els]
TS.diffs <- allDifferences(predictions = TS.preds, classify = "Sources:Species",
                           vcov = TS.vcov, tdf = den.df)
validAllDiffs(TS.diffs)
}

## Plot p-values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs")) {
  ## Recalculate the LSD values for predictions obtained using asreml or lmerTest
  TS.diffs <- recalcLSD.alldiffs(TS.diffs, LSDtype = "factor.combinations",
                              LSDby = "Sources")
}

### Description

If some or all denDF are not available, either because they are NA or because F.inc values were not calculated, this function allows the user to specify how approximate denDF values are to be obtained. This is done through the dDF.na and dDF.values arguments. Note that if denDF values are available in the Wald table then only those that are NA will be replaced. The P values are recalculated using F.com, if present in the wald.tab, otherwise F.inc is used. It is noted that, as of asreml version 4, wald.asreml has a kenadj argument.

#### Usage

```r
## S3 method for class 'asrtests'
recalcWaldTab(asrtests.obj, recalc.wald = FALSE,
              denDF = "numeric", dDF.na = "none",
              dDF.values = NULL, trace = FALSE, ...)
```

#### Arguments

- `asrtests.obj`: an `asrtests.object` containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.
- `recalc.wald`: A logical indicating whether to call wald.asreml to recalculate the pseudo-anova table for the model fit stored in the asreml object contained in `asrtests.obj`. 

---

**RecalcWaldTab.asrtests**

Recalculates the denDF, F.inc and P values for a table of Wald test statistics obtained using wald.asreml
recalcWaldTab.asrtests

`denDF` Specifies the method to use in computing approximate denominator degrees of freedom when `wald.asreml` is called. Can be `none` to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or `default`, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

`dDF.na` The method to use to obtain substitute denominator degrees of freedom. when the numeric or algebraic methods produce an NA. If `dDF.na = "none"`, no substitute denominator degrees of freedom are employed; if `dDF.na = "residual"`, the residual degrees of freedom from `asreml.obj$nedf` are used; if `dDF.na = "maximum"`, the maximum of those `denDF` that are available, excluding that for the Intercept, is used; if all `denDF` are NA, `asreml.obj$nedf` is used. If `dDF.na = "supplied"`, a vector of values for the denominator degrees of freedom is to be supplied in `dDF.values`. Any other setting is ignored and a warning message produced. Generally, substituting these degrees of freedom is anticonservative in that it is likely that the degrees of freedom used will be too large.

`dDF.values` A vector of values to be used when `dDF.na = "supplied"`. Its values will be used when `denDF` in a test for a fixed effect is NA. This vector must be the same length as the number of fixed terms, including (Intercept) whose value could be NA.

`trace` If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

... further arguments passed to `asreml` and to `wald.asreml`.

**Value**

A `wald.tab`: a 4- or 6-column data.frame containing a pseudo-anova table for the fixed terms produced by `wald.asreml`.

**Author(s)**

Chris Brien

**References**


**See Also**

`as.asrtests`, `testranfix.asrtests`

**Examples**

```r
## Not run:
wald.tab <- recalcWaldTab(current.asrt,
    dDF.na = "supplied",
    dDF.values = c(NA, rep(c(330, 346), c(4, 3))))

## End(Not run)
```
redoErrorIntervals.alldiffs

Adds or replaces the error intervals stored in a prediction component of an alldiffs.object.

Description

Given an alldiffs.object, adds or replaces error.intervals for its prediction component. If the backtransforms component is present, the transform.power, offset and scale will be retrieved from the backtransforms attributes, ignoring the values for the function’s arguments, and the backtransformed error.intervals will also be calculated.

Usage

## S3 method for class 'alldiffs'
redoErrorIntervals(alldiffs.obj, error.intervals = "Confidence", alpha = 0.05, avsed.tolerance = 0.25, accuracy.threshold = NA, LSDtype = NULL, LSDsupplied = NULL, LSDby = NULL, LSDstatistic = "mean", LSDaccuracy = "maxAbsDeviation", retain.zeroLSDs = FALSE, zero.tolerance = .Machine$double.eps ^ 0.5, ...)

Arguments

alldiffs.obj  An alldiffs.object.

error.intervals

A character string indicating the type of error interval, if any, to calculate in order to indicate uncertainty in the predicted values. Possible values are "none", "StandardError", "Confidence" and "halfLeastSignificant". The default is for confidence limits to be used. The "halfLeastSignificant" option results in half the Least Significant Difference (LSD) being added and subtracted to the predictions, the LSD being calculated using the square root of the mean of the variances of all or a subset of pairwise differences between the predictions. If the LSD is zero, as can happen when predictions are constrained to be equal, then the limits of the error intervals are set to NA. If LSDtype is set to overall, the avsed.tolerance is not NA and the range of the SEDs divided by the average of the SEDs exceeds avsed.tolerance then the error.intervals calculations and the plotting will revert to confidence intervals.

alpha

A numeric giving the significance level for LSDs or one minus the confidence level for confidence intervals. It is stored as an attribute to the alldiffs.object.

avsed.tolerance

A numeric giving the value of the SED range, the range of the SEDs divided by the square root of the mean of the variances of all or a subset of the pairwise differences, that is considered reasonable in calculating error.intervals. To have it ignored, set it to NA. It should be a value between 0 and 1. The following rules apply:

1. If avsed.tolerance is NA then mean LSDs of the type specified by LSDtype are calculated and used in error.intervals and plots.
2. Irrespective of the setting of LSDtype, if avsed.tolerance is not exceeded then the mean LSDs are used in error.intervals and plots.

3. If LSDtype is set to overall, avsed.tolerance is not NA, and avsed.tolerance is exceeded then error.intervals and plotting revert to confidence intervals.

4. If LSDtype is set to factor.combinations and avsed.tolerance is not exceeded for any factor combination then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

5. If LSDtype is set to per.prediction and avsed.tolerance is not exceeded for any prediction then the half LSDs are used in error.intervals and plots; otherwise, error.intervals and plotting revert to confidence intervals.

accuracy.threshold
A numeric specifying the value of the LSD accuracy measure, which measure is specified by LSDaccuracy, as a threshold value in determining whether the halfLeastSignificant error.interval for a predicted value is a reasonable approximation; this will be the case if the LSDs across all pairwise comparisons for which the interval’s LSD was computed, as specified by LSDtype and LSDby, are similar enough to the interval’s LSD, as measured by LSDaccuracy. If it is NA, it will be ignored. If it is not NA, a column of logicals named LSDwarning will be added to the predictions component of the alldiffs.object. The value of LSDwarning for a predicted.value will be TRUE if the value of the LSDaccuracy measure computed from the LSDs for differences between this predicted.value and the other predicted.values as compared to its assignedLSD exceeds the value of accuracy.threshold. Otherwise, the value of LSDwarning for a predicted.value will be FALSE.

LSDtype
A character string that can be overall, factor.combinations, per.prediction or supplied. It determines whether the values stored in a row of a LSD.frame are the values calculated (i) overall from the LSD values for all pairwise comparisons, (ii) the values calculated from the pairwise LSDs for the levels of each factor.combination, unless there is only one prediction for a level of the factor.combination, when a notional LSD is calculated, (iii) per.prediction, being based, for each prediction, on all pairwise differences involving that prediction, or (iv) as supplied values of the LSD, specified with the LSDsupplied argument; these supplied values are to be placed in the assignedLSD column of the LSD.frame stored in an alldiffs.object so that they can be used in LSD calculations.

If LSDtype is NULL (the default), the LSDtype attribute of the alldiffs.obj will be used; it is also NULL, then the LSDtype will be set to overall.

See LSD.frame for further information on the values in a row of this data.frame and how they are calculated.

LSDsupplied
A data.frame or a named numeric containing a set of LSD values that correspond to the observed combinations of the values of the LSDby variables in the predictions.frame or a single LSD value that is an overall LSD. If a data.frame, it may have (i) a column for the LSDby variable and a column of LSD values or (ii) a single column of LSD values with rownames being the combinations of the observed values of the LSDby variables. Any name can be used for the column of LSD values; assignedLSD is sensible, but not obligatory. Otherwise, a numeric containing the LSD values, each of which is named for the observed combination of the values of the LSDby variables to which it corresponds.
(Applying the function `dae::fac.combine` to the `predictions` component is one way of forming the required combinations for the (row) names.) The values supplied will be incorporated into `assignedLSD` column of the `LSD.frame` stored as the LSD component of the `alldiffs.object`.

**LSDby**
A character (vector) of variables names, being the names of the factors or numerics in the `classify`; for each combination of their levels and values, there will be or is a row in the `LSD.frame` stored in the LSD component of the `alldiffs.object` when LSDtype is `factor.combinations`.

**LSDstatistic**
A character nominating one or more of `minimum`, `q10`, `q25`, `mean`, `median`, `q75`, `q90` or maximum as the value(s) to be stored in the `assignedLSD` column in an `LSD.frame`; the values in the `assignedLSD` column are used in computing halfLeastSignificant error.intervals. Here `q10`, `q25`, `q75` and `q90` indicate the sample quantiles corresponding to probabilities of 0.1, 0.25, 0.75 and 0.9 for the group of LSDs from which a single LSD value is calculated. The function `quantile` is used to obtain them. The mean LSD is calculated as the square root of the mean of the squares of the LSDs for the group. The median is calculated using the `median` function. Multiple values are only produced for LSDtype set to `factor.combination`, in which case LSDby must not be `NULL` and the number of values must equal the number of observed combinations of the values of the variables specified by LSDby. If LSDstatistic is `NULL`, it is reset to `mean`.

**LSDaccuracy**
A character nominating one of `maxAbsDeviation`, `maxDeviation`, `q90Deviation` or `RootMeanSqDeviation` as the statistic to be calculated as a measure of the accuracy of `assignedLSD`. The option `q90Deviation` produces the sample quantile corresponding to a probability of 0.90. The deviations are the differences between the LSDs used in calculating the LSD statistics and each assigned LSD and the accuracy is expressed as a proportion of the assigned LSD value. The calculated values are stored in the column named `accuracyLSD` in an `LSD.frame`.

**retain.zeroLSDs**
A logical indicating whether to retain or omit LSDs that are zero when calculating the summaries of LSDs.

**zero.tolerance**
A numeric specifying the value such that if an LSD is less than it, it will be considered to be zero.

... further arguments passed to `recalcLSD.alldiffs`.

**Value**
An `alldiffs.object` with components `predictions`, `vcov`, `differences`, `p.differences` `sed`, `LSD` and, if present in `alldiffs.obj`, `backtransforms`.

If `error.intervals` is not "none", then the `predictions` component and, if present, the `backtransforms` component will contain columns for the lower and upper values of the limits for the interval. The names of these columns will consist of three parts separated by full stops: 1) the first part will be `lower` or `upper`; 2) the second part will be one of `Confidence`, `StandardError` or `halfLeastSignificant`; 3) the third component will be `limits`.

