Package ‘DHARMa’

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Title Residual Diagnostics for Hierarchical (Multi-Level / Mixed) Regression Models

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Description The 'DHARMa' package uses a simulation-based approach to create readily interpretable scaled (quantile) residuals for fitted (generalized) linear mixed models. Currently supported are linear and generalized linear (mixed) models from 'lme4' (classes 'lmerMod', 'glmerMod'), 'glmmTMB' and 'spaMM', generalized additive models ('gam' from 'mgcv'), 'glm' (including 'negbin' from 'MASS', but excluding quasi-distributions) and 'lm' model classes. Moreover, externally created simulations, e.g. posterior predictive simulations from Bayesian software such as 'JAGS', 'STAN', or 'BUGS' can be processed as well. The resulting residuals are standardized to values between 0 and 1 and can be interpreted as intuitively as residuals from a linear regression. The package also provides a number of plot and test functions for typical model misspecification problems, such as over/underdispersion, zero-inflation, and residual spatial and temporal autocorrelation.

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**createData**

_Simulate test data_

**Description**

This function creates synthetic dataset with various problems such as overdispersion, zero-inflation, etc.

**Usage**

```r
createData(sampleSize = 100, intercept = 0, fixedEffects = 1,
quadraticFixedEffects = NULL, numGroups = 10, randomEffectVariance = 1,
overdispersion = 0, family = poisson(), scale = 1, cor = 0,
roundPoissonVariance = NULL, pZeroInflation = 0, binomialTrials = 1,
temporalAutocorrelation = 0, spatialAutocorrelation = 0,
factorResponse = F, replicates = 1, hasNA = F)
```

**Arguments**

- `sampleSize`: sample size of the dataset
- `intercept`: intercept (linear scale)
- `fixedEffects`: vector of fixed effects (linear scale)
- `quadraticFixedEffects`: vector of quadratic fixed effects (linear scale)
- `numGroups`: number of groups for the random effect
- `randomEffectVariance`: variance of the random effect (intercept)
- `overdispersion`: if this is a numeric value, it will be used as the sd of a random normal variate that is added to the linear predictor. Alternatively, a random function can be provided that takes as input the linear predictor.
- `family`: family
- `scale`: scale if the distribution has a scale (e.g. sd for the Gaussian)
- `cor`: correlation between predictors
- `roundPoissonVariance`: if set, this creates a uniform noise on the poission response. The aim of this is to create heteroscedasticity
- `pZeroInflation`: probability to set any data point to zero
- `binomialTrials`: Number of trials for the binomial. Only active if family == binomial
- `temporalAutocorrelation`: strength of temporalAutocorrelation
- `spatialAutocorrelation`: strength of spatial Autocorrelation
- `factorResponse`: should the response be transformed to a factor (intended to be used for 0/1 data)
- `replicates`: number of datasets to create
- `hasNA`: should an NA be added to the environmental predictor (for test purposes)
createDHARMa

Create a DHARMa object from hand-coded simulations or Bayesian posterior predictive simulations

createDHARMa(simulatedResponse, observedResponse, 
  fittedPredictedResponse = NULL, integerResponse = F, seed = 123, 
  method = c("PIT", "traditional"))
createDHARMA

Arguments

simulatedResponse
matrix of observations simulated from the fitted model - row index for observations and column index for simulations

observedResponse
true observations

fittedPredictedResponse
optional fitted predicted response. For Bayesian posterior predictive simulations, using the median posterior prediction as fittedPredictedResponse is recommended. If not provided, the mean simulatedResponse will be used.

integerResponse
if T, noise will be added at to the residuals to maintain a uniform expectations for integer responses (such as Poisson or Binomial). Unlike in simulateResiduals, the nature of the data is not automatically detected, so this MUST be set by the user appropriately

seed
the random seed to be used within DHARMA. The default setting, recommended for most users, is keep the random seed on a fixed value 123. This means that you will always get the same randomization and thus the same result when running the same code. NULL = no new seed is set, but previous random state will be restored after simulation. FALSE = no seed is set, and random state will not be restored. The latter two options are only recommended for simulation experiments. See vignette for details.

method
the quantile randomization method used. The two options implemented at the moment are probability integral transform (PIT-) residuals (current default), and the "traditional" randomization procedure, that was used in DHARMA until version 0.3.0. For details, see getQuantile

Details

The use of this function is to convert simulated residuals (e.g. from a point estimate, or Bayesian p-values) to a DHARMA object, to make use of the plotting / test functions in DHARMA

Note

Either scaled residuals or (simulatedResponse AND observed response) have to be provided

Examples

```r
# READING IN HAND-CODED SIMULATIONS

testData = createData(sampleSize = 50, randomEffectVariance = 0)
fittedModel <- glm(observedResponse ~ Environment1, data = testData, family = "poisson")

# in DHARMA, using the simulate.glm function of glm
sims = simulateResiduals(fittedModel)
plot(sims, quantreg = FALSE)

# Doing the same with a handcoded simulate function.
# of course this code will only work with a 1-par glm model
```
simulateMyfit <- function(n=10, fittedModel){
  int = coef(fittedModel)[1]
  slo = coef(fittedModel)[2]
  pred = exp(int + slo * testData$Environment1)
  predSim = replicate(n, rpois(length(pred), pred))
  return(predSim)
}

sims = simulateMyfit(250, fittedModel)
dharmaRes <- createDHARMa(simulatedResponse = sims,
                           observedResponse = testData$observedResponse,
                           fittedPredictedResponse = predict(fittedModel, type = "response"),
                           integer = TRUE)
plot(dharmaRes, quantreg = FALSE)

---

**DHARMa**

**DHARMa - Residual Diagnostics for HierArchical (Multi-level / Mixed) Regression Models**

**Description**

The 'DHARMa' package uses a simulation-based approach to create readily interpretable scaled (quantile) residuals for fitted (generalized) linear mixed models. Currently supported are linear and generalized linear (mixed) models from 'lme4' (classes 'lmerMod', 'glmerMod'), 'glmmTMB' and 'spaMM', generalized additive models ('gam' from 'mgcv'), 'glm' (including 'negbin' from 'MASS', but excluding quasi-distributions) and 'lm' model classes. Moreover, externally created simulations, e.g. posterior predictive simulations from Bayesian software such as 'JAGS', 'STAN', or 'BUGS' can be processed as well. The resulting residuals are standardized to values between 0 and 1 and can be interpreted as intuitively as residuals from a linear regression. The package also provides a number of plot and test functions for typical model misspecification problems, such as over/underdispersion, zero-inflation, and residual spatial and temporal autocorrelation.

**Details**

See index / vignette for details

**See Also**

simulateResiduals

**Examples**

vignette("DHARMa", package="DHARMa")
getFitted

Get model fitted

Description
Wrapper to get the fitted value of a fitted model

Usage
getFitted(object, ...)

## Default S3 method:
getFitted(object, ...)

## S3 method for class 'gam'
getFitted(object, ...)

Arguments
object a fitted model
... additional parameters to be passed on, usually to the simulate function of the respective model class

Details
The purpose of this wrapper is to standardize the extraction of fitted values

Author(s)
Florian Hartig

See Also
getObservedResponse, getSimulations, getRefit, getFixedEffects

Examples
testData = createData(sampleSize = 400, family = gaussian())

fittedModel <- lm(observedResponse ~ Environment1, data = testData)

# response that was used to fit the model
getObservedResponse(fittedModel)

# predictions of the model for these points
getFitted(fittedModel)

# extract simulations from the model as matrix
getSimulations(fittedModel, nsim = 2)

# extract simulations from the model for refit (often requires different structure)
x = getSimulations(fittedModel, nsim = 2, type = "refit")

getRefit(fittedModel, x[[1]])

getRefit(fittedModel, getObservedResponse(fittedModel))
Description

Extract the response of a fitted model

Usage

getObservedResponse(object, ...)

## Default S3 method:
getObservedResponse(object, ...)

## S3 method for class 'HLfit'
getObservedResponse(object, ...)

Arguments

object a fitted model
...
additional parameters

Details

The purpose of this function is to safely extract the response (dependent variable) of the fitted model classes

Author(s)

Florian Hartig

See Also

getRefit, getSimulations, getFixedEffects, getFitted

Examples

testData = createData(sampleSize = 400, family = gaussian())
fittedModel <- lm(observedResponse ~ Environment1, data = testData)

# response that was used to fit the model
getObservedResponse(fittedModel)

# predictions of the model for these points
getFitted(fittedModel)

# extract simulations from the model as matrix
getSimulations(fittedModel, nsim = 2)

# extract simulations from the model for refit (often requires different structure)
x = getSimulations(fittedModel, nsim = 2, type = "refit")

getRefit(fittedModel, x[[1]])

getRefit(fittedModel, getObservedResponse(fittedModel))

getQuantile

calculate quantiles

Description

calculates residual quantiles from a given simulation

Usage

getQuantile(simulations, observed, integerResponse, method = c("PIT", "traditional"))

Arguments

simulations a matrix with simulations from a fitted model. Rows = observations, columns = replicate simulations
observed a vector with the observed data
integerResponse is the response integer-valued. Only has an effect for method = "traditional"
method the quantile randomization method used. See details

Details

The function calculates residual quantiles from the simulated data. For continuous distributions, this will simply the the value of the ecdf.

