# Package: gosset (via r-universe)

December 5, 2024

```
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```

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# Description

Akaike weights represent the relative likelihood of a model. It can be used in model averaging and selection.

# Usage

akaike\_weights(object)

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## **Arguments**

object a numerical vector with models goodness of fit coefficients

#### Value

A data frame containing the coefficients:

```
delta the delta overall change in the coefficients relative_logLik the relative log-likelihood akaike_weights the Akaike weights
```

#### Author(s)

Kauê de Sousa and Jacob van Etten

#### References

Wagenmakers E. J. & Farrell S. (2004). Psychonomic Bulletin and Review, 11(1), 192–196. doi:10.3758/BF03206482

```
data("airquality")
# try three model approaches
mod1 = glm(Temp \sim 1,
            data = airquality,
            family = poisson())
mod2 = glm(Temp \sim Ozone,
            data = airquality,
            family = poisson())
mod3 = glm(Temp ~ Ozone + Solar.R,
            data = airquality,
            family = poisson())
# models AICs together in a single vector
models = c(mod1 = AIC(mod1),
            mod2 = AIC(mod2),
            mod3 = AIC(mod3)
# calculate akaike weights
aw = akaike_weights(models)
# the higher the better
names(models[which.max(aw$akaike_weights)])
```

4 anova.PlackettLuce

anova.PlackettLuce

Likelihood-ratio test

# Description

Assesses the goodness of fit of competing statistical models

# Usage

```
## $3 method for class 'PlackettLuce'
anova(object, ...)
likelihood_ratio(x, split, ...)
```

# Arguments

object an object of class PlackettLuce
... additional arguments passed to methods
x an object of class rankings or grouped\_rankings
split a vector indicating the splitting rule for the test

## Author(s)

Joost van Heerwaarden and Kauê de Sousa

```
library("PlackettLuce")
example("beans", package = "PlackettLuce")
G = group(R, rep(seq_len(nrow(beans)), 4))
d = cbind(G, beans)

split = ifelse(d$maxTN < 18.7175, TRUE, FALSE)

likelihood_ratio(G, split)

mod = PlackettLuce(G)
anova(mod)</pre>
```

breadwheat 5

breadwheat

Preferred bread wheat varieties

#### **Description**

Data from decentralized on-farm trials of bread wheat (*Triticum aestivum* L.) varieties in Vaishali, India over the 2014's Rabi season. Farmers were asked to test three varieties of bread wheat randomly assigned as incomplete blocks of three varieties (out of 16 varieties) and assess which one had the best and worst performance considering four traits, germination, grain quality, yield and overall performance.

## Usage

breadwheat

#### **Format**

```
A data frame with 493 records and 19 variables:
variety_a The name of variety A in the comparison.
variety_b The name of variety B in the comparison.
variety_c The name of variety C in the comparison.
district The administrative region where the experiment was stablished.
village The village within the administrative region where the experiment was stablished.
participant_name The participant name (ommitted for protection and privacy).
age The participant age.
gender The participant gender M = Male; F = Female.
planting_date The date which the experiment started.
1 on The longitude in which the experiment was stablished.
lat The latitude in which the experiment was stablished.
germination_best The variety ranked as best for germination ("A", "B" or "C").
germination_worst The variety ranked as worst for germination ("A", "B" or "C").
grainquality_best The variety ranked as best for grain quality ("A", "B" or "C").
grainquality_worst The variety ranked as worst for grain quality ("A", "B" or "C").
yield_best The variety ranked as best for yield ("A", "B" or "C").
yield_worst The variety ranked as worst for yield ("A", "B" or "C").
overall_best The variety ranked as best for overall perfomance ("A", "B" or "C").
overall_worst The variety ranked as worst for overall perfomance ("A", "B" or "C").
```

## Source