The name of the response, the term, the `classify` and `tdf`, as well as the degrees of freedom of the standard error, will be set as attributes to the object. Also, if `error.intervals` is "halfLeastSignificant", then those of LSDtype, LSDby and LSDstatistic that are not `NULL` will be added as attributes of the object and of the `predictions.frame`; additionally, LSDvalues will be added as attribute of the `predictions.frame`, LSDvalues being the LSD values used in calculating the `error.intervals`.
Author(s)

Chris Brien

See Also

decallSD.alldiffs, exploreLSDs.alldiffs, pickLSDstatistics.alldiffs,
predictPresent.asreml.plotPredictions.data.frame.allDifferences.data.frame,
as.alldiffs.print.allDiffs.sort.allDiffs.subset.alldiffs,
as.Date.predict.asreml

Examples

data(WaterRunoff.dat)

## Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) # required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
random = ~ Benches:MainPlots,
keep.order=TRUE, data= WaterRunoff.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
TS.diffs <- predictPlus(classify = "Sources:Type",
            asreml.obj = current.asr,
            wald.tab = current.asrt$wald.tab,
            present = c("Sources", "Type", "Species"))

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE)) {
  m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                      (1|Benches:MainPlots),
                      data=na.omit(WaterRunoff.dat))
  TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Species)
  TS.preds <- summary(TS.emm)
  den.df <- min(TS.preds$df, na.rm = TRUE)
  ## Modify TS.preds to be compatible with a predictions.frame
  TS.preds <- as.predictions.frame(TS.preds,
            predictions = "emmean",
            se = "SE", interval.type = "CI",
            interval.names = c("lower.CL", "upper.CL"))

  ## Form an all.diffs object and check its validity
  els <- as.numeric(rownames(TS.preds))
  TS.vcov <- vcov(TS.emm)[els,els]
  TS.diffs <- allDifferences(predictions = TS.preds,
                classify = "Sources:Species",
                vcov = TS.vcov, tdf = den.df)
  validAlldiffs(TS.diffs)
}

## Plot p-values for predictions obtained using asreml or lmerTest
if (exists("TS.diffs"))
REMLRT.asreml

Perform a REML ratio test to compare two models.

Description

Extracts the REML log likelihood and the number of variance parameters from two asreml objects. It assumes that the first asreml object corresponds to the null hypothesis and the second asreml object to the alternative hypothesis for the test being conducted. That is, the second asreml object is the result of fitting a model that is a reduced version of the model for the first object. In the case where the reduced model is obtained by setting positively-constrained variance parameters in the full model to zero, the positive.zero argument should be set to TRUE so that the p-value is computed using a mixture of chi-square distributions as described in Self and Liang (1987).

The function checks that the models do not differ in either their fixed or sparse models.

Usage

```r
## S3 method for class 'asreml'
REMLRT(h0.asreml.obj, h1.asreml.obj,
    positive.zero = FALSE, bound.test.parameters = "none",
    DF = NULL, bound.exclusions = c("F","B","S","C"), ...)
```

Arguments

- **h0.asreml.obj**: asreml object containing the fit under the model for the null hypothesis.
- **h1.asreml.obj**: asreml object containing the fit under the model for the alternative hypothesis.
- **positive.zero**: Indicates whether the hypothesized values for the variance components being tested are on the boundary of the parameter space. For example, this is true for positively-constrained variance components that, under the reduced model, are zero. This argument does not need to be set if bound.test.parameters is set.
- **bound.test.parameters**: Indicates whether for the variance components being tested, at least some of the hypothesized values are on the boundary of the parameter space. The possibilities are "none", "onlybound" and "one-and-one". The default is "none", although if it is set to "none" and positive.zero is TRUE then bound.test.parameters is taken to be "onlybound". When bound.test.parameters is set to "one-and-one", it signifies that there are two parameters being tested, one of which is bound and the other is not. For example, the latter is true for testing a covariance and a positively-constrained variance component that, under the reduced model, are zero.
- **DF**: A numeric giving the difference between the two models in the number of variance parameters whose estimates are not of the type specified in bound.exclusions. If NULL then this is determined from the information in full.asreml.obj and reduced.asreml.obj.

```r
tS.diffs <- redoErrorIntervals.alldifTs(TS.diffs,
    error.intervals = "halfLeastSignificant")
```
bound.exclusions

A character specifying one or more bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters. If set to NULL then none will be excluded.

... Provision for passing arguments to functions called internally - not used at present.

Value

A data.frame containing the log of the likelihood ratio, its degrees of freedom, its p-value and the number of bound parameters in each of the two models being compared.

Note

If DF is not NULL, the supplied value is used. Otherwise DF is determined from the information in h1.asreml.obj and h0.asreml.obj. In this case, the degrees of freedom for the test are computed as the difference between the two models in the number of variance parameters whose estimates do not have a code for bound specified in bound.exclusions.

If ASReml-R version 4 is being used then the codes specified in bound.exclusions are not restricted to a subset of the default codes, but a warning is issued if a code other than these is specified. For ASReml-R version 3, only a subset of the default codes are allowed: F (Fixed), B (Boundary), C (Constrained) and S (Singular).

The test statistic is calculated as $2(\log(REML)_1 - \log(REML)_0)$.

This procedure is only appropriate when the null hypothesis is that (i) all parameters are on the boundary of the parameter space (ii) all parameters are in the interior of the parameter space, or (iii) there are two parameters, one of which is on the boundary and the other is not. Other cases have been discussed by Self and Liang (1987), but are not implemented here.

Author(s)

Chris Brien

References


See Also

infoCriteria.asreml, testranfix.asrtests

Examples

```r
## Not run:
REMLRT(ICV.max, ICV.red, bound.test.parameters = "onlybound")

## End(Not run)
```
renewClassify.alldiffs

Renews the components in an \texttt{alldiffs.object} according to a new classify.

Description

The classify is an attribute of an \texttt{alldiffs.object} and determines the order within the components of an unsorted \texttt{alldiffs.object}. This function resets the classify attribute and re-orders the components of \texttt{alldiffs.object} to be in standard order for the variables in a newclassify, using \texttt{allDifferences.data.frame}. The newclassify may be just a re-ordering of the variable names in the previous classify, or be based on a new set of variable names. The latter is particularly useful when \texttt{linTransform.alldiffs} has been used with a \texttt{matrix} and it is desired to replace the resulting Combination classify with a newclassify comprised of a more meaningful set of variables; first replace Combination in the predictions component with the new set of variables and then call renewClassify.

Usage

\begin{verbatim}
## S3 method for class 'alldiffs'
renewClassify(alldiffs.obj, newclassify, 
  sortFactor = NULL, sortParallelToCombo = NULL, 
  sortNestingFactor = NULL, sortOrder = NULL, decreasing = FALSE, ...)
\end{verbatim}

Arguments

- \texttt{alldiffs.obj} An \texttt{alldiffs.object}.
- \texttt{newclassify} A \texttt{character} string giving the variables that define the margins of the multiway table that was predicted, but ordered so that the predictions are in the desired order when they are arranged in standard order for the newclassify. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator. The number of combined values of the set of variable name(s) must equal the number of rows in the predictions component.
- \texttt{sortFactor} A \texttt{character} containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then \texttt{sortFactor} can be \texttt{NULL} and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then \texttt{sortFactor} must be set. In this case the \texttt{sortFactor} is sorted in the same order within each combination of the values of the \texttt{sortParallelToCombo} variables: the classify variables, excluding the \texttt{sortFactor}. There should be only one predicted value for each unique value of \texttt{sortFactor} within each set defined by a combination of the values of the classify variables, excluding the \texttt{sortFactor} factor. The order to use is determined by either \texttt{sortParallelToCombo} or \texttt{sortOrder}.
- \texttt{sortParallelToCombo} A \texttt{list} that specifies a combination of the values of the factors and numerics, excluding \texttt{sortFactor}, that are in classify. Each of the components of the supplied \texttt{list} is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the
predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.

sortNestingFactor
A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

sortOrder
A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the desired order of the levels in the reordered components of the alldiffs.object. The argument sortParallelToCombo is ignored.

The following creates a sortOrder vector lev for factor f based on the values in x: lev <- levels(f)[order(x)].

decreasing
A logical passed to order that determines whether the order is for increasing or decreasing magnitude of the predicted values.

... further arguments passed to allDifferences.data.frame; attributes transform.power, offset and scale cannot be passed.

Details
First, the components of the alldiffs.object is arranged in standard order for the newclassify. Then predictions are reordered according to the settings of sortFactor, sortParallelToCombo, sortOrder and decreasing (see sort.alldiffs for details).

Value
The alldiffs.object supplied with the following components, if present, sorted: predictions, vcov, backtransforms, differences, p.differences and sed. Also, the sortFactor and sortOrder attributes are set.

Author(s)
Chris Brien

See Also
as.alldiffs, allDifferences.data.frame, print.alldiffs, sort.alldiffs, redoErrorIntervals.alldiffs, recalcLSD.alldiffs, predictPlus.asreml, predictPresent.asreml

Examples

data(WaterRunoff.dat)

##Use asreml to get predictions and associated statistics

## Not run:
#Analyse pH
m1.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                random = ~ Benches:MainPlots,
reparamSigDevn.asrtests

Reparamterizes each random (deviations) term involving devn.fac to a fixed term and ensures that the same term, with trend.num replacing devn.fac, is included if any other term with trend.num is included in terms.
Description

This function reparameterizes each random (deviations) term involving devn.fac to a fixed term and ensures that the same term with trend.num replacing devn.fac is included if any other term with trend.num is included in terms. It also ensures that any term with spl(trend.num) replacing devn.fac in a term being reparameterized is removed from the model.

Usage

## S3 method for class 'asrtests'
reparamSigDevn(asrtests.obj, terms = NULL,
               trend.num = NULL, devn.fac = NULL,
               allow.unconverged = TRUE, allow.fixedcorrelation = TRUE,
               checkboundaryonly = FALSE,
               denDF = "numeric", IClikelihood = "none",
               trace = FALSE, update = TRUE,
               set.terms = NULL, ignore.suffices = TRUE,
               bounds = "P", initial.values = NA,...)