For discrete data, there are two options implemented.

The current default (available since DHARMa 0.3.1) are probability integral transform (PIT-) residuals (Smith, 1985; Dunn & Smyth, 1996; see also see also Warton, et al., 2017).

Before DHARMa 0.3.1, a different randomization procedure was used, in which the a U(-0.5, 0.5) distribution was added on observations and simulations for discrete distributions. For a completely discrete distribution, the two procedures should deliver equivalent results, but the second method has the disadvantage that a) one has to know if the distribution is discrete (DHARMa tries to recognize this automatically), and b) that it leads to inefficiencies for some distributions such as the the Tweedie, which are partly continuous, partly discrete (see e.g. https://github.com/florianhartig/DHARMa/issues/168).
**getRandomState**

**References**


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**getRandomState**  
*Record and restore a random state*

**Description**

The aim of this function is to record, manipulate and restore a random state.

**Usage**

getRandomState(seed = NULL)

**Arguments**

- **seed**: seed argument to set.seed(). NULL = no seed, but random state will be restored.  
  F = random state will not be restored

**Details**

This function is intended for two (not mutually exclusive tasks)

a) record the current random state

b) change the current random state in a way that the previous state can be restored

**Value**

a list with various infos about the random state that after function execution, as well as a function to restore the previous state before the function execution

**Author(s)**

Florian Hartig
Examples

# testing the function in standard settings

set.seed(13)
runif(1)
x = getRandomState(123)
runif(1)
x$restoreCurrent()
runif(1)

# values outside set /restore are identical to

set.seed(13)
runif(2)

# if no seed is set, this will also be restored

rm(.Random.seed) # now, there is no random seed
x = getRandomState(123)
exists(".Random.seed") # TRUE
runif(1)
x$restoreCurrent()
exists(".Random.seed") # False

# with seed = false

x = getRandomState(seed = FALSE)
exists(".Random.seed")
runif(1)
x$restoreCurrent()
exists(".Random.seed")

getRefit

Get model refit

Description

Wrapper to refit a fitted model
checks if the fitted model excluded NA values

Usage

getRefit(object, newresp, ...)

## Default S3 method:
getRefit(object, newresp, ...)

## S3 method for class 'lm'
getRefit(object, newresp, ...)

getRefit

Get model refit

Description

Wrapper to refit a fitted model
checks if the fitted model excluded NA values

Usage

getRefit(object, newresp, ...)

## Default S3 method:
getRefit(object, newresp, ...)

## S3 method for class 'lm'
getRefit(object, newresp, ...)

getRefit
getRefit

## S3 method for class 'glmmTMB'
getRefit(object, newresp, ...)

## S3 method for class 'HLfit'
getRefit(object, newresp, ...)

### Arguments

- **object**: a fitted model
- **newresp**: the new response that should be used to refit the model
- **...**: additional parameters to be passed on to the refit or update class that is used to refit the model

### Details

The purpose of this wrapper is to standardize the refit of a model. The behavior of this function depends on the supplied model. When available, it uses the refit method, otherwise it will use update. For glmmTMB: since version 1.0, glmmTMB has a refit function, but this didn’t work, so I switched back to this implementation, which is a hack based on the update function.

Checks if the fitted model excluded NA values

### Author(s)

Florian Hartig

### See Also

- `getObservedResponse`, `getSimulations`, `getFixedEffects`

### Examples

```r
testData = createData(sampleSize = 400, family = gaussian())

fittedModel <- lm(observedResponse ~ Environment1, data = testData)

# response that was used to fit the model
getObservedResponse(fittedModel)

# predictions of the model for these points
getFitted(fittedModel)

# extract simulations from the model as matrix
getSimulations(fittedModel, nsim = 2)

# extract simulations from the model for refit (often requires different structure)
x = getSimulations(fittedModel, nsim = 2, type = "refit")

getRefit(fittedModel, x[[1]])

getRefit(fittedModel, getObservedResponse(fittedModel))
```
getSimulations

Description

Wrapper to simulate from a fitted model

Usage

getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

## Default S3 method:
getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

## S3 method for class 'negbin'
getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

## S3 method for class 'lmerMod'
getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

## S3 method for class 'glmmTMB'
getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

## S3 method for class 'HLfit'
getSimulations(object, nsim = 1, type = c("normal", "refit"), ...)

Arguments

object a fitted model
nsim number of simulations
type if simulations should be prepared for getQuantile or for refit
... additional parameters to be passed on, usually to the simulate function of the respective model class

Details

The purpose of this wrapper for the simulate function is to return the simulations from a model in a standardized way.
The function is a wrapper for the simulate function is to return the simulations from a model in a standardized way.

Note: if the model was fit with weights, the function will throw a warning if used with a model class whose simulate function does not include the weights in the simulations. Note that the results may or may not be appropriate in this case, depending on how you use the weights.

Value

a matrix with simulations
Author(s)
Florian Hartig

See Also
getObservedResponse, getRefit, getFixedEffects, getFitted

Examples

testData = createData(sampleSize = 400, family = gaussian())
fittedModel <- lm(observedResponse ~ Environment1, data = testData)

# response that was used to fit the model
getObservedResponse(fittedModel)

# predictions of the model for these points
getFitted(fittedModel)

# extract simulations from the model as matrix
getSimulations(fittedModel, nsim = 2)

# extract simulations from the model for refit (often requires different structure)
x = getSimulations(fittedModel, nsim = 2, type = "refit")

getRefit(fittedModel, x[[1]])

getRefit(fittedModel, getObservedResponse(fittedModel))

hist.DHARMa

Hist of DHARMa residuals

Description
The function produces a histogram from a DHARMa output

Usage

```r
## S3 method for class 'DHARMa'
hist(x, breaks = seq(-0.02, 1.02, len = 53), col = c("red", rep("lightgrey", 50), "red"), main = "Hist of DHARMa residuals", xlab = "Residuals (outliers are marked red)", cex.main = 1, ...)
```

Arguments

- **x**: a DHARMa simulation output (class DHARMa)
- **breaks**: breaks for hist() function
- **col**: col for hist bars
main plot main
xlab plot xlab
cex.main plot cex.main
...
other arguments to be passed on to hist

See Also
plotSimulatedResiduals, plotResiduals

Examples

testData = createData(sampleSize = 200, family = poisson(),
  randomEffectVariance = 1, numGroups = 5)
fittedModel <- glm( observedResponse ~ Environment1,
  family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

######### main plotting function #############
# for all functions, quantreg = T will be more
# informative, but slower
plot(simulationOutput, quantreg = FALSE)

############# Distribution ####################
plotQQunif(simulationOutput = simulationOutput)
hist(simulationOutput )

############## residual plots ##################
# rank transformation, using a simulationOutput
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE)

# smooth scatter plot - usually used for large datasets, default for n > 10000
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE, smoothScatter = TRUE)

# residual vs predictors, using explicit values for pred, residual
plotResiduals(simulationOutput, form = testData$Environment1,
  quantreg = FALSE)

# if pred is a factor, or asFactor = T, will produce a boxplot
plotResiduals(simulationOutput, form = testData$group,
  quantreg = FALSE, asFactor = TRUE)

# All these options can also be provided to the main plotting function
plot(simulationOutput, quantreg = FALSE, rank = FALSE)

# If you want to plot summaries per group, use
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, asFactor = TRUE) # we see one residual point per RE

Description

Returns the outliers of a DHARMa object

Usage

outliers(object, lowerQuantile = 0, upperQuantile = 1, return = c("index", "logical"))

Arguments

object an object with simulated residuals created by simulateResiduals
lowerQuantile lower threshold for outliers. Default is zero = outside simulation envelope
upperQuantile upper threshold for outliers. Default is 1 = outside simulation envelope
return wheter to return an indices of outliers or a logical vector

Details

First of all, note that the standard definition of outlier in the DHARMa plots and outlier tests is an observation that is outside the simulation envelope. How far outside that is depends a lot on how many simulations you do. If you have 100 data points and to 100 simulations, you would expect to have one "outlier" on average, even with a perfectly fitting model. This is in fact what the outlier test tests.

Thus, keep in mind that for a small number of simulations, outliers are mostly a technical term: these are points that are outside our simulations, but we don’t know how far away they are.