```
van Etten, J., et. al. (2016). Experimental Agriculture, 55, 275-296. doi:10.1017/S0014479716000739 van Etten, J., et. al. (2019). PNAS 116(10) 4194-4199 doi:10.1073/pnas.1813720116
```

6 btpermute

btpermute

Variable selection with Permuted Inclusion Criterion

#### Description

Method of forward variable selection based on deviance for Bradley-Terry models using pairwise ranking data. The selection procedure consists of two steps, first, permuting the variables from the original predictors with n.iteractions, then performing a forward selection to retain the predictors with highest contribution to the model, see details.

## Usage

```
btpermute(
  contests = NULL,
  predictors = NULL,
  n.iterations = 15,
  seed = NULL,
  ...
)
```

# **Arguments**

contests a data frame with pairwise binary contests with these variables 'id', 'player1', 'player2', 'win1', 'win2';

in that order. The id should be equivalent to the index of each row in predictors

predictors a data frame with player-specific variables with row indices that should match

with the ids in contests. An id is not required, only the predictor variables, the

ids are the index for each row

n.iterations integer, number of iterations to compute

seed integer, the seed for random number generation. If NULL (the default), gosset

will set the seed randomly

... additional arguments passed to **BradleyTerry2** methods

#### **Details**

The selection procedure consists of two steps. In the first step, btpermute adds to the set of original (candidate) predictors variables an additional set of 'fake', permuted variables. This set of permuted predictors is created by assigning to each ranking the variables from another, randomly selected ranking. The permuted variables are not expected to have any predictive power for pairwise rankings. In the second step, btpermute adds predictors to the Bradley-Terry model in a forward selection procedure. Each predictors (real and permuted) is added to the null model individually, and btpermute retains which variable reduces model deviance most strongly. The two-step process is replicated n times with argument n.iterations. At each iteration, a new random permutation is generated and all variables are tested. Replicability can be controlled using argument seed. Across the n.iterations, the function identifies the predictor that appeared most often as the most deviance-reducing one. When this is a real variable, it is constantly added to the model and the forward selection procedure moves on – again creating new permutations, adding

cassava 7

real and fake variables individually, and examining model deviance. Variable selection stops when a permuted variable is found to be most frequently the most deviance-reducing predictors across n.iterations. In turn, variable selection continuous as long as any real variable has stronger explanatory power for pairwise rankings than the random variables.

#### Value

an object of class gosset\_btpermute with the final BTm() model, selected variables, seeds (random numbers) used for permutations and deviances

## Author(s)

Jonathan Steinke and Kauê de Sousa

#### References

Lysen, S. (2009) Permuted inclusion criterion: A variable selection technique. University of Pennsylvania

#### See Also

```
set_binomialfreq, BTm
Other model selection functions: crossvalidation()
```

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## **Description**

Data from decentralized consumer trials of gari-eba, a sub-product of cassava (*Manihot esculenta* Crantz). Consumer testing was carried out in 2022 in Cameroon and Nigeria using the tricot approach. Diverse consumers in villages, towns and cities evaluated the overall acceptability of garieba made from 13 cassava genotypes. Apart from overall preference of the samples, the following traits were evaluated for eba based on the triangulated insights obtained by earlier survey and participatory work in the three areas. Nigeria (Osun and Benue States): colour, smoothness, mouldability, stretchability, and taste. Cameroon (Littoral zone): colour, odour, taste, firmness and stretchability. Traits in common are: colour, taste and stretchability.

#### **Usage**

cassava

#### **Format**