Arguments

asrtests.obj  an `asrtests.object` containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.
terms  A character string vector giving the random terms that are to be reparameterized.
trend.num  A character string giving the name of the numeric covariate that corresponds to devn.fac and is potentially included in terms in the fitted model.devn.fac  A character string giving the name of the factor that corresponds to trend.num and is included in terms in the fitted model. The name must match those in the vparameters component of the asreml.obj component in the asrtests.obj.allow.unconverged  A logical indicating whether to accept a new model even when it does not converge. Initially all changes are made with allow.unconverged set to TRUE. If allow.unconverged has been set to FALSE in the call and the final fit does not converge, an attempt is made to achieve convergence by removing any boundary terms. If this is unsuccessful, the supplied asrtests.obj is returned.allow.fixedcorrelation  A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.checkboundaryonly  If TRUE then boundary and singular terms are not removed by `rmboundary.asrtests`; a warning is issued instead.denDF  Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are
calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

IClikelihood A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new asrtests.object. If none, none are included. Otherwise, if REML, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full, then the AIC and BIC based on the full likelihood are included. (See also infoCriteria.asreml.)

trace If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update If TRUE then update.asreml is called in removing and adding terms to the model. In doing this the arguments R.param and G.param are set to those in the asreml.object stored in the supplied asrtests.obj so that the values from the previous model are used as starting values. If FALSE then calls are made to asreml in which the only changes from the previous call are (i) that the models are updated and (ii) modifications specified via ... are made.

set.terms A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting.

ignore.suffices A logical vector specifying whether the suffices of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in terms.

bounds A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

initial.values A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

... further arguments passed to asreml via changeTerms.asrtests and as.asrtests.

Value

An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

Author(s)

Chris Brien

References

See Also

as.asrtests, changeTerms.asrtests, testranfix.asrtests, testresidual.asrtests, newfit.asreml, chooseModel.asrtests

Examples

```r
## Not run:
data(WaterRunoff.dat)
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = log.Turbidity ~ Benches + Sources + Type + Species +
                      Sources:Type + Sources:Species + Sources:Species:xDay +
                      Sources:Species:Date,
data = WaterRunoff.dat, keep.order = TRUE)
current.asrt <- as.asrtests(current.asr, NULL, NULL)

#Examine terms that describe just the interactions of Date and the treatment factors
terms.treat <- c("Sources", "Type", "Species", "Sources:Type", "Sources:Species")
date.terms <- sapply(terms.treat,
  FUN=function(term){paste("Date:'",term,sep="")},
simplify=TRUE)
date.terms <- c("Date", date.terms)
date.terms <- unname(date.terms)
treat.marginality <- matrix(c(1,0,0,0,0, 1,0,1,0,0, 1,0,1,1,0,1,0,0, 1,0,1,0,0,0, 1,0,1,1,0,1,0,0, 1,0,1,0,0,0, 1,0,1,1,0,1,0,0, 1,0,1,0,0,0, 1,0,1,1,0,1,0,0, 1,0,1,0,0,0, 1,0,1,1,0,1,0,0, 1,0,1,0,0,0, 1,0,1,1,0,1,0,0, nrow=6)
rownames(treat.marginality) <- date.terms
colnames(treat.marginality) <- date.terms
choose <- chooseModel(current.asrt, treat.marginality, denDF="algebraic")
current.asrt <- choose$asrtests.obj
current.asr <- current.asrt$asreml.obj
sig.date.terms <- choose$sig.terms

#Remove all Date terms left in the fixed model
terms <- "(Date/(Sources * (Type + Species)))"
current.asrt <- changeTerms(current.asrt, dropFixed = terms)

#if there are significant date terms, reparameterize to xDays + spl(xDays) + Date
if (length(sig.date.terms) != 0)
  {
    #add lin + spl + devn for each to fixed and random models
trend.date.terms <- sapply(sig.date.terms,
      FUN=function(term){paste("xDay","",term,"")},
simplify=TRUE)
trend.date.terms <- paste(trend.date.terms, collapse=" + ")
current.asrt <- changeTerms(current.asrt, addFixed=trend.date.terms)
trend.date.terms <- sapply(sig.date.terms,
      FUN=function(term){paste("spl(xDay)","",term,"")},
simplify=TRUE)
trend.date.terms <- paste(trend.date.terms, collapse=" + ")
current.asrt <- changeTerms(current.asrt, addRandom = trend.date.terms)
current.asrt <- rmboundary(current.asrt)
  }

#Now test terms for sig date terms
spl.terms <- sapply(terms.treat,
    FUN=function(term){paste("spl(xDay)","",term,"")},
simplify=TRUE)
spl.terms <- c("spl(xDay)",spl.terms)
```
lin.terms <- sapply(terms.treat, FUN=function(term){paste(term,":xDay",sep="")}, simplify=TRUE)
lin.terms <- c("xDay",lin.terms)
systematic.terms <- c(terms.treat, lin.terms, spl.terms, date.terms)
systematic.terms <- unname(systematic.terms)
treat.marginality <- matrix(c(1,0,0,0,0,0, 1,0,1,0,0,0, 1,0,1,1,0,0, 1,0,1,1,1,0, 1,1,1,1,1,0, 1,1,1,1,1,1), nrow=6)
systematic.marginality <- kronecker(matrix(c(1,0,0,0, 1,1,0,0, 1,1,1,0, 1,1,1,1), nrow=4),
                                treat.marginality)
rownames(systematic.marginality) <- systematic.terms
colnames(systematic.marginality) <- systematic.terms
choose <- chooseModel(current.asrt, systematic.marginality, denDF="algebraic", pos=TRUE)
current.asrt <- choose$asrtests.obj
#Check if any deviations are significant and, for those that are, go back to #fixed dates
current.asrt <- reparamSigDevn(current.asrt, choose$sig.terms, trend.num = "xDay", devn.fac = "Date", denDF = "algebraic")
## End(Not run)

**rmboundary.asrtests**

Removes any boundary or singular variance components from the fit stored in asrtests.obj and records their removal in an asrtests.object.

**Description**

Any terms specified in the random model that are estimated on the boundary or are singular and can be removed are removed from the fit stored in the asreml object stored in the asrtests.object. Terms that specify multiple parameters in the random model cannot be removed (e.g. terms specified using the at function with more than one level of the factor) and terms in residual model are not removed. Terms that can be removed are selected for removal in the following order based on whether they involve: (i) a dev function, (ii) only factors, (iii) an spl function, (iv) a pol function and (v) a lin function or a variable that is an integer or a numeric. It should be noted that this order of removal presumes that random deviation terms are specified via the dev function rather than via a random factor. Once the earliest of the above classes with a boundary term is identified, a term within this class is selected for removal. For all classes, except for factor-only terms, the smallest term with the largest number of variables/factors is removed. Amongst factor-only terms, the smallest term with the smallest number of variables/factors is removed. After each variance component is removed, a row for it is added to the test.summary data.frame and the model re-fitted. If there are further boundary or singular terms, one is removed using the above strategy. This process continues until there are no further boundary or singular variance components that are removable. Other types of boundary or singular terms, which cannot be removed, are reported in warning messages.
Usage

```r
## S3 method for class 'asrtests'
rmboundary(asrtests.obj, checkboundaryonly = FALSE,
            IClikelihood = "none", trace = FALSE, update = TRUE,
            set.terms = NULL, ignore.suffices = TRUE,
            bounds = "P", initial.values = NA, ...)
```

Arguments

- `asrtests.obj`: an `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary`.
- `checkboundaryonly`: If `TRUE` then boundary and singular terms are not removed by `rmboundary.asrtests`; a warning is issued instead.
- `IClikelihood`: A character that controls both the occurrence and the type of likelihood for information criterion in the `test.summary` of the new `asrtests.object`. If `none`, none are included. Otherwise, if `REML`, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if `full`, then the AIC and BIC based on the full likelihood are included. (See also `infoCriteria.asreml`.)
- `trace`: If `TRUE` then partial iteration details are displayed when ASReml-R functions are invoked; if `FALSE` then no output is displayed.
- `update`: If `TRUE` then `update.asreml` is called to fit the model with any boundary terms removed. In doing this the arguments `R.param` and `G.param` are set to those in the `asreml` object stored in `asrtests.obj` so that the values from the previous model are used as starting values. If `FALSE` then a call is made to `asreml` in which the only changes from the previous call are that (i) the terms for boundary variance components are removed from the models and (ii) modifications specified via `...` are made.
- `set.terms`: A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the `vparameters` component of the `asreml.obj` component in the `asrtests.object`.
- `ignore.suffices`: A logical vector specifying whether the suffices of the `asreml`-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of `terms`. If `TRUE` for an element of `terms`, the suffices are stripped from the `asreml`-assigned names. If `FALSE` for an element of `terms`, the element must exactly match an `asreml`-assigned name for a variance term. This vector must be of length one or the same length as `terms`. If it is of length one then the same action is applied to the `asreml`-assigned suffices for all the terms in `terms`.
- `bounds`: A character vector specifying the bounds to be applied to the terms specified in `set.terms`. This vector must be of length one or the same length as `set.terms`. If it is of length one then the same constraint is applied to all the terms in `set.terms`. If any of the bounds are equal to NA then they are left unchanged for those terms.
- `initial.values`: A character vector specifying the initial values for the terms specified in `terms`. This vector must be of length one or the same length as `set.terms`. If it is of length one then the same initial value is applied to all the terms in `set.terms`. If any of the initial values are equal to NA then they are left unchanged for those terms.

... further arguments passed to `asreml`. 
Value

An `asrtests.object` containing the components (i) `asreml.obj`, (ii) `wald.tab`, and (iii) `test.summary`.

Author(s)

Chris Brien

See Also

`as.asrtests`, `changeTerms.asrtests`, `testranfix.asrtests`, `testresidual.asrtests`, `newfit.asreml`, `reparamSigDevn.asrtests`, `chooseModel.asrtests`

Examples

```r
## Not run:
current.asrt <- rmboundary(current.asrt)
## End(Not run)
```

Description

Takes an unevaluated call and evaluates the call after setting the bounds and initial values for the terms specified in `terms`. The elements of `terms` are matched with those generated by `asreml` and used, for example, in the `varcomp` component of a `summary.asreml` object. These names generally include descriptive suffices. To match an element of `terms` that includes such a suffix, set `ignore.suffices` to `FALSE` so that a literal match between the element and the assigned names is sought.

Usage

```r
## S3 method for class 'call'
setvarianceterms(call, terms, ignore.suffices = TRUE,
                  bounds = "P", initial.values = NA, ...)
```

Arguments

- `call` an unevaluated call to `asreml`. One way to create such a call is to use the `call` function with its `name` argument set to "asreml". Another is to obtain it from the `call` component of an `asreml` object (e.g. `call <- asreml.obj$call`).
- `terms` A character vector specifying the terms that are to have bounds and/or initial values specified. The names must match those in the `vparameters` component of the `asreml.obj` component in the `asrtests.object`.

setvarianceterms.call allows the setting of bounds and initial values for terms in the random and residual arguments of an asreml call, with the resulting call being evaluated.
ignore.suffices

A logical vector specifying whether the suffices of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in terms.

bounds

A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

The codes used by ASReml are:

- B - fixed at a boundary;
- F - fixed by the user;
- P - positive definite;
- C - Constrained by user;
- U - unbounded.

initial.values

A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

... additional arguments to be added to the call, or arguments in the call with changed values.

Value

An asreml object.