If you are seriously interested in HOW FAR outside the expected distribution a data point is, you should increase the number of simulations in simulateResiduals to be sure to get the tail of the data distribution correctly. In this case, it may make sense to adjust lowerQuantile and upperQuantile, e.g. to 0.025, 0.975, which would define outliers as values outside the central 95%

Also, note that outliers are particularly concerning if they have a strong influence on the model fit. One could test the influence, for example, by removing them from the data, or by some measures of leverage, e.g. generalisations for Cook’s distance as in Pinho, L. G. B., Nobre, J. S., & Singer, J. M. (2015). Cook’s distance for generalized linear mixed models. Computational Statistics & Data Analysis, 82, 126–136. doi:10.1016/j.csda.2014.08.008. At the moment, however, no such function is provided in DHARMa.
plot.DHARMa

DHARMa standard residual plots

Description

This function creates standard plots for the simulated residuals

Usage

```r
## S3 method for class 'DHARMa'
plot(x, rank = TRUE, ...)
```

Arguments

- `x` an object with simulated residuals created by `simulateResiduals`
- `rank` if T (default), the values of pred will be rank transformed. This will usually make patterns easier to spot visually, especially if the distribution of the predictor is skewed.
- `...` further options for `plotResiduals`. Consider in particular parameters quantreg, rank and asFactor. xlab, ylab and main cannot be changed when using `plotSimulatedResiduals`, but can be changed when using `plotResiduals`.

Details

The function creates two plots. To the left, a qq-uniform plot to detect deviations from overall uniformity of the residuals (calling `plotQQunif`), and to the right, a plot of residuals against predicted values (calling `plotResiduals`). Outliers are highlighted in red (for more on outliers, see `testOutliers`). For a correctly specified model, we would expect

a) a straight 1-1 line in the uniform qq-plot -> evidence for an overall uniform (flat) distribution of the residuals

b) uniformity of residuals in the vertical direction in the res against predictor plot

Deviations of this can be interpreted as for a linear regression. See the vignette for detailed examples.

To provide a visual aid in detecting deviations from uniformity in y-direction, the plot of the residuals against the predicted values also performs an (optional) quantile regression, which provides 0.25, 0.5 and 0.75 quantile lines across the plots. These lines should be straight, horizontal, and at y-values of 0.25. 0.5 and 0.75. Note, however, that some deviations from this are to be expected by chance, even for a perfect model, especially if the sample size is small. See further comments on this plot, its interpretation and options, in `plotResiduals`

The quantile regression can take some time to calculate, especially for larger datasets. For that reason, quantreg = F can be set to produce a smooth spline instead. This is default for n > 2000.

See Also

`plotResiduals, plotQQunif`
Examples

testData = createData(sampleSize = 200, family = poisson(),
                       randomEffectVariance = 1, numGroups = 5)
fittedModel <- glm(observableResponse ~ Environment1,
                    family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

######### main plotting function#################
# for all functions, quantreg = T will be more
# informative, but slower
plot(simulationOutput, quantreg = FALSE)

############## Distribution  ########################
plotQQunif(simulationOutput = simulationOutput)
hist(simulationOutput)

############## residual plots  ########################
# rank transformation, using a simulationOutput
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE)

# smooth scatter plot - usually used for large datasets, default for n > 10000
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE, smoothScatter = TRUE)

# residual vs predictors, using explicit values for pred, residual
plotResiduals(simulationOutput, form = testData$Environment1,
              quantreg = FALSE)

# if pred is a factor, or asFactor = T, will produce a boxplot
plotResiduals(simulationOutput, form = testData$group,
              quantreg = FALSE, asFactor = TRUE)

# All these options can also be provided to the main plotting function
plot(simulationOutput, quantreg = FALSE, rank = FALSE)

# If you want to plot summaries per group, use
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, asFactor = TRUE)  # we see one residual point per RE
**plotQQunif**

**Description**

Convenience function to draw conventional residual plots

**Usage**

plotConventionalResiduals(fittedModel)

**Arguments**

- **fittedModel** a fitted model object

---

**plotQQunif**  
Quantile-quantile plot for a uniform distribution

**Description**

The function produces a uniform quantile-quantile plot from a DHARMa output

**Usage**

plotQQunif(simulationOutput, testUniformity = T, testOutliers = T, testDispersion = T, ...)

**Arguments**

- **simulationOutput** a DHARMa simulation output (class DHARMa)
- **testUniformity** if T, the function testUniformity will be called and the result will be added to the plot
- **testOutliers** if T, the function testOutliers will be called and the result will be added to the plot
- **testDispersion** if T, the function testDispersion will be called and the result will be added to the plot
- ... arguments to be passed on to qqunif

**Details**

the function calls qqunif from the R package gap to create a quantile-quantile plot for a uniform distribution.

**See Also**

plotSimulatedResiduals, plotResiduals
Examples

testData = createData(sampleSize = 200, family = poisson(),
                      randomEffectVariance = 1, numGroups = 5)
fitModel <- glm(observableResponse ~ Environment1,
                 family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fitModel)

######### main plotting function #############
# for all functions, quantreg = T will be more
# informative, but slower
plot(simulationOutput, quantreg = FALSE)

############# Distribution ########################
plotQQunif(simulationOutput = simulationOutput)
hist(simulationOutput )

############# residual plots ####################
# rank transformation, using a simulationOutput
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE)

# smooth scatter plot - usually used for large datasets, default for n > 10000
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE, smoothScatter = TRUE)

# residual vs predictors, using explicit values for pred, residual
plotResiduals(simulationOutput, form = testData$Environment1,
              quantreg = FALSE)

# if pred is a factor, or asFactor = T, will produce a boxplot
plotResiduals(simulationOutput, form = testData$group,
              quantreg = FALSE, asFactor = TRUE)

# All these options can also be provided to the main plotting function
plot(simulationOutput, quantreg = FALSE, rank = FALSE)

# If you want to plot summaries per group, use
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, asFactor = TRUE) # we see one residual point per RE

plotResiduals

Generic res ~ pred scatter plot with spline or quantile regression on top
Description

The function creates a generic residual plot with either spline or quantile regression to highlight patterns in the residuals. Outliers are highlighted in red.

Usage

plotResiduals(simulationOutput, form = NULL, quantreg = NULL, rank = F, asFactor = NULL, smoothScatter = NULL, quantiles = c(0.25, 0.5, 0.75), ...)

Arguments

- `simulationOutput`: an object, usually a DHARMa object, from which residual values can be extracted. Alternatively, a vector with residuals or a fitted model can be provided, which will then be transformed into a DHARMa object.
- `form`: optional predictor against which the residuals should be plotted. Default is to use the predicted(simulationOutput).
- `quantreg`: whether to perform a quantile regression on 0.25, 0.5, 0.75 on the residuals. If `F`, a spline will be created instead. Default NULL chooses `T` for nObs < 2000, and `F` otherwise.
- `rank`: if `T`, the values provided in `form` will be rank transformed. This will usually make patterns easier to spot visually, especially if the distribution of the predictor is skewed. If `form` is a factor, this has no effect.
- `asFactor`: should a numeric predictor provided in `form` be treated as a factor. Default is to choose this for < 10 unique values, as long as enough predictions are available to draw a boxplot.
- `smoothScatter`: if `T`, a smooth scatter plot will plotted instead of a normal scatter plot. This makes sense when the number of residuals is very large. Default NULL chooses `T` for nObs < 10000, and `F` otherwise.
- `quantiles`: for a quantile regression, which quantiles should be plotted
- `...`: additional arguments to plot / boxplot.

Details

The function plots residuals against a predictor (by default against the fitted value, extracted from the DHARMa object, or any other predictor).

Outliers are highlighted in red (for information on definition and interpretation of outliers, see `testOutliers`).

To provide a visual aid in detecting deviations from uniformity in y-direction, the plot function calculates an (optional) quantile regression, which compares the empirical 0.25, 0.5 and 0.75 quantiles (default) in y direction (red solid lines) with the theoretical 0.25, 0.5 and 0.75 quantiles (dashed black line).

Asymptotically (i.e. for lots of data / residuals), if the model is correct, theoretical and the empirical quantiles should be identical (i.e. dashed and solid lines should match). A p-value for the deviation is calculated for each quantile line. Significant deviations are highlighted by red color.
If form is a factor, a boxplot will be plotted instead of a scatter plot. The distribution for each factor level should be uniformly distributed, so the box should go from 0.25 to 0.75, with the median line at 0.5. Again, chance deviations from this will increases when the sample size is smaller. You can run null simulations to test if the deviations you see exceed what you would expect from random variation. If you want to create box plots for categorical predictors (e.g. because you only have a small number of unique numeric predictor values), you can convert your predictor with as.factor(pred).

**Value**

if quantile tests are performed, the function returns them invisibly.

**Note**

The quantile regression can take some time to calculate, especially for larger datasets. For that reason, quantreg = F can be set to produce a smooth spline instead.

**See Also**

plotQQunif

**Examples**

testData = createData(sampleSize = 200, family = poisson(),
    randomEffectVariance = 1, numGroups = 5)
fittedModel <- glm(observedResponse ~ Environment1,
    family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# for all functions, quantreg = T will be more informative, but slower
plot(simulationOutput, quantreg = FALSE)

# rank transformation, using a simulationOutput
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE)

# smooth scatter plot - usually used for large datasets, default for n > 10000
plotResiduals(simulationOutput, rank = TRUE, quantreg = FALSE, smoothScatter = TRUE)

# residual vs predictors, using explicit values for pred, residual
plotResiduals(simulationOutput, form = testData$Environment1,
plotSimulatedResiduals

DHARMa standard residual plots

Description

DEPRECATED, use plot() instead

Usage

plotSimulatedResiduals(simulationOutput, ...)