```
A data frame with 1000 records and 27 variables
id The tricot package id.
option_a The name of genotype A in the comparison.
option_b The name of genotype B in the comparison.
option_c The name of genotype C in the comparison.
country The country where the experiment was conducted.
gender The participant gender.
age The participant age.
consumption Indicates how often the participant consumes eba.
consumptionform Indicates in which meal the participant consumes eba.
colour_pos The sample ranked as best for colour ("A", "B" or "C").
colour_neg The sample ranked as worst for colour ("A", "B" or "C").
smoothness_pos The sample ranked as best for smoothness ("A", "B" or "C").
smoothness_neg The sample ranked as worst for smoothness ("A", "B" or "C").
mouldability_pos The sample ranked as best for mouldability ("A", "B" or "C").
mouldability_neg The sample ranked as worst for mouldability ("A", "B" or "C").
stretchability_pos The sample ranked as best for stretchability ("A", "B" or "C").
stretchability_neg The sample ranked as worst for stretchability ("A", "B" or "C").
taste_pos The sample ranked as best for taste ("A", "B" or "C").
taste_neg The sample ranked as worst for taste ("A", "B" or "C").
odour_pos The sample ranked as best for odour ("A", "B" or "C").
odour_neg The sample ranked as worst for odour ("A", "B" or "C").
firmness_pos The sample ranked as best for firmness ("A", "B" or "C").
firmness_neg The sample ranked as worst for firmness ("A", "B" or "C").
overall_pos The sample ranked as best overall ("A", "B" or "C").
```

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```
overall_neg The sample ranked as worst overall ("A", "B" or "C").

reason_like Open question for the reason why the participant ranked the best sample as "best".

reason_dislike Open question for the reason why the participant ranked the worst sample as "worst".
```

#### **Source**

Olaosebikan, et. al. (2023). Journal of the Science of Food and Agriculture. doi:10.1002/jsfa.12867

compare

Compare agreement between two methods

# **Description**

Measures the agreement between two methods

## Usage

```
compare(x, y, ...)
## Default S3 method:
compare(x, y, labels = NULL, ...)
## S3 method for class 'PlackettLuce'
compare(x, y, ...)
```

## **Arguments**

```
    a numeric vector, or an object of class PlackettLuce
    a numeric vector, or an object of class PlackettLuce
    additional arguments passed to methods
    optional, a vector with the same length x to plot values
```

#### Value

a ggplot with the agreement

#### References

```
Bland, M. J., and Altman, D. G. (1986). Lancet (8476):307-10.
```

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#### **Examples**

```
set.seed(1)
x = runif(10, -1, 2)
set.seed(2)
y = runif(10, -1, 2)
compare(x, y)
```

crossvalidation

Cross-validation

## **Description**

Methods for measuring the performance of a predictive model on sets of test data in Bradley-Terry model from **psychotree**, Generalized Linear and Generalized Nonlinear models from **gnm**, and Plackett-Luce model from **PlackettLuce** 

## Usage

```
crossvalidation(formula, data, k = 10, folds = NULL, seed = NULL, ...)
## S3 method for class 'bttree'
deviance(object, newdata = NULL, ...)
## S3 method for class 'bttree'
deviance(object, newdata = NULL, ...)
## S3 method for class 'pltree'
deviance(object, newdata = NULL, ...)
## S3 method for class 'gnm'
AIC(object, newdata = NULL, ...)
## S3 method for class 'gnm'
deviance(object, newdata = NULL, ...)
```

#### **Arguments**

formula	an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted, of the form $y$ $x1 + + xn$
data	a data frame (or object coercible by as.data.frame to a data frame) containing the variables in the model
k	an integer for the number of bins in the cross-validation
folds	an optional vector or list of vectors specifying the $k$ -folds in the cross-validation

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seed integer, the seed for random number generation. If NULL (the default), gosset

will set the seed randomly

... additional arguments passed the methods of the chosen model

object a model object

newdata a data.frame with test data

#### Value

an object of class gosset\_cv with the cross-validation goodness-of-fit estimates, which are:

AIC Akaike Information Criterion

deviance Model deviance logLik Log-Likelihood

MaxLik Maximum likelihood pseudo R-squared CraggUhler Cragg and Uhler's pseudo R-squared

McFadden McFadden pseudo R-squared kendallTau the Kendall correlation coefficient