Author(s)

Chris Brien

References


See Also

update.asreml

Examples

```r
## Not run:
m1.call <- call("asreml",
  fixed = Height ~ (Block + Irrig)*csDay.num,
  random= ~ spl(csDay.num)/(Irrig+Block)
  + dev(csDay.num)
  + str(~Block:Plot/csDay.num, ~us(2):id(20))
```
simulate.asreml

+ Block:Plot:spl(csDay.num),
  data=quote(dat)) ##use quote to stop evaluation of dat here
terms <- c("Block:Plot\+Block:Plot:csDay.num!us(2);2:1", "R!variance")
m1.asreml <- setvarianceterms(m1.call, terms, bounds=c("U","P"),
  initial=c(NA,3), ignore.suffices=c(FALSE,TRUE))

summary(m1.asreml)

## End(Not run)

simulate.asreml

Produce sets of simulated data from a multivariate normal distribution
and save quantities related to the simulated data

Description

Produce in parallel sets of simulated data corresponding to an asreml model, along with its fitted
values and residuals. A variance matrix $V$, corresponding to the random and residual models must
be supplied. What to save is specified by the which argument.

Usage

## S3 method for class 'asreml'
simulate(object, nsim=100, seed = NULL, means=NULL, V, tolerance = 1E-10,
  update = TRUE, trace = FALSE, which="data", units = "ignore",
  ncores = detectCores(), ...)

Arguments

object An asreml object from a call to asreml in which the data argument has been set.
means The vector of means to be used in generating simulated data sets. If it is NULL,
  the fitted values based on object are used. It must be the same length as the
  response variable for object.
V The fitted variance matrix, i.e. having the pattern and values that conform to
  the model fit stored in the supplied object.
nsim The number of data sets to be simulated.
seed A single value, interpreted as an integer, that specifies the starting value of the
  random number generator. The "L’Ecuyer-CMRG" random generator is used
  and nextRNGStream is used to seed each core from the original seed.
tolerance The value such that eigenvalues less than it are considered to be zero.
update If TRUE then the arguments R.param and G.param are set to those in the asreml
  object supplied in object so that the values from the original model are used
  as starting values. If FALSE then calls are made to asreml in which the only
  changes from the previous call are (i) the model is fitted to simulated data and
  (ii) modifications specified via ... are made, except that changes cannot be
  made to any of the models.
trace If TRUE then partial iteration details are displayed when ASReml-R functions are
  invoked; if FALSE then no output is displayed.
which

The quantities from the simulated data set to be stored. Any combination of "response", "residuals" and "fitted", or "all". If residuals and/or fitted is specified, those for the analysis stored in object will be added to the data.frame nominated in the data argument of object and the modified data.frame added as a component named data in the list that is the value returned by the function.

units

A character indicating whether the BLUPs for units are added to the residuals when this reserved factor is included in the random model. Possible values are addtoresiduals and ignore.

ncores

A numeric specifying the number of cores to use in doing the simulations.

... Other arguments specifying the number of cores to use in doing the simulations.

Details

Generate nsim sets of data and analyse them using asreml using the model in object, performing the generation and analysis of several sets in parallel. Note, if the analysis for a data set does not converge in maxiter iterations, it is discarded and a replacement data set generated. The value of maxiter can be specified in the call to simulate.asreml. The fitted values and residuals are extracted as required. If aom = TRUE when the simulated data are analysed, standardised conditional residuals are stored. If which includes residuals or fitted, the specified quantities for the observed data are added to the data.frame on which the fit in object is based.

Value

A list with the following components whose presence depends on the setting of which:

1. observed: present if which includes residuals or fitted, in which case it will be the data.frame on which the fit in object is based, with residuals and/or fitted.
2. data: present if which includes data, a data.frame containing the simulated data sets.
3. fitted: present if which includes fitted, a data.frame containing the fitted values from the analyses of the simulated data sets.
4. residuals: present if which includes residuals, a data.frame containing the residuals from the analyses of the simulated data sets.

Author(s)

Chris Brien

See Also

asreml, variofaces.asreml, plotVariofaces.data.frame, set.seed.

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety, 
   random = ~ Row + Column + units, 
   residual = ~ ar1(Row):ar1(Column), 
   data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL)
```
current.asrt <- rmboundary.asrtests(current.asrt)
# Form variance matrix based on estimated variance parameters
s2 <- current.asr$sigma2
gamma.Row <- current.asr$gammas[1]
gamma.unit <- current.asr$gammas[2]
rho.r <- current.asr$gammas[4]
rho.c <- current.asr$gammas[5]
row.ar1 <- mat.ar1(order=10, rho=rho.r)
col.ar1 <- mat.ar1(order=15, rho=rho.c)
V <- gamma.Row * fac.sumop(Wheat.dat$Row) +
    gamma.unit * diag(1, nrow=150, ncol=150) +
    mat.dirprod(col.ar1, row.ar1)
V <- s2*V

#Produce residuals from 100 simulated data sets
resid <- simulate(current.asr, V=V, which="residuals", ncores = 2)

## End(Not run)

### sort.alldiffs

sort.alldiffs <- function(x, decreasing = FALSE, classify = NULL,
                         sortFactor = NULL, sortParallelToCombo = NULL,
                         sortNestingFactor = NULL, sortOrder = NULL,...)

Arguments

- x: An alldiffs.object.
- decreasing: A logical passed to order that determines whether the order is for increasing or decreasing magnitude of the predicted values.
- classify: A character string giving the variables that define the margins of the multiway table that was predicted. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator. If NULL, it will be obtained from the classify attribute of the as.alldiffs object supplied through x.
sortFactor  A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.

sortParallelToCombo  A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the supplied list is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.

sortNestingFactor  A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.

sortOrder  A character vector whose length is the same as the number of levels for sortFactor in the predictions component of the alldiffs.object. It specifies the desired order of the levels in the reordered components of the alldiffs.object. The argument sortParallelToCombo is ignored.

The following creates a sortOrder vector levs for factor f based on the values in x: levs <- levels(f)[order(x)].

Details

The basic technique is to change the order of the levels of the sortFactor within the predictions and, if present, backtransforms components so that they are ordered for a subset of predicted values, one for each levels of the sortFactor. When the classify term consists of more than one variable then a subset of one combination of the values of variables other than the sortFactor, the sortParallelToCombo combination, must be chosen for determining the order of the sortFactor levels. Then the sorting of the rows (and columns) will be in parallel within each combination of the values of sortParallelToCombo variables: the classify term, excluding the sortFactor.

Value

The alldiffs.object supplied with the following components, if present, sorted: predictions, vcov, backtransforms, differences, p.differences and sed. Also, the sortFactor and sortOrder attributes are set.

Author(s)

Chris Brien
See Also

as.alldiffs, allDifferences.data.frame, print.alldiffs,
sort.predictions.frame, renewClassify.alldiffs, redoErrorIntervals.alldiffs,
recalcLSD.alldiffs, predictPlus.asreml, predictPresent.asreml

Examples

## Halve WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18")

## Use asreml to get predictions and associated statistics

## Not run:
# Analyse pH
m1.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                 random = ~ Benches:MainPlots,
                 keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(m1.asr, NULL, NULL)
current.asrt <- as.asrtests(m1.asr)
current.asrt <- rmboundary(current.asrt)
m1.asr <- current.asrt$asreml.obj

# Get predictions and associated statistics
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
                                asreml.obj = m1.asr, tables = "none",
                                wald.tab = current.asrt$wald.tab,
                                present = c("Type", "Species", "Sources"))

# Use sort.alldiffs and save order for use with other response variables
TS.diffs.sort <- sort(TS.diffs, sortFactor = "Sources",
                      sortOrder = "sortOrder")

# Analyse Turbidity
m2.asr <- asreml(fixed = Turbidity ~ Benches + (Sources * (Type + Species)),
                 random = ~ Benches:MainPlots,
                 keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(m2.asr)

# Use pH sort.order to sort Turbidity alldiffs object
diffs2.sort <- predictPlus(m2.asr, classify = "Sources:Type",
                           pairwise = FALSE, error.intervals = "Stand",
                           tables = "none", present = c("Type", "Species", "Sources"),
                           sortFactor = "Sources",
                           sortOrder = sort.order)

## End(Not run)

## Use lmeTest and emmmeans to get predictions and associated statistics

if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
    # Analyse pH
    m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
                              (1|Benches:MainPlots),
                              ...}

data=na.omit(tmp))
TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
      se = "SE", interval.type = "CI",
      interval.names = c("lower.CL", "upper.CL"))
## Form an all.diffs object and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els,els]
TS.diffs <- allDifferences(predictions = TS.preds,
      classify = "Sources:Type",
      vcov = TS.vcov, tdf = den.df)
validAlldiffs(TS.diffs)
#Use sort.alldiffs and save order for use with other response variables
TS.diffs.sort <- sort(TS.diffs, sortFactor = "Sources",
      sortParallelToCombo = list(Type = "Control"))
sort.order <- attr(TS.diffs.sort, which = "sortOrder")
#Analyse Turbidity
m2.lmer <- lmerTest::lmer(Turbidity ~ Benches + (Sources * (Type + Species)) +
   (1|Benches:MainPlots),
   data=na.omit(tmp))
TS.emm <- emmeans::emmeans(m2.lmer, specs = ~ Sources:Type)
TS.preds <- summary(TS.emm)
den.df <- min(TS.preds$df, na.rm = TRUE)
## Modify TS.preds to be compatible with a predictions.frame
TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
      se = "SE", interval.type = "CI",
      interval.names = c("lower.CL", "upper.CL"))
## Form an all.diffs object, sorting it using the pH sort.order and check its validity
els <- as.numeric(rownames(TS.preds))
TS.vcov <- vcov(TS.emm)[els,els]
TS.diffs2.sort <- allDifferences(predictions = TS.preds,
      classify = "Sources:Type",
      vcov = TS.vcov, tdf = den.df,
      sortFactor = "Sources",
      sortOrder = sort.order)
validAlldiffs(TS.diffs2.sort)
}

---

**sort.predictions.frame**

Sorts a **predictions.frame** according to the predicted values associated with a factor.

**Description**

Sorts the rows of a **predictions.frame** according to the predicted values in the predictions.frame. These predicted values are generally obtained using `predict.asreml` by specifying a `classify`
term comprised of one or more variables. Generally, the values associated with one variable are sorted in parallel within each combination of values of the other variables. When there is more than one variable in the classify term, the sorting is controlled using one or more of sortFactor, sortParallelToCombo and sortOrder. If there is only one variable in the classify then the predictions.frame is sorted according to the order of the complete set of predictions.

Usage

## S3 method for class 'predictions.frame'

\[
\text{sort}(x, \text{decreasing} = \text{FALSE}, \text{classify}, \text{sortFactor} = \text{NULL}, \\
\text{sortParallelToCombo} = \text{NULL}, \text{sortNestingFactor} = \text{NULL}, \\
\text{sortOrder} = \text{NULL}, \ldots)
\]

Arguments

- **x** A predictions.frame.
- **decreasing** A logical passed to order that determines whether the order is for increasing or decreasing magnitude of the predicted values.
- **classify** A character string giving the variables that define the margins of the multiway table that was predicted. Multiway tables are specified by forming an interaction type term from the classifying variables, that is, separating the variable names with the : operator.
- **sortFactor** A character containing the name of the factor that indexes the set of predicted values that determines the sorting of the components. If there is only one variable in the classify term then sortFactor can be NULL and the order is defined by the complete set of predicted values. If there is more than one variable in the classify term then sortFactor must be set. In this case the sortFactor is sorted in the same order within each combination of the values of the sortParallelToCombo variables: the classify variables, excluding the sortFactor. There should be only one predicted value for each unique value of sortFactor within each set defined by a combination of the values of the classify variables, excluding the sortFactor factor. The order to use is determined by either sortParallelToCombo or sortOrder.
- **sortParallelToCombo** A list that specifies a combination of the values of the factors and numerics, excluding sortFactor, that are in classify. Each of the components of the supplied list is named for a classify variable and specifies a single value for it. The combination of this set of values will be used to define a subset of the predicted values whose order will define the order of sortFactor. Each of the other combinations of the values of the factors and numerics will be sorted in parallel. If sortParallelToCombo is NULL then the first value of each classify variable, except for the sortFactor factor, in the predictions component is used to define sortParallelToCombo. If there is only one variable in the classify then sortParallelToCombo is ignored.
- **sortNestingFactor** A character containing the name of the factor that defines groups of the sortFactor within which the predicted values are to be ordered. If there is only one variable in the classify then sortNestingFactor is ignored.
- **sortOrder** A character vector whose length is the same as the number of levels for sortFactor in the predictions.frame. It specifies the desired order of the levels in the reordered the predictions.frame. The argument sortParallelToCombo is ignored.
The following creates a sortOrder vector `levs` for factor `f` based on the values in `x`: `levs <- levels(f)[order(x)]`.

... further arguments passed to or from other methods. Not used at present.

**Details**

The basic technique is to change the order of the levels of the `sortFactor` within the `predictions.frame` so that they are ordered for a subset of predicted values, one for each levels of the `sortFactor`. When the `classify` term consists of more than one variable then a subset of one combination of the values of variables other than the `sortFactor`, the `sortParallelToCombo` combination, must be chosen for determining the order of the `sortFactor` levels. Then the sorting of the rows (and columns) will be in parallel within each combination of the values of `sortParallelToCombo` variables: the `classify` term, excluding the `sortFactor`.

**Value**

The sorted `predictions.frame`. Also, the `sortFactor` and `sortOrder` attributes are set.

**Author(s)**

Chris Brien

**See Also**

`as.predictions.frame`, `print.predictions.frame`, `sort.alldiffs`, `predictPlus.asreml`, `predictPresent.asreml`

**Examples**