Arguments

- simulationOutput: an object with simulated residuals created by simulateResiduals
- ...: further options for plotResiduals. Consider in particular parameters quantreg, rank and asFactor. xlab, ylab and main cannot be changed when using plotSimulatedResiduals, but can be changed when using plotResiduals.

Note

This function is deprecated. Use plot.DHARMa

See Also

plotResiduals, plotQQunif
**print.DHARMa**

| print.DHARMa | *Print simulated residuals* |

**Description**

Print simulated residuals

**Usage**

```r
## S3 method for class 'DHARMa'
print(x, ...)
```

**Arguments**

- `x`: an object with simulated residuals created by `simulateResiduals`
- `...`: optional arguments for compatibility with the generic function, no function implemented

**recalculateResiduals**

| recalculateResiduals | *Recalculate residuals with grouping* |

**Description**

The purpose of this function is to recalculate scaled residuals per group, based on the simulations done by `simulateResiduals`

**Usage**

```r
recalculateResiduals(simulationOutput, group = NULL, aggregateBy = sum,
                     seed = 123, method = c("PIT", "traditional"))
```

**Arguments**

- `simulationOutput`: an object with simulated residuals created by `simulateResiduals`
- `group`: group of each data point
- `aggregateBy`: function for the aggregation. Default is `sum`. This should only be changed if you know what you are doing. Note in particular that the expected residual distribution might not be flat any more if you choose general functions, such as `sd` etc.
the random seed to be used within DHARMa. The default setting, recommended for most users, is keep the random seed on a fixed value 123. This means that you will always get the same randomization and thus the same result when running the same code. NULL = no new seed is set, but previous random state will be restored after simulation. FALSE = no seed is set, and random state will not be restored. The latter two options are only recommended for simulation experiments. See vignette for details.

the quantile randomization method used. The two options implemented at the moment are probability integral transform (PIT-) residuals (current default), and the "traditional" randomization procedure, that was used in DHARMa until version 0.3.0. For details, see getQuantile

an object of class DHARMa, similar to what is returned by simulateResiduals, but with additional outputs for the new grouped calculations. Note that the relevant outputs are 2x in the object, the first is the grouped calculations (which is returned by $name access), and later another time, under identical name, the original output. Moreover, there is a function `aggregateByGroup`, which can be used to aggregate predictor variables in the same way as the variables calculated here

Examples

library(lme4)

df <- data.frame(x = rnorm(50), y = rnorm(50))
model <- lme4::lmer(y ~ x + (1|group), data = df)
simulationOutput <- simulateResiduals(fittedModel = model)

# standard plot
plot(simulationOutput)

# one of the possible test, for other options see ?testResiduals
testOutliers(simulationOutput)

# for various other plots and tests, see the help / vignette

# the calculated residuals can be accessed via
residuals(simulationOutput)

# transform residuals to other pdf, see ?residuals.DHARMa for details
residuals(simulationOutput, quantileFunction = qnorm, outlierValues = c(-7,7))

# calculating summaries per group
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, quantreg = FALSE)

# create residuals with refitting, see ?simulateResiduals for details
# n=10 is very low, set higher when using this for real
simulationOutput <- simulateResiduals(fittedModel = fittedModel,
residuals.DHARMa

n = 10, refit = TRUE)
plot(simulationOutput, quantreg = FALSE)

residuals.DHARMa

Return residuals of a DHARMa simulation

Description

Return residuals of a DHARMa simulation

Usage

## S3 method for class 'DHARMa'
residuals(object, quantileFunction = NULL, outlierValues = NULL, ...)

Arguments

object

an object with simulated residuals created by simulateResiduals

quantileFunction

optional - a quantile function to transform the uniform 0/1 scaling of DHARMa
to another distribution

outlierValues

if a quantile function with infinite support (such as dnorm) is used, residuals thatare 0/1 are mapped to -Inf / Inf. outlierValues allows to convert -Inf / Inf values
to an optional min / max value.

...

optional arguments for compatibility with the generic function, no function implemented

Details

the function accesses the slot $scaledResiduals in a fitted DHARMa object, and optionally transforms the standard DHARMa quantile residuals (which have a uniform distribution) to a particular pdf.

Note

some of the papers on simulated quantile residuals transforming the residuals (which are natively uniform) back to a normal distribution. I presume this is because of the larger familiarity of most users with normal residuals. Personally, I never considered this desirable, for the reasons explained in https://github.com/florianhartig/DHARMa/issues/39, but with this function, I wanted to give users the option to plot normal residuals if they so wish.
Examples

```r
library(lme4)

testData = createData(sampleSize = 200, overdispersion = 0.5, family = poisson())
fittedModel <- glmer(observedResponse ~ Environment1 + (1|group),
  family = "poisson", data = testData)

simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# standard plot
plot(simulationOutput)

# one of the possible test, for other options see ?testResiduals
testOutliers(simulationOutput)

# for various other plots and tests, see the help / vignette
# the calculated residuals can be accessed via
residuals(simulationOutput)

# transform residuals to other pdf, see ?residuals.DHARMa for details
residuals(simulationOutput, quantileFunction = qnorm, outlierValues = c(-7,7))

# calculating summaries per group
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, quantreg = FALSE)

# create residuals with refitting, see ?simulateResiduals for details
# n=10 is very low, set higher when using this for real
simulationOutput <- simulateResiduals(fittedModel = fittedModel,
  n = 10, refit = TRUE)
plot(simulationOutput, quantreg = FALSE)
```

---

**runBenchmarks**  
**Benchmark calculations**

**Description**

This function runs statistical benchmarks, including Power / Type I error simulations for an arbitrary test with a control parameter

**Usage**

```r
runBenchmarks(calculateStatistics, controlValues = NULL, nRep = 10, alpha = 0.05, parallel = F, ...)
```
simulateResiduals

Arguments

calculateStatistics
the statistics to be benchmarked. Should return one value, or a vector of values. If controlValues are given, must accept a parameter control

controlValues
a vector with a control parameter (e.g. to vary the strength of a problem the test should be specific to)

nRep
number of replicates per level of the controlValues

alpha
significance level

parallel
whether to use parallel computations. Possible values are F, T (sets the cores automatically to number of available cores -1), or an integer number for the number of cores that should be used for the cluster

... additional parameters to calculateStatistics

Note

The benchmark function in DHARMa are intended for development purposes, and for users that want to test / confirm the properties of functions in DHARMa. If you are running an applied data analysis, they are probably of little use.

simulateResiduals

Create simulated residuals

Description

The function creates scaled residuals by simulating from the fitted model. Residuals can be extracted with residuals.DHARMa. See testResiduals for an overview of residual tests, plot.DHARMa for an overview of available plots.

Usage

simulateResiduals(fittedModel, n = 250, refit = F, integerResponse = NULL, plot = F, seed = 123, method = c("PIT", "traditional"),...)

Arguments

fittedModel
a fitted model of a class supported by DHARMa

n
number of simulations. Default is 100. A more save value would be 250 or even 1000. The smaller the number, the higher the stochastic error on the residuals. Also, for very small n, discretization artefacts can influence the tests.

refit
if FALSE, new data will be simulated and scaled residuals will be created by comparing observed data with new data. If TRUE, the model will be refit on the simulated data (parametric bootstrap), and scaled residuals will be created by comparing observed with refitted residuals.
integerResponse

if TRUE, noise will be added at to the residuals to maintain a uniform expectations for integer responses (such as Poisson or Binomial). Usually, the model will automatically detect the appropriate setting, so there is no need to adjust this setting.

plot

if TRUE, plotResiduals will be directly run after the residuals have been calculated.

seed

the random seed to be used within DHARMa. The default setting, recommended for most users, is keep the random seed on a fixed value 123. This means that you will always get the same randomization and thus teh same result when running the same code. NULL = no new seed is set, but previous random state will be restored after simulation. FALSE = no seed is set, and random state will not be restored. The latter two options are only recommended for simulation experiments. See vignette for details.

method

the quantile randomization method used. The two options implemented at the moment are probability integral transform (PIT-) residuals (current default), and the "traditional" randomization procedure, that was used in DHARMa until version 0.3.0. For details, see getQuantile

... parameters to pass to the simulate function of the model object. An important use of this is to specify whether simulations should be conditional on the current random effect estimates, e.g. via re.form. Note that not all models support syntax to specify conditionao or unconditional simulations. See also details

Details

There are a number of important considerations when simulating from a more complex (hierarchical) model:

**Re-simulating random effects / hierarchical structure:** in a hierarchical model, we have several stochastic processes aligned on top of each other. Specifically, in a GLMM, we have a lower level stochastic process (random effect), whose result enters into a higher level (e.g. Poisson distribution). For other hierarchical models such as state-space models, similar considerations apply.