#### Author(s)

Kauê de Sousa, Jacob van Etten and David Brown

#### References

```
Elder J. F. (2003). Journal of Computational and Graphical Statistics, 12(4), 853–864. doi:10.1198/1061860032733
```

```
James G., et al. (2013). doi:10.1007/9781461471387
```

Whitlock M. C. (2005). Journal of Evolutionary Biology, 18(5), 1368–1373. doi:10.1111/j.1420-9101.2005.00917.x

#### See Also

```
bttree, gnm, pltree
```

Other model selection functions: btpermute()

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```
# Plackett-Luce Model
if(require("PlackettLuce")) {
# beans data from PlackettLuce
data("beans", package = "PlackettLuce")
G = rank_tricot(data = beans,
                 items = c(1:3),
                 input = c(4:5),
                 additional.rank = beans[c(6:8)],
                 group = TRUE)
beans = cbind(G, beans)
# take seasons as bins
k = length(unique(beans$season))
folds = as.integer(as.factor(beans$season))
cv = crossvalidation(G ~ maxTN,
                      data = beans,
                      k = k,
                      folds = folds,
                      minsize = 100)
}
```

gosset

Tools for Data Analysis in Experimental Agriculture

## **Description**

Methods to analyse experimental agriculture data, from data synthesis to model selection and visualisation. The package is named after W.S. Gosset aka 'Student', a pioneer of modern statistics in small sample experimental design and analysis.

# Author(s)

Kauê de Sousa and Jacob van Etten and David Brown and Jonathan Steinke

#### See Also

#### **Useful links:**

gosset paper: doi:10.2139/ssrn.4236267

• Development repository: https://github.com/AgrDataSci/gosset

• Static documentation: https://AgrDataSci.github.io/gosset/

• Report bugs: https://github.com/AgrDataSci/gosset/issues

kendallTau 13

kendallTau

Kendall rank correlation coefficient

#### **Description**

Compute Kendall rank correlation coefficient between two objects. Kendall is a coefficient used in statistics to measure the ordinal association between two measured quantities. A tau test is a non-parametric hypothesis test for statistical dependence based on the tau coefficient. The 'kendallTau' function applies the "kendall" method from 'stats::cor' with some previous treatment in the data, such as converting floating numbers into ranks (from the higher being the first and negative being the last) and the possibility to remove zeros from incomplete ranks

Perform a pairwise permutation test to assess statistical differences in Kendall's Tau correlation between two or more groups.

## Usage

```
kendallTau(x, y, null.rm = TRUE, average = TRUE, na.omit = FALSE, ...)
## Default S3 method:
kendallTau(x, y, null.rm = TRUE, ...)
## S3 method for class 'matrix'
kendallTau(x, y, null.rm = TRUE, average = TRUE, na.omit = FALSE, ...)
## S3 method for class 'rankings'
kendallTau(x, y, ...)
## S3 method for class 'grouped_rankings'
kendallTau(x, y, ...)
## S3 method for class 'paircomp'
kendallTau(x, y, ...)
kendallTau_bootstrap(x, y, nboot = 100, seed = NULL, ...)
kendallTau_permute(x, y, split, n.permutations = 500)
```

#### **Arguments**

X	a numeric vector, matrix or data frame
У	a vector, matrix or data frame with compatible dimensions to x
null.rm	logical, to remove zeros from x and y
average	logical, if FALSE returns the kendall and N-effective for each entry
na.omit	logical, if TRUE ignores entries with kendall = NA when computing the average
	further arguments affecting the Kendall tau produced. See details

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nboot integer, the size of the bootstrap sample

seed integer, the seed for random number generation. If NULL (the default), gosset

will set the seed randomly

split a vector indicating the splitting rule for the test n.permutations an integer, the number of permutations to perform

#### Value

The Kendall correlation coefficient and the Effective N, which is the equivalent N needed if all items were compared to all items. Used for significance testing.

A data.frame containing:

observed\_diff observed absolute differences in Kendall's tau for all group pairs.

p\_values p-values from the permutation test for all group pairs.

#### Author(s)

Kauê de Sousa and Jacob van Etten

Kauê de Sousa

#### References