```r
##Halve WaterRunoff data to reduce time to execute
data(WaterRunoff.dat)
tmp <- subset(WaterRunoff.dat, Date == "05-18")
##Use asreml to get predictions and associated statistics
## Not run:
#Analyse pH
m1.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
                 random = ~ Benches:MainPlots,
                 keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(m1.asr, NULL, NULL)
current.asrt <- as.asrtests(m1.asr)
current.asrt <- rmboundary(current.asrt)
m1.asr <- current.asrt$asreml.obj

#Get predictions and associated statistics
TS.diffs <- predictPlus.asreml(classify = "Sources:Type",
                                 asreml.obj = m1.asr, tables = "none",
                                 wald.tab = current.asrt$wald.tab,
                                 present = c("Type","Species","Sources"))

#Use sort.predictions.frame and save order for use with other response variables
TS.preds <- TS.diffs$predictions
TS.preds.sort <- sort(TS.preds, sortFactor = "Sources",
                      sortParallelToCombo = list(Type = "Control"))
```
sort.order <- attr(TS.preds.sort, which = "sortOrder")

# Analyse Turbidity
m2.asr <- asreml(fixed = Turbidity ~ Benches + (Sources * (Type + Species)),
    random = ~ Benches:MainPlots,
    keep.order=TRUE, data= tmp)
current.asrt <- as.asrtests(m2.asr)
# Use pH sort.order to sort Turbidity alldiffs object
TS.diffs2 <- predictPlus(m2.asr, classify = "Sources:Type",
    pairwise = FALSE, error.intervals = "Stand",
    tables = "none", present = c("Type","Species","Sources"))
TS.preds2 <- TS.diffs2$predictions
TS.preds2.sort <- sort(TS.preds, sortFactor = "Sources", sortOder = sort.order)

## Use lmeTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) &
    requireNamespace("emmeans", quietly = TRUE))
{
    # Analyse pH
    m1.lmer <- lmerTest::lmer(pH ~ Benches + (Sources * (Type + Species)) +
            (1|Benches:MainPlots),
            data=na.omit(tmp))
    TS.emm <- emmeans::emmeans(m1.lmer, specs = ~ Sources:Type)
    TS.preds <- summary(TS.emm)
    den.df <- min(TS.preds$df, na.rm = TRUE)
    ## Modify TS.preds to be compatible with a predictions.frame
    TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
        se = "SE", interval.type = "CI",
        interval.names = c("lower.CL", "upper.CL"))

    # Use sort.predictions.frame and save order for use with other response variables
    TS.preds.sort <- sort(TS.preds, classify = "Sources:Type", sortFactor = "Sources",
        sortParallelToCombo = list(Type = "Control"))
    sort.order <- attr(TS.preds.sort, which = "sortOrder")

    # Analyse Turbidity
    m2.lmer <- lmerTest::lmer(Turbidity ~ Benches + (Sources * (Type + Species)) +
            (1|Benches:MainPlots),
            data=na.omit(tmp))
    TS.emm <- emmeans::emmeans(m2.lmer, specs = ~ Sources:Type)
    TS.preds <- summary(TS.emm)
    den.df <- min(TS.preds$df, na.rm = TRUE)
    ## Modify TS.preds to be compatible with a predictions.frame
    TS.preds <- as.predictions.frame(TS.preds, predictions = "emmean",
        se = "SE", interval.type = "CI",
        interval.names = c("lower.CL", "upper.CL"))
}

subset.alldiffs

Subsets the components in an alldiffs.object according to the supplied condition.
subset.alldiffs

Description

Subsets each of the components of an alldiffs.object. The subset is determined by applying the condition to the prediction component to determine which of its rows are to be included in the subset. Then, if present, this subset is applied to the rows of backtransforms and to the rows and columns of differences, p.differences and sed components. In addition, if sed is present, recalcLSD.alldiffs is called to recalculate the values in the LSD.frame stored in the LSD component, with any arguments supplied via the ... argument passed on it.

The select argument of subset is not implemented, but can be achieved for variables in the classify using the rmClassifyVars argument.

Usage

## S3 method for class 'alldiffs'
subset(x, subset = rep(TRUE, nrow(x$predictions)),
       rmClassifyVars = NULL, ...)

Arguments

x
An alldiffs.object.

subset
A logical that determines rows of the predictions component of x to be included in the subset. By default all rows are included.

rmClassifyVars
A character that contains the names of the variables in the classify attribute of x that are to be removed from the predictions data.frame and the names of the dimensions of the other components of x. In doing this, the combinations of the remaining classify variables must uniquely index the predictions.

... further arguments passed to recalcLSD.alldiffs.

Value

An alldiffs.object with the following components of the supplied alldiffs.object subsetted, if present in the original object: predictions, vcov, backtransforms, differences, p.differences and sed. In addition, if sed is present, the LSD.frame in the LSD component will be recalculated.

Author(s)

Chris Brien

See Also

as.alldiffs, allDifferences.data.frame, print.alldiffs, sort.alldiffs,
redoErrorIntervals.alldiffs, recalcLSD.alldiffs,
predictPlus.asreml, predictPresent.asreml

Examples

data(WaterRunoff.dat)

##Use asreml to get predictions and associated statistics

## Not run:
asreml.options(keep.order = TRUE) #required for asreml-R4 only
current.asr <- asreml(fixed = pH ~ Benches + (Sources * (Type + Species)),
       random = ~ Benches:MainPlots,
subset.list

Forms a list that contains a subset of the components of the supplied list.

### Description

Select components of a list specified by a list of numbers or names, or by a logical indicating for each component of the list whether or not it is to be retained.

### Usage