In such a situation, we have to decide if we want to re-simulate all stochastic levels, or only a subset of those. For example, in a GLMM, it is common to only simulate the last stochastic level (e.g. Poisson) conditional on the fitted random effects. This is often referred to as a conditional simulation. For controlling how many levels should be re-simulated, the simulateResidual function allows to pass on parameters to the simulate function of the fitted model object. Please refer to the help of the different simulate functions (e.g. ?simulate.merMod) for details. For merMod (lme4) model objects, the relevant parameters are parameters are use.u and re.form

If the model is correctly specified, the simulated residuals should be flat regardless how many hierarchical levels we re-simulate. The most thorough procedure would therefore be to test all possible options. If testing only one option, I would recommend to re-simulate all levels, because this essentially tests the model structure as a whole. This is the default setting in the DHARMa package. A potential drawback is that re-simulating the lower-level random effects creates more variability, which may reduce power for detecting problems in the upper-level stochastic processes. In particular dispersion tests may produce different results when switching from conditional to unconditional simulations, and often the conditional simulation is more sensitive.
**Integer responses**: a second complication is the treatment of inter responses. Imaging we have observed a 0, and we predict 30% zeros - what is the quantile that we should display for the residual? To deal with this problem and maintain a uniform response, the option integerResponse adds a uniform noise from -0.5 to 0.5 on the simulated and observed response, which creates a uniform distribution - you can see this via hist(ecdf(runif(10000))(runif(10000))).

DHARMa will try to automatically if the fitted model has an integer or discrete distribution via the family argument. However, in some cases the family does not allow to uniquely identify the distribution type. For example, a tweedie distribution can be inter or continuous. Therefore, DHARMa will additionally check the simulation results for repeated values, and will change the distribution type if repeated values are found (a message is displayed in this case).

**Refitting or not**: a third issue is how residuals are calculated. simulateResiduals has two options that are controlled by the refit parameter:

1. if refit = FALSE (default), new data is simulated from the fitted model, and residuals are calculated by comparing the observed data to the new data
2. if refit = TRUE, a parametric bootstrap is performed, meaning that the model is refit on the new data, and residuals are created by comparing observed residuals against refitted residuals. I advise against using this method per default (see more comments in the vignette), unless you are really sure that you need it.

**Residuals per group**: In many situations, it can be useful to look at residuals per group, e.g. to see how much the model over/underpredicts per plot, year or subject. To do this, use recalculateResiduals, together with a grouping variable (see also help)

**Transformation to other distributions**: DHARMa calculates residuals for which the theoretical expectation (assuming a correctly specified model) is uniform. To transfer this residuals to another distribution (e.g. so that a correctly specified model will have normal residuals) see residuals.DHARMa.

**Value**

An S3 class of type "DHARMa", essentially a list with various elements. Implemented S3 functions include plot, print and residuals.DHARMa. Residuals returns the calculated scaled residuals.

**See Also**

testResiduals, plot.DHARMa, plotResiduals, print.DHARMa, residuals.DHARMa, recalculateResiduals

**Examples**

```r
library(lme4)

testData = createData(sampleSize = 200, overdispersion = 0.5, family = poisson())
fittedModel <- glmer( observedResponse ~ Environment1 + (1|group),
                      family = "poisson", data = testData)

simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# standard plot
plot(simulationOutput)
```
# one of the possible test, for other options see ?testResiduals
testOutliers(simulationOutput)

# for various other plots and tests, see the help / vignette
# the calculated residuals can be accessed via
residuals(simulationOutput)

# transform residuals to other pdf, see ?residuals.DHARMa for details
residuals(simulationOutput, quantileFunction = qnorm, outlierValues = c(-7,7))

# calculating summaries per group
simulationOutput = recalculateResiduals(simulationOutput, group = testData$group)
plot(simulationOutput, quantreg = FALSE)

# create residuals with refitting, see ?simulateResiduals for details
# n=10 is very low, set higher when using this for real
simulationOutput <- simulateResiduals(fittedModel = fittedModel, n = 10, refit = TRUE)
plot(simulationOutput, quantreg = FALSE)

---

**testDispersion**  
*DHARMa dispersion tests*

**Description**

This function performs a simulation-based test for over/underdispersion

**Usage**

```r
testDispersion(simulationOutput, alternative = c("two.sided", "greater", "less"), plot = T, ...)
```

**Arguments**

- `simulationOutput`  
an object of class DHARMa with simulated quantile residuals, either created via `simulateResiduals` or by `createDHARMa` for simulations created outside DHARMa

- `alternative`  
a character string specifying whether the test should test if observations are "greater", "less" or "two.sided" compared to the simulated null hypothesis. Greater corresponds to overdispersion.

- `plot`  
whether to plot output

- `...`  
arguments to pass on to `testGeneric`
Details

The function implements two tests, depending on whether it is applied on a simulation with refit = F, or refit = T.

If refit = F, the function tests the sd of the data against the sd of the simulated data.

If refit = T, the function compares the approximate deviance (via squared pearson residuals) with the same quantity from the models refitted with simulated data. Applying this is much slower than the previous alternative. Given the computational cost, I would suggest that most users will be satisfied with the standard dispersion test.

Note

The results of the dispersion test can differ depending on whether it is evaluated on conditional (= conditional on fitted random effects) or unconditional (= REs are re-simulated) simulations. You can change between conditional or unconditional simulations in `simulateResiduals` if this is supported by the regression package that you use. The default in DHARMa is to use unconditional simulations, but I have often found that conditional simulations are more sensitive to dispersion problems. I recommend trying both, as neither test should be positive if the dispersion is correct.

Author(s)

Florian Hartig

See Also

testResiduals, testUniformity, testOutliers, testZeroInflation, testGeneric, testTemporalAutocorrelation, testSpatialAutocorrelation, testQuantiles

Examples

testData = createData(sampleSize = 200, overdispersion = 0.5, randomEffectVariance = 0)
fittedModel <- glm(observedResponse ~ Environment1, family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# the plot function runs 4 tests
# i) KS test ii) Dispersion test iii) Outlier test iv) quantile test
plot(simulationOutput, quantreg = TRUE)

# testResiduals tests distribution, dispersion and outliers
testResiduals(simulationOutput)

###### Individual tests #######

# KS test for correct distribution of residuals
testUniformity(simulationOutput)

# Dispersion test
testDispersion(simulationOutput) # tests under and overdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion
# if model is refitted, a different test will be called
simulationOutput2 <- simulateResiduals(fittedModel = fittedModel, refit = TRUE, seed = 12)
testDispersion(simulationOutput2)

# often useful to test dispersion per group (e.g. binomial data, see vignette)
simulationOutput3 = recalculateResiduals(simulationOutput, group = testData$group)
testDispersion(simulationOutput3)

# Outlier test (number of observations outside simulation envelope)
testOutliers(simulationOutput)

# testing zero inflation
testZeroInflation(simulationOutput)

# testing generic summaries
countOnes <- function(x) sum(x == 1) # testing for number of 1s
testGeneric(simulationOutput, summary = countOnes) # 1-inflation
testGeneric(simulationOutput, summary = countOnes, alternative = "less") # 1-deficit

means <- function(x) mean(x) # testing if mean prediction fits
testGeneric(simulationOutput, summary = means)

spread <- function(x) sd(x) # testing if mean sd fits
testGeneric(simulationOutput, summary = spread)

---

**testGeneric**

*Generic simulation test of a summary statistic*

**Description**

This function tests if a user-defined summary differs when applied to simulated / observed data.

**Usage**

```r
testGeneric(simulationOutput, summary, alternative = c("two.sided", "greater", "less"), plot = T, methodName = "DHARMa generic simulation test")
```

**Arguments**

- `simulationOutput`
  - an object of class DHARMa with simulated quantile residuals, either created via `simulateResiduals` or by `createDHARMa` for simulations created outside DHARMa

- `summary`
  - a function that can be applied to simulated / observed data. See examples below
alternative  a character string specifying whether the test should test if observations are
"greater", "less" or "two.sided" compared to the simulated null hypothesis
plot         whether to plot the simulated summary
methodName   name of the test (will be used in plot)

Details
This function tests if a user-defined summary differs when applied to simulated / observed data.
the function can easily be remodeled to apply summaries on the residuals, by simply defining f =
function(x) summary(x - predictions), as done in testDispersion

Note
The function that you supply is applied on the data as it is represented in your fitted model, which
may not always correspond to how you think. This is important in particular when you use k/n
binomial data, and want to test for 1-inflation. As an example, if have k/20 observations, and you
provide your data via cbind (y, y-20), you have to test for 20-inflation (because this is how the data
is represented in the model). However, if you provide data via y/20, and weights = 20, you should
test for 1-inflation. In doubt, check how the data is internally represented in model.frame(model),
or via simulate(model)

Author(s)
Florian Hartig

See Also
testResiduals, testUniformity, testOutliers, testDispersion, testZeroInflation, testTemporalAutocorrelation,
testSpatialAutocorrelation, testQuantiles