```
Kendall M. G. (1938). Biometrika, 30(1-2), 81-93. doi:10.1093/biomet/30.12.81
```

#### See Also

cor

kendallTau

Other goodness-of-fit functions: kendallW(), pseudoR2()

```
# Vector based example same as stats::cor(x, y, method = "kendall")
# but showing N-effective
x = c(1, 2, 3, 4, 5)
y = c(1, 1, 3, 2, NA)
w = c(1, 1, 3, 2, 5)
kendallTau(x, y)
kendallTau(x, w)
# Matrix and PlacketLuce ranking example
library("PlackettLuce")
R = matrix(c(1, 2, 4, 3,
```

kendallTau 15

```
1, 4, 2, 3,
             1, 2, NA, 3,
             1, 2, 4, 3,
             1, 3, 4, 2,
             1, 4, 3, 2), nrow = 6, byrow = TRUE)
colnames(R) = LETTERS[1:4]
G = group(as.rankings(R), 1:6)
mod = pltree(G ~ 1, data = G)
preds = predict(mod)
kendallTau(R, preds)
# Also returns raw values (no average)
kendallTau(R, preds, average = FALSE)
# Choose to ignore entries with NA
R2 = matrix(c(1, 2, 4, 3,
              1, 4, 2, 3,
              NA, NA, NA, NA,
              1, 2, 4, 3,
              1, 3, 4, 2,
              1, 4, 3, 2), nrow = 6, byrow = TRUE)
kendallTau(R, R2, average = FALSE)
kendallTau(R, R2, average = TRUE)
kendallTau(R, R2, average = TRUE, na.omit = TRUE)
set.seed(42)
x = rnorm(100)
y = rnorm(100)
split = rep(c("Group1", "Group2", "Group3"), length.out = 100)
kendallTau_permute(x, y, split)
data("breadwheat", package = "gosset")
x = rank_tricot(breadwheat,
                 items = paste0("variety_", letters[1:3]),
                 input = c("yield_best", "yield_worst"),
                 validate.rankings = TRUE)
y = rank_tricot(breadwheat,
                items = paste0("variety_", letters[1:3]),
input = c("overall_best", "overall_worst"),
                 validate.rankings = TRUE)
kendallTau_permute(x, y,
```

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```
split = rep(c("Group1", "Group2", "Group3"), length.out = nrow(breadwheat)),
    n.permutations = 100)
```

kendallW

Kendall's W (coefficient of concordance)

# Description

Compute Kendall's W, also known as coefficient of concordance among observed rankings and those predicted by the Plackett-Luce model.

## Usage

```
kendallW(x, ...)
## Default S3 method:
kendallW(x, y, ...)
## S3 method for class 'pltree'
kendallW(x, newdata = NULL, ...)
```

#### **Arguments**

x an object of class pltree

... additional arguments passed to methods

y an object of class pltree

newdata data for predictions

## **Details**

It is as wrapper of the function DescTools::KendallW, adapted to compute the Kendall's W on the observed and predicted rankings.

Kendall's W values range between 0 (no agreement) to 1 (full agreement)

#### Value

Kendall's W (coefficient of concordance)

## Author(s)

David Brown and Jacob van Etten

#### See Also

Other goodness-of-fit functions: kendallTau(), pseudoR2()

kenyachoice 17

kenyachoice

Kenyan farmers' preferences for agricultural and livelihood practices

## Description

Data from a preference experiment in Makueni County, Kenya. Twenty-six smallholder farmers ordered 9 different livelihood improvement practices by their personal preference. Full rankings were broken down into multiple pairwise rankings (kenyachoice[["contests"]]). To each respondent, a set of nine socio-economic variables is available (kenyachoice[["predictors"]]).