```r
# S3 method for class 'list'
subset(x, select = 1:length(x), ...)
```
Arguments

x  An \texttt{list} object.

select  A \texttt{numeric} or \texttt{character} that lists or names the components of the \texttt{list} \texttt{x} that are to be retained in the subset. It can also be a \texttt{logical} that is the same length as \texttt{x} and indicates whether or not a component is to be retained.

...  further arguments passed to or from other methods. Not used at present.

Value

A \texttt{list} with just the subset of the components from \texttt{x}. If the components of \texttt{x} are named, then these names are retained in the subset \texttt{list}.

Author(s)

Chris Brien

See Also

\texttt{subset.alldiffs}

Examples

\begin{verbatim}
x <- list(1:3,letters[1:3],LETTERS[1:3])
y <- subset.list(x, select = c(1,3))
y <- subset.list(x, select = c(TRUE,FALSE,TRUE))

names(x) <- LETTERS[1:3]
y <- subset.list(x, select = c(1,3))
z <- subset.list(x, select = LETTERS[c(1,3)])
x <- list(1:3,letters[1:3],LETTERS[1:3])
names(x)[c(1,3)] <- LETTERS[c(1,3)]
z <- subset.list(x, select = c(1,2))
v <- subset.list(x)
\end{verbatim}

testranfix.asrtests  

Tests for a single fixed or random term in model fitted using \texttt{asreml} and records the result in an \texttt{asrtests.object}.

Description

Tests for a single term, using a REML ratio test (REMLRT) for a random term or based on Wald statistics for a fixed term. The term must be in the fitted model. A random term is removed from the model fit and a REMLRT is performed using \texttt{REMLRT.asreml}. It compares the fit of the model in \texttt{asreml.obj} and the newly fitted model without the term. If the newly fitted model is retained, any boundary terms are then removed using \texttt{rmboundary.asrtests}. For a fixed term, the probability of the Wald statistics is extracted from the pseudo-anova table produced by \texttt{wald.asreml}. If this is available in the \texttt{asrtests.object}, it is used; otherwise \texttt{wald.asreml} is called to add it to the \texttt{asrtests.object}. Whether nonsignificant terms are dropped is controlled by \texttt{drop.ran.ns} for random terms and \texttt{drop.fix.ns} for fixed terms. A row is added to the \texttt{test.summary data.frame} for the term that is tested.
Usage

## S3 method for class 'asrtests'
testranfix(asrtests.obj, term=NULL, alpha = 0.05,
  allow.unconverged = TRUE, allow.fixedcorrelation = TRUE,
  checkboundaryonly = FALSE,
  drop.ran.ns = TRUE, positive.zero = FALSE,
  bound.test.parameters = "none",
  bound.exclusions = c("F","B","S","C"), REMLDF = NULL,
  drop.fix.ns = FALSE, denDF="numeric", dDF.na = "none",
  dDF.values = NULL, IClikelihood = "none",
  trace = FALSE, update = TRUE,
  set.terms = NULL, ignore.suffices = TRUE,
  bounds = "P", initial.values = NA, ...)

Arguments

asrtests.obj An \texttt{asrtests.object} containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary.

term A single model term that is valid in asreml, stored as a character. The names of fixed terms must match those in the wald.tab component of the asrtests.obj, while the names of random terms must match those in the vparameters component of the asreml.obj component in the asrtests.obj.

alpha The significance level for the test.

allow.unconverged A logical indicating whether to accept a new model even when it does not converge. If FALSE, it will be checked whether convergence can be achieved with the removal of any boundary random terms; random terms will be retested if terms are removed. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

allow.fixedcorrelation A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

checkboundaryonly If TRUE then boundary and singular terms are not removed by \texttt{rmboundary.asrtests}; a warning is issued instead.

drop.ran.ns A logical indicating whether to drop a random term from the model when it is nonsignificant. Note that multiple terms specified using a single asreml::at function will only be dropped as a whole. If the term was specified using an asreml::at function with a single level, then it can be removed and either the level itself or its \texttt{numeric} position in the levels returned by the \texttt{levels} function can be specified in term.

positive.zero Indicates whether the hypothesized values for the variance components being tested are on the boundary of the parameter space. For example, this is true for positively-constrained variance components that, under the reduced model, are zero. This argument does not need to be set if bound.test.parameters is set.
bound.test.parameters
Indicates whether for the variance components being tested, at least some of the hypothesized values are on the boundary of the parameter space. The possibilities are "none", "onlybound" and "one-and-one". The default is "none", although if it is set to "none" and positive.zero is TRUE then bound.test.parameters is taken to be "onlybound". When bound.test.parameters is set to "one-and-one", it signifies that there are two parameters being tested, one of which is bound and the other is not. For example, the latter is true for testing a covariance and a positively-constrained variance component that, under the reduced model, are zero.

bound.exclusions
A character specifying one or more bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in using `REMLRT.asreml`. If set to NULL then none will be excluded.

REMLDF
A numeric giving the difference in the number of variance parameters whose estimates are not of the type specified in bound.exclusions for two models being compared in a REML ratio test using `REMLRT.asreml`. If NULL then this is determined from the information in the asreml object for the two models.

drop.fix.ns
A logical indicating whether to drop a fixed term from the model when it is nonsignificant. Note that multiple terms specified using a single asreml::at function can only be dropped as a whole. If the term was specified using an asreml::at function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the levels function can be specified.

denDF
Specifies the method to use in computing approximate denominator degrees of freedom when wald.asreml is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

dDF.na
The method to use to obtain substitute denominator degrees of freedom when the numeric or algebraic methods produce an NA. If dDF.na = "none", no substitute denominator degrees of freedom are employed; if dDF.na = "residual", the residual degrees of freedom from asreml.obj$nedf are used; if dDF.na = "maximum", the maximum of those denDF that are available, excluding that for the Intercept, is used; if all denDF are NA, asreml.obj$nedf is used. If dDF.na = "supplied", a vector of values for the denominator degrees of freedom is to be supplied in dDF.values. Any other setting is ignored and a warning message produced. Generally, substituting these degrees of freedom is anticonservative in that it is likely that the degrees of freedom used will be too large.

dDF.values
A vector of values to be used when dDF.na = "supplied". Its values will be used when dDF in a test for a fixed effect is NA. This vector must be the same length as the number of fixed terms, including (Intercept) whose value could be NA.

IClikelihood
A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new asrtests.object. If none, none are included. Otherwise, if REML and family is set to asr_guassian (the default), then the AIC and BIC based on the Restricted Maximum Likelihood are included; if full and family is set to asr_guassian, then the AIC and BIC based on the full likelihood are included. If family is asr_binomial or
asr_poisson, with dispersion set to 1, the deviance is extracted from object and used to calculate the AIC and BIC. (See also infoCriteria.asreml.)

trace
If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update
If TRUE then update.asreml is called to fit the model to be tested. In doing this the arguments R.param and G.param are set to those in the asreml object stored in asrtests.obj so that the values from the previous model are used as starting values. If FALSE then a call is made to asreml in which the only changes from the previous call are that (i) models are modified for the supplied terms and (ii) modifications specified via ... are made.

set.terms
A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the vparameters component of the asreml.obj component in the new asrtests.object.

ignore.suffices
A logical vector specifying whether the suffices of the asreml-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the asreml-assigned names. If FALSE for an element of terms, the element must exactly match an asreml-assigned name for a variance term. This vector must be of length one or the same length as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in terms.

bounds
A character vector specifying the bounds to be applied to the terms specified in set.terms. This vector must be of length one or the same length as set.terms. If it is of length one then the same constraint is applied to all the terms in set.terms. If any of the bounds are equal to NA then they are left unchanged for those terms.

initial.values
A character vector specifying the initial values for the terms specified in terms. This vector must be of length one or the same length as terms. If it is of length one then the same initial value is applied to all the terms in terms. If any of the initial.values are equal to NA then they are left unchanged for those terms.

Further arguments passed to asreml, wald.asreml and as.asrtests.

Value
An asrtests.object containing the components (i) asreml.obj, (ii) wald.tab, and (iii) test.summary. If the term is not in the model, then the supplied asreml object will be returned. Also, reml.test will have the likelihood ratio and the p-value set to NA and the degrees of freedom to zero. Similarly, the row of test.summary for the term will have its name, DF set to NA, p-value set to NA, and action set to Absent.

Author(s)
Chris Brien

References
testresidual.asrtests

Fits a new residual formula, tests whether the change is significant and records the result in an asrtests.object.

Usage

## S3 method for class 'asrtests'
testresidual(asrtests.obj, terms=NULL, label = "R model", simpler = FALSE, alpha = 0.05, allow.unconverged = TRUE, allow.fixedcorrelation = TRUE, checkboundarystart = FALSE, positive.zero = FALSE, bound.test.parameters = "none", bound.exclusions = c("F","B","S","C"), REMLDF = NULL, denDF="numeric", IClikelihood = "none", update = TRUE, trace = FALSE, set.terms = NULL, ignore.suffices = TRUE, bounds = "P", initial.values = NA, ...)
Arguments

asrtests.obj

an asrtests.object for a fitted model that is a list containing the components (i) asreml.obj, (ii) wald.tab (iii) test.summary.

terms

A model for the residual argument in asreml-R4 (the rcov formula in older versions of asreml), stored as a character. To remove the model, enter “-(.)”.

label

A character string to use as the label in test.summary and which indicates what is being tested.

simpler

A logical indicating whether the new model to be fitted is simpler than the already fitted model whose fit is stored in asrtests.obj.

alpha

The significance level for the test.

allow.unconverged

A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit of the new model does not converge, the supplied asreml object is returned. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

allow.fixedcorrelation

A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

cHECKboundaryonly

If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead.

positive.zero

Indicates whether the hypothesized values for the variance components being tested are on the boundary of the parameter space. For example, this is true for positively-constrained variance components that, under the reduced model, are zero. This argument does not need to be set if bound.test.parameters is set.

bound.test.parameters

Indicates whether for the variance components being tested, at least some of the hypothesized values are on the boundary of the parameter space. The possibilities are “none”, “onlybound” and “one-and-one”. The default is "none", although if it is set to "none" and positive.zero is TRUE then bound.test.parameters is taken to be "onlybound". When bound.test.parameters is set to "one-and-one", it signifies that there are two parameters being tested, one of which is bound and the other is not. For example, the latter is true for testing a covariance and a positively-constrained variance component that, under the reduced model, are zero.

bound.exclusions

A character specifying one or more bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in using REMLRT.asreml. If set to NULL then none will be excluded.

REMLDF

A numeric giving the difference in the number of variance parameters whose estimates are not of the type specified in bound.exclusions for two models being compared in a REML ratio test using REMLRT.asreml. If NULL then this is determined from the information in the asreml object for the two models.
denDF  Specifies the method to use in computing approximate denominator degrees of freedom when \texttt{wald.asreml} is called. Can be \texttt{none} to suppress the computations, \texttt{numeric} for numerical methods, \texttt{algebraic} for algebraic methods or \texttt{default}, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

\texttt{IClikelihood}  A character that controls both the occurrence and the type of likelihood for information criterion in the \texttt{test.summary} of the new \texttt{asrtests.object}. If \texttt{none}, none are included. Otherwise, if \texttt{REML}, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if \texttt{full}, then the AIC and BIC based on the full likelihood are included. (See also \texttt{infoCriteria.asreml}.)

\texttt{update}  If \texttt{TRUE} then \texttt{update.asreml} is called to fit the model with the residual (\texttt{rcov}) model supplied in \texttt{terms}. In doing this the arguments \texttt{R.param} and \texttt{G.param} are set to those in the \texttt{asreml} object stored in \texttt{asrtests.obj} so that the values from the previous model are used as starting values. If \texttt{FALSE} then a call is made to \texttt{asreml} in which the only changes from the previous call are that (i) residual (\texttt{rcov}) model is that specified in \texttt{terms} and (ii) modifications specified via \ldots are made.

\texttt{trace}  If \texttt{TRUE} then partial iteration details are displayed when ASReml-R functions are invoked; if \texttt{FALSE} then no output is displayed.

\texttt{set.terms}  A \texttt{character} vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the \texttt{vparameters} component of the \texttt{asreml.obj} component in the new \texttt{asrtests.object}.

\texttt{ignore.suffixes}  A \texttt{logical} vector specifying whether the suffixes of the \texttt{asreml}-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of \texttt{terms}. If \texttt{TRUE} for an element of \texttt{terms}, the suffixes are stripped from the \texttt{asreml}-assigned names. If \texttt{FALSE} for an element of \texttt{terms}, the element must exactly match an \texttt{asreml}-assigned name for a variance term. This vector must be of length one or the same length as \texttt{terms}. If it is of length one then the same action is applied to the \texttt{asreml}-assigned suffixes for all the terms in \texttt{terms}.

\texttt{bounds}  A \texttt{character} vector specifying the bounds to be applied to the terms specified in \texttt{set.terms}. This vector must be of length one or the same length as \texttt{set.terms}. If it is of length one then the same constraint is applied to all the terms in \texttt{set.terms}. If any of the bounds are equal to \texttt{NA} then they are left unchanged for those terms.

\texttt{initial.values}  A character vector specifying the initial values for the terms specified in \texttt{terms}. This vector must be of length one or the same length as \texttt{terms}. If it is of length one then the same initial value is applied to all the terms in \texttt{terms}. If any of the initial.values are equal to \texttt{NA} then they are left unchanged for those terms.

\ldots  Further arguments passed to \texttt{asreml}, \texttt{wald.asreml} and \texttt{as.asrtests}.

\textbf{Value}

An \texttt{asrtests.object} containing the components (i) \texttt{asreml.obj}, (ii) \texttt{wald.tab}, and (iii) \texttt{test.summary}. If the term is not in the model, then the supplied \texttt{asreml.obj} will be returned. Also, \texttt{reml.test} will have the likelihood ratio and the p-value set to \texttt{NA} and the degrees of freedom to zero. Similarly, the row of \texttt{test.summary} for the term will have its name, a p-value set to \texttt{NA}, and action set to \texttt{Absent}.  

\texttt{testresidual.asrtests}
Author(s)

Chris Brien

References


See Also

c ran.asrtests, chooseModel.asrtests, REMLRT.asreml, rmboundary.asrtests, newfit.asreml, testswapran.asrtests, changeModelOnIC.asrtests, changeTerms.asrtests, reparamSigDevn.asrtests

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                       random = ~ Row + Column + units,
                       residual = ~ ar1(Row):ar1(Column),
                       data=Wheat.dat)
current.asr <- as.asrtests(current.asr, NULL, NULL)
current.asr <- rmboundary(current.asr)
# Test Row autocorrelation
current.asr <- testresidual(current.asr, "~ Row:ar1(Column)",
                            label="Row autocorrelation", simpler=TRUE)
print(current.asr)
## End(Not run)
```