Examples
testData = createData(sampleSize = 200, overdispersion = 0.5, randomEffectVariance = 0)
fittedModel <- glm(observedResponse ~ Environment1, family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# the plot function runs 4 tests
# i) KS test ii) Dispersion test iii) Outlier test iv) quantile test
plot(simulationOutput, quantreg = TRUE)

# testResiduals tests distribution, dispersion and outliers
testResiduals(simulationOutput)

######### Individual tests #########

# KS test for correct distribution of residuals
testUniformity(simulationOutput)

# Dispersion test
testDispersion(simulationOutput) # tests under and overdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion

# if model is refitted, a different test will be called
simulationOutput2 <- simulateResiduals(fittedModel = fittedModel, refit = TRUE, seed = 12)
testDispersion(simulationOutput2)

# often useful to test dispersion per group (e.g. binomial data, see vignette)
simulationOutput3 = recalculateResiduals(simulationOutput, group = testData$group)
testDispersion(simulationOutput3)

# Outlier test (number of observations outside simulation envelope)
testOutliers(simulationOutput)

# testing zero inflation
testZeroInflation(simulationOutput)

# testing generic summaries
countOnes <- function(x) sum(x == 1) # testing for number of 1s
testGeneric(simulationOutput, summary = countOnes) # 1-inflation
testGeneric(simulationOutput, summary = countOnes, alternative = "less") # 1-deficit

means <- function(x) mean(x) # testing if mean prediction fits
testGeneric(simulationOutput, summary = means)

spread <- function(x) sd(x) # testing if mean sd fits
testGeneric(simulationOutput, summary = spread)

---

testOutliers

Test for outliers

Description

This function tests if the number of observations outside the simulation envelope are larger or smaller than expected

Usage

testOutliers(simulationOutput, alternative = c("two.sided", "greater", "less"), margin = c("both", "upper", "lower"), plot = T)

Arguments

simulationOutput

an object of class DHARMa with simulated quantile residuals, either created via simulateResiduals or by createDHARMa for simulations created outside DHARMa
alternative a character string specifying whether the test should test if observations are "greater", "less" or "two.sided" (default) compared to the simulated null hypothesis

margin whether to test for outliers only at the lower, only at the upper, or both sides (default) of the simulated data distribution

plot if T, the function will create an additional plot

Details

DHARMa residuals are created by simulating from the fitted model, and comparing the simulated values to the observed data. It can occur that all simulated values are higher or smaller than the observed data, in which case they get the residual value of 0 and 1, respectively. I refer to these values as simulation outliers, or simply outliers.

Because no data was simulated in the range of the observed value, we don’t know "how strongly" these values deviate from the model expectation, so the term "outlier" should be used with a grain of salt - it’s not a judgment about the magnitude of a deviation from an expectation, but simply that we are outside the simulated range, and thus cannot say anything more about the location of the residual.

Note also that the number of outliers will decrease as we increase the number of simulations. Under the null hypothesis that the model is correct, we expect nData / (nSim +1) outliers at each margin of the distribution. For a reason, consider that if the data and the model distribution are identical, the probability that a given observation is higher than all simulations is 1/(nSim +1).

Based on this null expectation, we can test for an excess or lack of outliers. Per default, testOutliers() looks for both, so if you get a significant p-value, you have to check if you have too many or too few outliers. An excess of outliers is to be interpreted as too many values outside the simulation envelope. This could be caused by overdispersion, or by what we classically call outliers. A lack of outliers would be caused, for example, by underdispersion.

Author(s)

Florian Hartig

See Also
testResiduals, testUniformity, testDispersion, testZeroInflation, testGeneric, testTemporalAutocorrelation, testSpatialAutocorrelation, testQuantiles

Examples

set.seed(123)

testData = createData(sampleSize = 200, overdispersion = 1, randomEffectVariance = 0)
fittedModel <- glm(observedResponse ~ Environment1 , family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# default outlier test (with plot)
testOutliers(simulationOutput)

# note that default is to test outliers at both margins for both an excess and a lack
# of outliers. Here we see that we mostly have an excess of outliers at the upper
# margin. You see that it is an exces because the frequency of outliers is 0.055,
# while expected is 0.008

# Let's see what would have happened if we would just have checked the lower margin
testOutliers(simulationOutput, margin = "lower", plot = FALSE)

# OK, now the frequency of outliers is 0, so we have too few, but this is n.s. against
# the expectation

# just for completeness, what would have happened if we would have checked both
# margins, but just for a lack of outliers (i.e. underdispersion)
testOutliers(simulationOutput, alternative = "less", plot = FALSE)

testOverdispersion  Simulated overdisperstion tests

Description

Simulated overdispersion tests

Usage

testOverdispersion(simulationOutput, ...)

Arguments

simulationOutput

an object of class DHARMa with simulated quantile residuals, either created
via simulateResiduals or by createDHARMa for simulations created outside
DHARMa

...  additional arguments to testDispersion

Details

Deprecated, switch your code to using the testDispersion function
testOverdispersionParametric

Parametric overdispersion tests

Description

Parametric overdispersion tests

Usage

testOverdispersionParametric(...)

Arguments

... arguments will be ignored, the parametric tests is no longer recommend

Details

Deprecated, switch your code to using the testDispersion function. The function will do nothing, arguments will be ignored, the parametric tests is no longer recommend

testPDistribution

Plot distribution of p-values

Description

Plot distribution of p-values

Usage

testPDistribution(x, plot = T,
main = "p distribution \n expected is flat at 1", ...)  

Arguments

x vector of p values
plot should the values be plottet
main title for the plot
... additional arguments to hist

Author(s)

Florian Hartig
testQuantiles  

Test for quantiles

Description

This function tests

Usage

testQuantiles(simulationOutput, predictor = NULL, quantiles = c(0.25, 0.5, 0.75), plot = T)

Arguments

- simulationOutput: an object of class DHARMa with simulated quantile residuals, either created via simulateResiduals or by createDHARMa for simulations created outside DHARMa
- predictor: an optional predictor variable to be used, instead of the predicted response (default)
- quantiles: the quantiles to be tested
- plot: if T, the function will create an additional plot

Details

The function fits quantile regressions (via package qgam) on the residuals, and compares their location to the expected location (because of the uniform distribution, the expected location is 0.5 for the 0.5 quantile).

A significant p-value for the splines means the fitted spline deviates from a flat line at the expected location (p-values of intercept and spline are combined via Benjamini & Hochberg adjustment to control the FDR).

The p-values of the splines are combined into a total p-value via Benjamini & Hochberg adjustment to control the FDR.

Author(s)

Florian Hartig

See Also

testResiduals, testUniformity, testDispersion, testZeroInflation, testGeneric, testTemporalAutocorrelation, testSpatialAutocorrelation, testOutliers
Examples

testData = createData(sampleSize = 200, overdispersion = 0.0, randomEffectVariance = 0)
fittedModel <- glm(1~Environment1, family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# run the quantile test
x = testQuantiles(simulationOutput)
x # the test shows a combined p-value, corrected for multiple testing
x$pvals # p-values for the individual quantiles
x$qgamFits # access the fitted quantile regression
summary(x$qgamFits[[1]]) # summary of the first fitted quantile

# possible to test user-defined quantiles
x = testQuantiles(simulationOutput, quantiles = c(0.7))

# example with missing environmental predictor
fittedModel <- glm(1~Environment1, family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)
testQuantiles(simulationOutput, predictor = testData$Environment1)

# the quantile test is automatically performed in
## Not run:
plot(simulationOutput)
plotResiduals(simulationOutput)
## End(Not run)

---

testResiduals  DHARMa general residual test

Description

Calls both uniformity and dispersion test

Usage

testResiduals(simulationOutput, plot = T)

Arguments

simulationOutput

an object of class DHARMa with simulated quantile residuals, either created
via simulateResiduals or by createDHARMa for simulations created outside
DHARMa

plot

if T, plots functions of the tests are called
Details

This function is a wrapper for the various test functions implemented in DHARMa. Currently, this function calls the `testUniformity` and the `testDispersion` functions. All other tests (see list below) have to be called by hand.