#### Usage

kenyachoice

#### **Format**

A list with two dataframes. kenyachoice[["contests"]] contains the pairwise rankings from farmers choices. kenyachoice[["predictors"]] contains the socioeconomic data for each farmer. Codes for contests are described:

- B Opening a business
- D Dry planting
- G Collective crop marketing
- J Finding off-farm job
- M Machine tillage
- 0 Renting out traction animals
- R Mulching
- T Terracing
- Z Zai pits

#### Source

Steinke, J., et. al. (2019). Computers and Electronics in Agriculture, 162, 991–1000. doi:10.1016/j.compag.2019.05.026

18 nicabean

nicabean

Common bean on-farm trial in Nicaragua

### **Description**

Data from decentralized on-farm trials of common bean (*Phaseolus vulgaris* L.) varieties in Nicaragua over five seasons between 2015 and 2016. Following the tricot approach, farmers were asked to test three varieties of common bean randomly assigned as incomplete blocks of three varieties (out of 10 varieties) and assess which of those three had the best and worst performance in nine traits (Vigor, Architecture, Resistance to Pests, Resistance to Diseases, Tolerance to Drought, Yield, Marketability, Taste, and Overall Appreciation).

## Usage

nicabean

#### **Format**

```
A list with two data frames, nicabean[["trial"]] contains the trial data:
id the plot id
item the variety name
trait the trait for the given variety and plot id
rank the rank for the given variety and trait, with 1 being higher and 3 the lowest
nicabean[["bean_covar"]] contains the covariates associated with the data:
id the plot id
adm0 the country name where trials were set
longitude the longitude of the trial plot
latitude the latitude of the trial plot
trial the trial name as registered on ClimMob
variety_a the variety assigned as label A in the incomplete block
variety_b the variety assigned as label B in the incomplete block
variety_c the variety assigned as label C in the incomplete block
planting_date the planting date
gender the farmer gender
age the farmer age
```

# Source

```
van Etten, J., et. al. (2016). Experimental Agriculture, 55, 275-296. doi:10.1017/S0014479716000739 van Etten, J., et. al. (2019). PNAS 116(10) 4194-4199 doi:10.1073/pnas.1813720116
```

pairwise\_probs 19

pairwise\_probs

Compute pairwise probabilities

### **Description**

Implements the Luce's Choice Axiom to calculate pairwise probabilities in a set of choice probabilities. The Luce's Choice Axiom states that the probability of selecting one item over another from a pool of many items is not affected by the presence or absence of other items in the pool.

#### Usage

```
pairwise_probs(object, relative.probs = TRUE, ...)
```

## **Arguments**

```
object a named numeric vector with probabilities
relative.probs logical, TRUE to return matrix with relative probs (prob - 0.5), otherwise true values are returned
... additional arguments passed to methods
```

#### Value

a matrix with pairwise probabilities

```
library("PlackettLuce")
library("ggplot2")
R = matrix(c(1, 2, 3, 0,
             4, 1, 2, 3,
             2, 1, 3, 4,
             1, 2, 3, 0,
             2, 1, 3, 0,
             1, 0, 3, 2), nrow = 6, byrow = TRUE)
colnames(R) = c("apple", "banana", "grape", "pear")
mod = PlackettLuce(R)
coefs = coefficients(mod, log = FALSE)
pair_worth = pairwise_probs(coefs)
pair_worth
# plot the results
lvls = dimnames(pair_worth)[[1]]
pair_dat = data.frame(player1 = rep(lvls, times = length(lvls)),
```