Description

Tests, using a REMLRT, the significance of the difference between the current random model and one in which oldterms are dropped and newterms are added. The result is recorded in an asrtests.object.

Tests a new random model using asreml by removing oldterms and adding newterms. If simpler = FALSE the model to be fitted must be more complex than the one whose fit has been stored in asrtests.obj. That is, the new model must have more parameters. However, if simpler = TRUE the model to be fitted must be simpler than the one whose fit has been stored in asrtests.obj in that it must have fewer parameters. The test is a REML ratio test that is performed using REMLRT.asreml, which is only valid if the models are nested. It compares the newly fitted model with the fit of the model in asrtest.obj. A row is added to the test.summary data.frame using the supplied label. If the newly fitted model is retained, any boundary terms are then removed using rmboundary.asrtests. If the models are not nested, then using changeModelOnIC.asrtests may be the more appropriate approach for comparing models.
## Usage

```r
testswapran(asrtests.obj, oldterms = NULL, newterms = NULL, label = "Swap in random model", simpler = FALSE, alpha = 0.05, allow.unconverged = TRUE, allow.fixedcorrelation = TRUE, checkboundaryonly = FALSE, positive.zero = FALSE, bound.test.parameters = "none", bound.exclusions = c("F","B","S","C"), REMLDF = NULL, denDF="numeric", IClikelihood = "none", trace = FALSE, update = TRUE, set.terms = NULL, ignore.suffices = TRUE, bounds = "P", initial.values = NA, ...)
```

### Arguments

- **asrtests.obj**
  - an *asrtests.object* for a fitted model that is a list containing the components (i) asreml.obj, (ii) wald.tab (iii) test.summary.

- **oldterms**
  - Terms, stored as a character, that are to be removed from the random model using asreml. The names of the terms must match those in the vparameters component of the asreml.obj component in asrtests.obj. Note that multiple terms specified using a single asreml::at function can only be dropped as a whole. If the term was specified using an asreml::at function with a single level, then it can be removed and either the level itself or its numeric position in the levels returned by the levels function can be specified.

- **newterms**
  - Terms, stored as a character, that are to be added to the random model using asreml.

- **simpler**
  - A logical indicating whether the new model to be fitted after the changes made as a result of swapping oldterms for newterms, is simpler than the already fitted model whose fit is stored in asrtests.obj.

- **alpha**
  - The significance level for the test.

- **allow.unconverged**
  - A logical indicating whether to accept a new model even when it does not converge. If FALSE and the fit of the new model does not converge, the supplied asrtests.obj is returned. Also, if FALSE and the fit of the new model has converged, but that of the old model has not, the new model will be accepted.

- **allow.fixedcorrelation**
  - A logical indicating whether to accept a new model even when it contains correlations in the model whose values have been designated as fixed, bound or singular. If FALSE and the new model contains correlations whose values have not been able to be estimated, the supplied asrtests.obj is returned. The fit in the asreml.obj component of the supplied asrtests.obj will also be tested and a warning issued if both fixed correlations are found in it and allow.fixedcorrelation is FALSE.

- **checkboundaryonly**
  - If TRUE then boundary and singular terms are not removed by rmboundary.asrtests; a warning is issued instead.

- **label**
  - A character string to use as the label in test.summary and which indicates what is being tested.

- **positive.zero**
  - Indicates whether the hypothesized values for the variance components being tested are on the boundary of the parameter space. For example, this is true for
positively-constrained variance components that, under the reduced model, are zero. This argument does not need to be set if bound.test.parameters is set.

bound.test.parameters
Indicates whether for the variance components being tested, at least some of the hypothesized values are on the boundary of the parameter space. The possibilities are "none", "onlybound" and "one-and-one". The default is "none", although if it is set to "none" and positive.zero is TRUE then bound.test.parameters is taken to be "onlybound". When bound.test.parameters is set to "one-and-one", it signifies that there are two parameters being tested, one of which is bound and the other is not. For example, the latter is true for testing a covariance and a positively-constrained variance component that, under the reduced model, are zero.

bound.exclusions
A character specifying one or more bound (constraint) codes that will result in a variance parameter being excluded from the count of estimated variance parameters in using \texttt{REMLRT.asreml}. If set to NULL then none will be excluded.

REMLDF
A numeric giving the difference in the number of variance parameters whose estimates are not of the type specified in bound.exclusions for two models being compared in a REML ratio test using \texttt{REMLRT.asreml}. If NULL then this is determined from the information in the \texttt{asreml} object for the two models.

denDF
Specifies the method to use in computing approximate denominator degrees of freedom when \texttt{wald.asreml} is called. Can be none to suppress the computations, numeric for numerical methods, algebraic for algebraic methods or default, the default, to automatically choose numeric or algebraic computations depending on problem size. The denominator degrees of freedom are calculated according to Kenward and Roger (1997) for fixed terms in the dense part of the model.

IClikelihood
A character that controls both the occurrence and the type of likelihood for information criterion in the test.summary of the new \texttt{asrtests.object}. If none, none are included. Otherwise, if \texttt{REML}, then the AIC and BIC based on the Restricted Maximum Likelihood are included; if \texttt{full}, then the AIC and BIC based on the full likelihood are included. (See also \texttt{infoCriteria.asreml}.)

trace
If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

update
If TRUE then update.asreml is called to change the model. In doing this the arguments \texttt{R.param} and \texttt{G.param} are set to those in the \texttt{asreml} object stored in \texttt{asrtests.obj} so that the values from the previous model are used as starting values. If FALSE then a call is made to \texttt{asreml} in which the only changes from the previous call are that (i) models are modified for the supplied \texttt{oldterms} and \texttt{newterms}, and (ii) modifications specified via \texttt{...} are made.

set.terms
A character vector specifying the terms that are to have bounds and/or initial values set prior to fitting. The names must match those in the \texttt{vparameters} component of the \texttt{asreml.obj} component in the \texttt{asrtests.object}.

ignore.suffices
A logical vector specifying whether the suffices of the \texttt{asreml}-assigned names of the variance terms (i.e. the information to the right of an "!", other than "R!") is to be ignored in matching elements of terms. If TRUE for an element of terms, the suffices are stripped from the \texttt{asreml}-assigned names. If FALSE for an element of terms, the element must exactly match an \texttt{asreml}-assigned name for a variance term. This vector must be of length one or the same length...
as terms. If it is of length one then the same action is applied to the asreml-assigned suffices for all the terms in `terms`.

**bounds**
A character vector specifying the bounds to be applied to the terms specified in `set.terms`. This vector must be of length one or the same length as `set.terms`. If it is of length one then the same constraint is applied to all the terms in `set.terms`. If any of the bounds are equal to NA then they are left unchanged for those terms.

**initial.values**
A character vector specifying the initial values for the terms specified in `terms`. This vector must be of length one or the same length as `terms`. If it is of length one then the same initial value is applied to all the terms in `terms`. If any of the initial.values are equal to NA then they are left unchanged for those terms.

Further arguments passed to asreml, wald.asreml and as.asrtests.

**Value**
An asrtests.object for a fitted model that is a list containing the components (i) asreml.obj, (ii) wald.tab (iii) test.summary. If the term is not in the model, then the supplied asreml object will be returned. Also, reml.test will have the likelihood ratio and the p-value set to NA and the degrees of freedom to zero. Similarly, the row of test.summary for the term will have its name, a p-value set to NA, and action set to Absent.

**Author(s)**
Chris Brien

**References**

**See Also**

as.asrtests, chooseModel.asrtests, REMLRT.asreml, rmboundary.asrtests, newfit.asreml, testresidual.asrtests, changeModelOnIC.asrtests, changeTerms.asrtests, reparamSigDevn.asrtests

**Examples**

```r
## Not run:
current.asrt <- testswapran(current.asrt, oldterms = "str(~ Cart/xDays, ~us(2):id(184))", 
newterms = "Cart/xDays", pos = FALSE, 
label = "Intercept/Slope correlation",
simpler = TRUE)
print(current.asrt)
## End(Not run)
```
validAlldiffs

Checks that an object is a valid alldiffs object.

Description
Checks that an object is an alldiffs.object of S3-class alldiffs containing the components asreml.obj, wald.tab and test.summary.

Usage
validAlldiffs(object)

Arguments
object an alldiffs.object.

Value
TRUE or a character describing why the object is not a valid alldiffs.object.

Author(s)
Chris Brien

See Also
alldiffs.object, is.alldiffs, as.alldiffs, validPredictionsFrame, validAsrtests

Examples
data(Oats.dat)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & requireNamespace("emmmeans", quietly = TRUE))
{
  m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots), data=Oats.dat)
  Var.emm <- emmmeans::emmmeans(m1.lmer, specs = ~ Nitrogen:Variety)
  Var.preds <- summary(Var.emm)
  den.df <- min(Var.preds$df)
  ## Modify Var.preds to be compatible with a predictions.frame
  Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean", se = "SE", interval.type = "CI", interval.names = c("lower.CL", "upper.CL"))
  Var.vcov <- vcov(Var.emm)
  Var.sed <- NULL
  ## Form an all.diffs object
  Var.diffs <- as.alldiffs(predictions = Var.preds, classify = "Nitrogen:Variety", sed = Var.sed, vcov = Var.vcov, tdf = den.df)
  ## check the validity of Var.diffs
validAsrtests

Checks that an object is a valid asrtests object.