Author(s)

Florian Hartig

See Also

`testUniformity`, `testOutliers`, `testDispersion`, `testZeroInflation`, `testGeneric`, `testTemporalAutocorrelation`, `testSpatialAutocorrelation`, `testQuantiles`

Examples

def testData = createData(sampleSize = 200, overdispersion = 0.5, randomEffectVariance = 0)
def fittedModel <- glm(observedResponse ~ Environment1, family = "poisson", data = testData)
def simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# the plot function runs 4 tests
# i) KS test i) Dispersion test iii) Outlier test iv) quantile test
plot(simulationOutput, quantreg = TRUE)

def testResiduals tests distribution, dispersion and outliers
def testResiduals(simulationOutput)

######## Individual tests ########

# KS test for correct distribution of residuals
def testUniformity(simulationOutput)

# Dispersion test
def testDispersion(simulationOutput) # tests under and overdispersion
def testDispersion(simulationOutput, alternative = "less") # only underdispersion
def testDispersion(simulationOutput, alternative = "less") # only underdispersion

# if model is refitted, a different test will be called
def simulationOutput2 <- simulateResiduals(fittedModel = fittedModel, refit = TRUE, seed = 12)
def testDispersion(simulationOutput2)

# often useful to test dispersion per group (e.g. binomial data, see vignette)
def simulationOutput3 = recalculateResiduals(simulationOutput, group = testData$group)
def testDispersion(simulationOutput3)

# Outlier test (number of observations outside simulation envelope)
def testOutliers(simulationOutput)

# testing zero inflation
def testZeroInflation(simulationOutput)

# testing generic summaries
`testSimulatedResiduals`  

```r
countOnes <- function(x) sum(x == 1) # testing for number of 1s
testGeneric(simulationOutput, summary = countOnes) # 1-inflation
testGeneric(simulationOutput, summary = countOnes, alternative = "less") # 1-deficit

means <- function(x) mean(x) # testing if mean prediction fits
testGeneric(simulationOutput, summary = means)

spread <- function(x) sd(x) # testing if mean sd fits
testGeneric(simulationOutput, summary = spread)
```

---

**testSimulatedResiduals**  
*Residual tests*

**Description**

Residual tests

**Usage**

```r
testSimulatedResiduals(simulationOutput)
```

**Arguments**

- `simulationOutput`
  - an object of class DHARMa with simulated quantile residuals, either created via `simulateResiduals` or by `createDHARMa` for simulations created outside DHARMa

**Details**

Deprecated, switch your code to using the `testResiduals` function

**Author(s)**

Florian Hartig
testSpatialAutocorrelation

Test for spatial autocorrelation

Description

This function performs a standard test for spatial autocorrelation on the simulated residuals

Usage

testSpatialAutocorrelation(simulationOutput, x = NULL, y = NULL,
distMat = NULL, alternative = c("two.sided", "greater", "less"),
plot = T)

Arguments

- simulationOutput: an object of class DHARMa with simulated quantile residuals, either created via simulateResiduals or by createDHARMa for simulations created outside DHARMa
- x: the x coordinate, in the same order as the data points. If not provided, random values will be created
- y: the y coordinate, in the same order as the data points. If not provided, random values will be created
- distMat: optional distance matrix. If not provided, a distance matrix will be calculated based on x and y. See details for explanation
- alternative: a character string specifying whether the test should test if observations are "greater", "less" or "two.sided" compared to the simulated null hypothesis
- plot: whether to plot output

Details

The function performs Moran.I test from the package ape, based on the provided distance matrix of the data points.

There are several ways to specify this distance. If a distance matrix (distMat) is provided, calculations will be based on this distance matrix, and x,y coordinates will only used for the plotting (if provided) If distMat is not provided, the function will calculate the euclidian distances between x,y coordinates, and test Moran.I based on these distances.

If no x/y values are provided, random values will be created. The sense of being able to run the test with x/y = NULL (random values) is to test the rate of false positives under the current residual structure (random x/y corresponds to H0: no spatial autocorrelation), e.g. to check if the test has nominal error rates for particular residual structures.

Testing for spatial autocorrelation requires unique x,y values - if you have several observations per location, either use the recalculateResiduals function to aggregate residuals per location, or extract the residuals from the fitted object, and plot / test each of them independently for spatially repeated
subgroups (a typical scenario would repeated spatial observation, in which case one could plot/test each time step separately for temporal autocorrelation). Note that the latter must be done by hand, outside testSpatialAutocorrelation.

**Note**

Important to note for all autocorrelation tests (spatial/temporal): the autocorrelation tests are valid to check for residual autocorrelation in models that don’t assume such a correlation (in this case, you can use conditional or unconditional simulations), or if there is remaining residual autocorrelation after accounting for it in a spatial/temporal model (in that case, you have to use conditional simulations), but if checking unconditional simulations from a model with an autocorrelation structure on data that corresponds to this model, they will be significant, even if the model fully accounts for this structure.

This behavior is not really a bug, but rather originates from the definition of the quantile residuals: quantile residuals are calculated independently per data point, i.e. without consideration of any correlation structure between data points that may exist in the simulations. As a result, the simulated distributions from a unconditional simulation will typically not reflect the correlation structure that is present in each single simulation, and the same is true for the subsequently calculated quantile residuals.

The bottomline here is that spatial/temporal/other autoregressive models should either be tested based on conditional simulations, or (ideally) custom tests should be used that are not based on quantile residuals, but rather compare the correlation structure in the simulated data with the correlation structure in the observed data.

**Author(s)**

Florian Hartig

**See Also**

testResiduals, testUniformity, testOutliers, testDispersion, testZeroInflation, testGeneric, testTemporalAutocorrelation, testQuantiles

**Examples**

testData = createData(sampleSize = 40, family = gaussian())
fittedModel <- lm(observedResponse ~ Environment1, data = testData)
res = simulateResiduals(fittedModel)

# Standard use
testSpatialAutocorrelation(res, x = testData$x, y = testData$y)

# If x and y is not provided, random values will be created
testSpatialAutocorrelation(res)

# Alternatively, one can provide a distance matrix
dM = as.matrix(dist(cbind(testData$x, testData$y)))
testSpatialAutocorrelation(res, distMat = dM)

# if there are multiple observations with the same x values,
# create first ar group with unique values for each location
# then aggregate the residuals per location, and calculate
# spatial autocorrelation on the new group

res2 = recalculateResiduals(res, group = testData$group)
testSpatialAutocorrelation(res)

# careful when using REs to account for spatially clustered (but not grouped)
# data. This originates from https://github.com/florianhartig/DHARMa/issues/81

# Assume our data is divided into clusters, where observations are close together
# but not at the same point, and we suspect that observations in clusters are
# autocorrelated

clusters = 100
subsamples = 10
size = clusters * subsamples

testData = createData(sampleSize = size, family = gaussian(), numGroups = clusters)
testData$x = rnorm(clusters)[testData$group] + rnorm(size, sd = 0.01)
testData$y = rnorm(clusters)[testData$group] + rnorm(size, sd = 0.01)

# It's a good idea to use a RE to take out the cluster effects. This accounts
# for the autocorrelation within clusters

library(lme4)
fittedModel <- lmer(observedResponse ~ Environment1 + (1|group), data = testData)

# DHARMa default is to re-simulated REs - this means spatial pattern remains
# because residuals are still clustered

res = simulateResiduals(fittedModel)
testSpatialAutocorrelation(res, x = testData$x, y = testData$y)

# However, it should disappear if you just calculate an aggregate residuals per cluster
# Because at least how the data are simulated, cluster are spatially independent

res2 = recalculateResiduals(res, group = testData$group)
testSpatialAutocorrelation(res2,
  x = aggregate(testData$x, list(testData$group), mean)$x,
  y = aggregate(testData$y, list(testData$group), mean)$x)

# For lme4, it's also possible to simulated residuals conditional on fitted
# REs (re.form). Conditional on the fitted REs (i.e. accounting for the clusters)
# the residuals should now be independent. The remaining RSA we see here is
# probably due to the RE shrinkage

res = simulateResiduals(fittedModel, re.form = NULL)
testSpatialAutocorrelation(res, x = testData$x, y = testData$y)
testTemporalAutocorrelation

Test for temporal autocorrelation

Description

This function performs a standard test for temporal autocorrelation on the simulated residuals

Usage

testTemporalAutocorrelation(simulationOutput, time = NULL, alternative = c("two.sided", "greater", "less"), plot = T)

Arguments

simulationOutput
an object with simulated residuals created by simulateResiduals
time
the time, in the same order as the data points. If not provided, random values will be created
alternative
a character string specifying whether the test should test if observations are "greater", "less" or "two.sided" compared to the simulated null hypothesis
plot
whether to plot output

Details

The function performs a Durbin-Watson test on the uniformly scaled residuals, and plots the residuals against time. The DB test was originally be designed for normal residuals. In simulations, I didn’t see a problem with this setting though. The alternative is to transform the uniform residuals to normal residuals and perform the DB test on those.

If no time values are provided, random values will be created. The sense of being able to run the test with time = NULL (random values) is to test the rate of false positives under the current residual structure (random time corresponds to H0: no spatial autocorrelation), e.g. to check if the test has nominal error rates for particular residual structures (note that Durbin-Watson originally assumes normal residuals, error rates seem correct for uniform residuals, but may not be correct if there are still other residual problems).

Testing for temporal autocorrelation requires unique time values - if you have several observations per time value, either use the recalculateResiduals function to aggregate residuals per time step, or extract the residuals from the fitted object, and plot / test each of them independently for temporally repeated subgroups (typical choices would be location / subject etc.). Note that the latter must be done by hand, outside testSpatialAutocorrelation.

Note

Important to note for all autocorrelation tests (spatial / temporal): the autocorrelation tests are valid to check for residual autocorrelation in models that don’t assume such a correlation (in this case,
you can use conditional or unconditional simulations), or if there is remaining residual autocorrelation after accounting for it in a spatial/temporal model (in that case, you have to use conditional simulations), but if checking unconditional simulations from a model with an autocorrelation structure on data that corresponds to this model, they will be significant, even if the model fully accounts for this structure.