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```
player2 = rep(lvls, each = length(lvls)),
                       worth = as.vector(pair_worth))
pair_dat
pair_dat$player1 = factor(pair_dat$player1, levels = lvls)
pair_dat$player2 = factor(pair_dat$player2, levels = rev(lvls))
pair_dat$worth = round(pair_dat$worth, 2)
ggplot(pair_dat,
       aes(x = player2,
           y = player1,
           fill = worth,
           label = worth)) +
  geom_tile(show.legend = FALSE) +
  geom_text() +
  scale_fill_gradient2(low = "#b2182b",
                        high = "#2166ac",
                        na.value = "white") +
  scale_x_discrete(position = "top") +
  theme_bw() +
  \label{theme} theme(axis.text = element\_text(color = "grey10"),
        strip.text.x = element_text(color = "grey10"),
        axis.text.x = element_text(angle = 90, hjust = 0),
        panel.grid = element_blank()) +
  labs(x = "",
y = "",
       fill = "")
```

plot.pltree

Get node labels and rules used in a party tree

# Description

Returns the covariates used to split a recursive partitioning tree and the rules that were applied to build the tree

# Usage

```
## S3 method for class 'pltree'
plot(x, log = TRUE, ref = NULL, ...)
## S3 method for class 'PlackettLuce'
plot(x, ...)
node_labels(x)
```

plot.pltree 21

```
node_rules(x)
top_items(x, top = 5)
```

#### **Arguments**

X	an object of class party or PlackettLuce
log	logical, if TRUE log-worth coefficients are displayed instead of worth
ref	optional, character or integer for the reference item when $log = TRUE$
	additional arguments passed to methods. See details
top	an integer for the number of items to return

#### **Details**

multcomp = TRUE adds multi-comparison letters from multcompView ci.level = numeric for the confidence interval levels

#### Value

a vector with the node labels, a data.frame with node rules, a ggplot object

#### Author(s)

Kauê de Sousa

22 pseudoR2

pseudoR2

Pseudo R-squared

## **Description**

Regression coefficient to evaluate goodness-of-fit in a given model when ordinary least squares (OLS) are not available. The algorithm computes estimates from the maximum likelihood through an iterative process. These estimates are called 'pseudo R-squared' because they look like 'R-squared' in the sense that they are on a similar scale (from 0 to 1), with higher values indicating better model fit.

#### Usage

```
pseudoR2(object, ...)
## Default S3 method:
pseudoR2(object, ...)
## S3 method for class 'pltree'
pseudoR2(object, newdata = NULL, ...)
## S3 method for class 'bttree'
pseudoR2(object, ...)
```

pseudoR2 23

# Arguments

object a model object of class glm, gnm, lm, pltree or bttree
... additional arguments affecting the R-squared produced

newdata a data.set with testing data

#### Value

A data frame containing the pseudo R-squared coefficients:

logLik log-likelihood

logLikNull Null log-likelihood

MaxLik maximum likelihood pseudo R-squared CraggUhler Cragg and Uhler's pseudo R-squared

McFadden pseudo R-squared

#### Author(s)

Kauê de Sousa and Jacob van Etten

#### References

Agresti A. (2002). Categorical Data Analysis. John Wiley & Sons, Inc., Hoboken, NJ, USA. doi:10.1002/0471249688

Hunter D. R. (2004). The Annals of Statistics, 32(1), 384-406. http://www.jstor.org/stable/3448514

Cragg, J. G., & Uhler, R. S. (1970). The Canadian Journal of Economics 3(3), 386-406. doi:10.2307/133656

McFadden, D. (1973). Conditional logit analysis of qualitative choice behavior.

#### See Also

```
Other goodness-of-fit functions: kendallTau(), kendallW()
```

24 rank\_numeric

pseudo\_rank

Add pseudo-rank to missing values

# **Description**

Add pseudo-rank to missing values

## Usage

```
pseudo_rank(object, ...)
```

# Arguments

```
object a matrix or PlackettLuce rank
... additional arguments passed to methods
```

#### Value

a matrix or PlackettLuce rank

# **Examples**

rank\_numeric

Plackett-Luce rankings from numeric values

## Description

Group and coerce numeric values into Plackett-Luce rankings.

25 rank\_numeric

#### Usage