Description
Checks that an object is an asrtests.object of S3-class asrtests containing the components asreml.obj, wald.tab and test.summary.

Usage
validAsrtests(object)

Arguments
object an asrtests.object.

Value
TRUE or a character describing why the object is not a valid asrtests.object.

Author(s)
Chris Brien

See Also
asrtests.object, is.asrtests, as.asrtests, validPredictionsFrame, validAlldiffs

Examples
## Not run:
library(dae)
library(asreml)
library(asremlPlus)
## use ?Wheat.dat for data set details
data(Wheat.dat)

# Fit initial model
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety,
                      random = ~ Row + Column + units,
                      residual = ~ ar1(Row):ar1(Column),
                      data=Wheat.dat)

# Load current fit into an asrtests object
current.asrt <- as.asrtests(current.asr, NULL, NULL)

# check validity of current.asrt
validAsrtests(current.asrt)

## End(Not run)
validPredictionsFrame

Checks that an object is a valid predictions.frame.

Description

Checks that an object is a predictions.frame of S3-class data.frame that contains the columns predicted.value, standard.error and est.status.

Usage

validPredictionsFrame(object)

Arguments

object an predictions.frame.

Value

TRUE or a character describing why the object is not a valid predictions.frame.

Author(s)

Chris Brien

See Also

predictions.frame, is.predictions.frame, as.predictions.frame, validAsrtests, validAlldiffs

Examples

data(Oats.dat)

## Use asreml to get predictions and associated statistics
## Not run:
m1.asr <- asreml(Yield ~ Nitrogen*Variety, 
                random=~Blocks/Wplots, 
                data=Oats.dat)
current.asrt <- as.asrtests(m1.asr)
Var.pred <- asreml::predict.asreml(m1.asr, classify="Nitrogen:Variety", 
                                sed=TRUE)
if (getASRemlVersionLoaded(nchar = 1) == "3")
  Var.pred <- Var.pred$predictions
Var.preds <- as.predictions.frame(Var.pred$pvals, se = "std.error", 
                                 est.status = "status")

## End(Not run)

## Use lmerTest and emmmeans to get predictions and associated statistics
if (requireNamespace("lmerTest", quietly = TRUE) & 
    requireNamespace("emmeans", quietly = TRUE))
{
  # Code here
}
m1.lmer <- lmerTest::lmer(Yield ~ Nitrogen*Variety + (1|Blocks/Wplots),
                          data=Oats.dat)
Var.emm <- emmeans::emmeans(m1.lmer, specs = ~ Nitrogen:Variety)
Var.preds <- summary(Var.emm)
Var.preds <- as.predictions.frame(Var.preds, predictions = "emmean",
                                 se = "SE", interval.type = "CI",
                                 interval.names = c("lower.CL", "upper.CL"))
}
if (exists("Var.preds")){
  ## Check the class and validity of the predictions.frame
  is.predictions.frame(Var.preds)
  validPredictionsFrame(Var.preds)
}

variofaces.asreml  
Plots empirical variogram faces, including envelopes, as described by 

Description

A function that produces a plot for each face of an empirical 2D variogram based on residuals produced after the fitting of a model using the function asreml. It also adds envelopes to the plot by simulating data sets in parallel from a multivariate normal distribution with expectation equal to the fitted values obtained from the fixed and spline terms and variance matrix equal to the fitted variance matrix (Stefanova, Smith & Cullis, 2009). The plot is controlled by the residual model, which must consist of two factors corresponding to the two physical dimensions underlying the data. It can also have a third term involving the at or dsnum function that defines sections of the data, such as experiments in different environments. In this case, the two variogram faces are produced for each section.

Usage

## S3 method for class 'asreml'
variofaces(asreml.obj, means=NULL, V=NULL, nsim=100, seed = NULL,
            extra.matrix = NULL, ignore.terms = NULL, fixed.spline.terms = NULL,
            bound.exclusions = c("F","B","S","C"), tolerance=1E-10,
            units = "ignore", update = TRUE, trace = FALSE,
            graphics.device=NULL, ncores = detectCores(), ...)

Arguments

asreml.obj  
An asreml object from a call to asreml in which the data argument has been set.

means  
The vector of means to be used in generating simulated data sets. If it is NULL, the fitted values based on object are used. It must be the same length as the response variable for object.

V  
The fitted variance matrix, i.e. having the appropriate pattern and values given the model fitted to the observed data and the estimates of the parameters obtained. If V is NULL then estimateV.asreml is called to obtain it from asreml.obj
The number of data sets to be simulated in obtaining the envelopes.

A single value, interpreted as an integer, that specifies the starting value of the random number generator. The "L'Ecuyer-CMRG" random generator is used and nextRNGStream is used to seed each core from the original seed.

A matrix of order equal to the number of observations that is to be added to the variance matrix, the latter based on the information in asreml.obj. It is assumed that the sigma-parameterized values of the variance parameter estimates, such as is given in the varcomp component of summary.asreml, have been used in calculating extra.matrix; the values in the vparameters component of G.param and R.param may be either gamma- or sigma-parameterized. The argument extra.matrix can be used in conjunction with ignore.terms as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.

A character giving terms from either the random or residual models that are to be ignored in that their contributions to the variance is not to be included in the estimated matrix. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!). This can be used in conjunction with estimateV.asreml as a workaround to include components of the variance matrix for variance functions that have not been implemented in estimateV.

A character vector giving one or more spline terms in the random model that are regarded as fixed and so are to be ignored because they are not regarded as contributing to the variance. The term names are those given in the vparameters component of the asreml object or the varcomp component produced by summary.asreml, but only up to the first exclamation mark (!).

A character specifying one or more bound codes that will result in a variance parameter in the random model being excluded from contributing to the variance. If set to NULL then none will be excluded.

The value such that eigenvalues less than it are considered to be zero.

A character indicating whether the BLUPs for units are added to the residuals when this reserved factor is included in the random model. Possible values are addtoreiduals and ignore. If standardized conditional residuals are plotted and the BLUPs for units are to be added then it is the standardized BLUPs that are added.

If TRUE then the arguments R.param and G.param are set to those in the asreml object supplied in object so that the values from the original model are used as starting values. If FALSE then calls are made to asreml in which the only changes from the previous call are (i) the model is fitted to simulated data and (ii) modifications specified via ... are made, except that changes cannot be made to any of the models.

If TRUE then partial iteration details are displayed when ASReml-R functions are invoked; if FALSE then no output is displayed.

A character specifying a graphics device for plotting. The default is graphics.device = NULL, which will result in plots being produced on the current graphics device. Setting it to "windows", for example, will result in a windows graphics device being opened.

A numeric specifying the number of cores to use in doing the simulations.
Other arguments that are passed down to the function `asreml`. Changes to the models are not allowed. Other changes are dangerous and generally should be avoided.

Details

The residual model is scanned to ensure that it involves only two factors not included in the at function, and to see if it has a third factor in an at function. If so, the faces of the 2D variogram, each based on one of the two non-at factors, are derived from the residuals in the supplied `asreml` object using `asreml.variogram`, this yielding the observed variogram faces. If `aom` was set to `TRUE` for the `asreml` object, the standardized conditional residuals are used. Then `nsim` data sets are generated by adding the fitted values, extracted from the `asreml` object, to a vector of values randomly generated from a normal distribution with expectation zero and variance matrix $V$. Each data set is analyzed using the model in `object` and several sets are generated and analyzed in parallel. The variogram values for the faces are obtained using `asreml.variogram` stored. Note, if the analysis for a data set does not converge in `maxiter` iterations, it is discarded and a replacement data set generated. The value of `maxiter` can be specified in the call to `variofaces.asreml`. Plots are produced for each face and include the observed values and the 2.5%, 50% & 97.5% quantiles.

Value

A list with the following components:

1. **face1**: a `data.frame` containing the variogram values on which the plot for the first dimension is based.
2. **face2**: a `data.frame` containing the variogram values on which the plot for the second dimension is based.

Author(s)

Chris Brien

References


See Also

`asremlPlus-package`, `asreml.plotVariofaces.data.frame`, `simulate.asreml`, `set.seed`.

Examples

```r
## Not run:
data(Wheat.dat)
current.asr <- asreml(yield ~ Rep + WithinColPairs + Variety, 
                      random = ~ Row + Column + units, 
                      residual = ~ ar1(Row):ar1(Column), 
                      data=Wheat.dat)
current.asrt <- as.asrtests(current.asr, NULL, NULL) 
current.asrt <- rmboundary.asrtests(current.asrt) 
# Form variance matrix based on estimated variance parameters
s2 <- current.asr$sigma2
gamma.Row <- current.asr$gammas[1]
gamma.unit <- current.asr$gammas[2]
```
rho.r <- current.asr$gammas[4]
rho.c <- current.asr$gammas[5]
row.ar1 <- mat.ar1(order=10, rho=rho.r)
col.ar1 <- mat.ar1(order=15, rho=rho.c)
V <- gamma.Row * fac.sumop(Wheat.dat$Row) +
    gamma.unit * diag(1, nrow=150, ncol=150) +
    mat.dirprod(col.ar1, row.ar1)
V <- s2*V

#Produce variogram faces plot (Stefanaova et al, 2009)
variofaces(current.asr, V=V, ncores = 2)
## End(Not run)

---

WaterRunoff.dat

Data for an experiment to investigate the quality of water runoff over time

Description

This data is from an experiment to investigate the quality of water runoff. However, it has been modified to hide the true identity of the Species and Sources. It is used to provide executable examples of the functions listed under Examples.

Usage

data(WaterRunoff.dat)

Format

A data.frame containing 440 observations of 13 variables.

Author(s)

Chris Brien

Source

Kazemi, F. (pers. comm.)

See Also

chooseModel.asrtests, reparamSigDevn.asrtests, plotPredictions.data.frame, predictPlus.asreml, predictPresent.asreml
Wheat.dat

Data for a 1976 experiment to investigate 25 varieties of wheat

Description

The data appears in Gilmour et al. (1995) and is from a field experiment designed to compare the performance of 25 varieties of spring wheat. An analysis of it using asreml is presented by Butler et al. (2018, Section 7.6), although they suggest that it is a barley experiment. It is used in the Wheat vignettes [Enter vignette(package = “asremlPlus”)] as an executable example of the use of the asremlPlus to analyse a data set.

The experiment was conducted at Slate Hall Farm, UK, in 1976 and was designed as a balanced lattice square with 6 replicates laid out in a $10 \times 15$ rectangular grid. The columns in the data frame are: Rep, Row, Column, WithinColPairs, Variety, yield. The response variable is the grain yield.

Usage

data(Wheat.dat)

Format

A data.frame containing 150 observations of 6 variables.

Author(s)

Chris Brien

Source


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