This behavior is not really a bug, but rather originates from the definition of the quantile residuals: quantile residuals are calculated independently per data point, i.e. without consideration of any correlation structure between data points that may exist in the simulations. As a result, the simulated distributions from an unconditional simulation will typically not reflect the correlation structure that is present in each single simulation, and the same is true for the subsequently calculated quantile residuals.

The bottomline here is that spatial / temporal / other autoregressive models should either be tested based on conditional simulations, or (ideally) custom tests should be used that are not based on quantile residuals, but rather compare the correlation structure in the simulated data with the correlation structure in the observed data.

Author(s)
Florian Hartig

See Also
testResiduals, testUniformity, testOutliers, testDispersion, testZeroInflation, testGeneric, testSpatialAutocorrelation, testQuantiles

Examples
testData = createData(sampleSize = 40, family = gaussian())
fittedModel <- lm(observedResponse ~ Environment1, data = testData)
res = simulateResiduals(fittedModel)

# Standard use
testTemporalAutocorrelation(res, time = testData$time)

# If no time is provided, random values will be created
testTemporalAutocorrelation(res)

# If you have several observations per time step
timeSeries1 = createData(sampleSize = 40, family = gaussian())
timeSeries1$location = 1
timeSeries2 = createData(sampleSize = 40, family = gaussian())
timeSeries2$location = 2
testData = rbind(timeSeries1, timeSeries2)
fittedModel <- lm(observedResponse ~ Environment1, data = testData)
res = simulateResiduals(fittedModel)

# for this, you cannot do testTemporalAutocorrelation(res, time = testData$time)
# because here we would have observations with the same time, i.e.
# zero difference in time. We have two options a) aggregate observations
# b) calculate / test per subset. Testing per subset might also be useful
# if you have several locations, regardless of whether the times are
# identical, because you would expect the autocorrelation structure to be
# independent per location

# testing grouped residuals

res = recalculateResiduals(res, group = testData$time)
testTemporalAutocorrelation(res, time = unique(testData$time))

# plotting and testing per subgroup

# extract subgroup
testData$Residuals = res$scaledResiduals
temp = testData[testData$location == 1,]

# plots and tests
plot(Residuals ~ time, data = temp)
lmtest::dwtest(temp$Residuals ~ 1, order.by = temp$time)

---

testUniformity  Test for overall uniformity

**Description**

This function tests the overall uniformity of the simulated residuals in a DHARMa object

**Usage**

testUniformity(simulationOutput, alternative = c("two.sided", "less", "greater"), plot = T)

**Arguments**

- **simulationOutput**
  - an object of class DHARMa with simulated quantile residuals, either created via simulateResiduals or by createDHARMa for simulations created outside DHARMa

- **alternative**
  - a character string specifying whether the test should test if observations are "greater", "less" or "two.sided" compared to the simulated null hypothesis. See ks.test for details

- **plot**
  - if T, plots calls plotQQunif as well

**Details**

The function applies a ks.test for uniformity on the simulated residuals.
Author(s)
Florian Hartig

See Also

testResiduals, testOutliers, testDispersion, testZeroInflation, testGeneric, testTemporalAutocorrelation, testSpatialAutocorrelation, testQuantiles

Examples

testData = createData(sampleSize = 200, overdispersion = 0.5, randomEffectVariance = 0)
fittedModel <- glm(phenomenon ~ Environment1 + Environment2, family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# the plot function runs 4 tests
# i) KS test ii) Dispersion test iii) Outlier test iv) quantile test
plot(simulationOutput, quantreg = TRUE)

testResiduals tests distribution, dispersion and outliers
testResiduals(simulationOutput)

######## Individual tests ########

# KS test for correct distribution of residuals
testUniformity(simulationOutput)

# Dispersion test
testDispersion(simulationOutput) # tests under and overdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion

# if model is refitted, a different test will be called
simulationOutput2 <- simulateResiduals(fittedModel = fittedModel, refit = TRUE, seed = 12)
testDispersion(simulationOutput2)

# often useful to test dispersion per group (e.g. binomial data, see vignette)
simulationOutput3 = recalculateResiduals(simulationOutput, group = testData$group)
testDispersion(simulationOutput3)

# Outlier test (number of observations outside simulation envelope)
testOutliers(simulationOutput)

# testing zero inflation
testZeroInflation(simulationOutput)

# testing generic summaries
countOnes <- function(x) sum(x == 1) # testing for number of 1s
testGeneric(simulationOutput, summary = countOnes) # 1-inflation
testGeneric(simulationOutput, summary = countOnes, alternative = "less") # 1-deficit

means <- function(x) mean(x) # testing if mean prediction fits
testGeneric(simulationOutput, summary = means)
spread <- function(x) sd(x) # testing if mean sd fits
testGeneric(simulationOutput, summary = spread)

---

testZeroInflation | Tests for zero-inflation

Description
This function compares the observed number of zeros with the zeros expected from simulations.

Usage
testZeroInflation(simulationOutput, ...)

Arguments
- simulationOutput:
  an object of class DHARMa with simulated quantile residuals, either created via `simulateResiduals` or by `createDHARMa` for simulations created outside DHARMa.
- ...
  further arguments to `testGeneric`.

Details
The plot shows the expected distribution of zeros against the observed values, the ratioObsSim shows observed vs. simulated zeros. A value < 1 means that the observed data has less zeros than expected, a value > 1 means that it has more zeros than expected (aka zero-inflation). Per default, the function tests both sides.

Some notes about common problems / questions:
* Zero-inflation tests after fitting the model are crucial to see if you have zero-inflation. Just because there are a lot of zeros doesn’t mean you have zero-inflation, see Warton, D. I. (2005). Many zeros does not mean zero inflation: comparing the goodness-of-fit of parametric models to multivariate abundance data. Environmetrics 16(3), 275-289.
* That being said, zero-inflation tests are often not a reliable guide to decide whether to add a zi term or not. In general, model structures should be decided on ideally a priori, if that is not possible via model selection techniques (AIC, BIC, WAIC, Bayes Factor). A zero-inflation test should only be run after that decision, and to validate the decision that was taken.

Note
This function is a wrapper for `testGeneric`, where the summary argument is set to function(x) sum(x == 0).
Author(s)

Florian Hartig

See Also

testResiduals, testUniformity, testOutliers, testDispersion, testGeneric, testTemporalAutocorrelation, testSpatialAutocorrelation, testQuantiles

Examples

testData = createData(sampleSize = 200, overdispersion = 0.5, randomEffectVariance = 0)
fittedModel <- glm( observedResponse ~ Environment1 , family = "poisson", data = testData)
simulationOutput <- simulateResiduals(fittedModel = fittedModel)

# the plot function runs 4 tests
# i) KS test i) Dispersion test iii) Outlier test iv) quantile test
plot(simulationOutput, quantreg = TRUE)

# testResiduals tests distribution, dispersion and outliers
interpretResiduals(simulationOutput)

######### Individual tests ########

# KS test for correct distribution of residuals
testUniformity(simulationOutput)

# Dispersion test
testDispersion(simulationOutput) # tests under and overdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion
testDispersion(simulationOutput, alternative = "less") # only underdispersion

# if model is refitted, a different test will be called
simulationOutput2 <- simulateResiduals(fittedModel = fittedModel, refit = TRUE, seed = 12)
testDispersion(simulationOutput2)

# often useful to test dispersion per group (e.g. binomial data, see vignette)
simulationOutput3 = recalculateResiduals(simulationOutput, group = testData$group)
testDispersion(simulationOutput3)

# Outlier test (number of observations outside simulation envelope)
testOutliers(simulationOutput)

# testing zero inflation
testZeroInflation(simulationOutput)

# testing generic summaries
countOnes <- function(x) sum(x == 1) # testing for number of 1s
testGeneric(simulationOutput, summary = countOnes) # 1-inflation
testGeneric(simulationOutput, summary = countOnes, alternative = "less") # 1-deficit

means <- function(x) mean(x) # testing if mean prediction fits
testGeneric(simulationOutput, summary = means)
spread <- function(x) sd(x) # testing if mean sd fits
testGeneric(simulationOutput, summary = spread)

transformQuantiles

Transform quantiles to pdf (deprecated)

Description

The purpose of this function was to transform the DHARMa quantile residuals (which have a uniform distribution) to a particular pdf. Since DHARMa 0.3.0, this functionality is integrated in the residuals.DHARMa function. Please switch to using this function.

Usage

transformQuantiles(res, quantileFunction = qnorm, outlierValue = 7)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>res</td>
<td>an object with simulated residuals created by simulateResiduals</td>
</tr>
<tr>
<td>quantileFunction</td>
<td>optional - a quantile function to transform the uniform 0/1 scaling of DHARMa to another distribution</td>
</tr>
<tr>
<td>outlierValue</td>
<td>if a quantile function with infinite support (such as dnorm) is used, residuals that are 0/1 are mapped to -Inf / Inf. outlierValues allows to convert -Inf / Inf values to an optional min / max value.</td>
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