```
rank_numeric(
  data,
  items,
  input,
  id = NULL,
  group = FALSE,
  ascending = FALSE,
)
```

# **Arguments**

data a data.frame with columns specified by items and input values a character or numerical vector for indexing the column(s) containing the item items names in data a character or numerical vector for indexing the column(s) containing the values input in data to be ranked id an index of data indicating the ids for "long" data logical, if TRUE return an object of class "grouped\_rankings" group ascending logical, only for floating point numbers, to compute rankings from lower to

higher values

additional arguments passed to methods

## Value

a PlackettLuce "rankings" object, which is a matrix of dense rankings

# Author(s)

Kauê de Sousa

## See Also

#### rankings

```
Other rank functions: rank_tricot(), set_binomialfreq(), set_paircomp()
```

```
# A matrix with 10 rankings of 5 items (A, B, C, D, E)
# with numeric values as "rank"
set.seed(123)
df = cbind(id = rep(1:10, each = 5),
            items = rep(LETTERS[1:5], times = 10),
            input = runif(50, 1, 3))
# return an object of class 'rankings'
R = rank_numeric(df,
```

26 rank\_tricot

rank\_tricot

Build Plackett-Luce rankings from tricot dataset

# Description

Create an object of class "rankings" from tricot data. Tricot stands for "triadic comparison of technology options". Is an approach to carry out large decentralized agronomic field experiments as incomplete blocks. Each incomplete block contains a set of three randomised technologies out of a larger set.

#### Usage

```
rank_tricot(
  data,
  items,
  input,
  group = FALSE,
  validate.rankings = FALSE,
  additional.rank = NULL,
  ...
)
```

## **Arguments**

data a data.frame with columns specified by items and input values

items a character or numerical vector for indexing the column(s) containing the item

names in data

27 rank\_tricot

a character or numerical vector for indexing the column(s) containing the values input in data to be ranked logical, if TRUE return an object of class "grouped\_rankings" group validate.rankings logical, if TRUE implements a check on ranking consistency looking for possible ties, NA or letters other than A, B, C. These entries are set to 0 additional.rank optional, a data frame for the comparisons between tricot items and the local

additional arguments passed to methods. See details

#### **Details**

full.output: logical, to return a list with a "rankings", a "grouped\_rankings" and the ordered items

#### Value

a PlackettLuce "rankings" object, which is a matrix of dense rankings

#### Author(s)

Kauê de Sousa and Jacob van Etten, with ideas from Heather Turner

#### References

```
van Etten J., et al. (2016). Experimental Agriculture, 55(S1), 275–296. doi:10.1017/S0014479716000739
```

#### See Also

```
rankings, breadwheat
Other rank functions: rank_numeric(), set_binomialfreq(), set_paircomp()
```

```
library("PlackettLuce")
data("beans", package = "PlackettLuce")
# Using a subset of the bean data
beans = beans[1:5, 1:5]
beans[1, 1] = NA
beans[3, 4:5] = NA
beans[5, 5] = "Tie"
# The default approach do not validate rankings
# accepting any entry used in the argument input
R1 = rank_tricot(beans,
                 items = c(1:3),
                 input = c(4:5),
                 group = FALSE)
```

28 regret

```
# Using validate.rankings = TRUE, the rankings
# are only considered for those entries without
# NAs, Ties and with any of the letters A, B, C
# this do not affect the lenght of the final ranking
R2 = rank_tricot(beans,
               items = c(1:3),
               input = c(4:5),
               validate.rankings = TRUE,
               group = FALSE)
coef(PlackettLuce(R1))
coef(PlackettLuce(R2))
# pass the comparison with local item as an additional rankings, then
# each of the 3 varieties are compared separately with the local item
# and return an object of class grouped_rankings
data("beans", package = "PlackettLuce")
G = rank_tricot(data = beans,
               items = c(1:3),
               input = c(4:5),
               group = TRUE,
                additional.rank = beans[c(6:8)])
head(G)
```

regret

Regret-based values for risk assessment

## **Description**

Regret is an important heuristic in the behavioural sciences. Minimizing worst regret (the loss under the worst possible outcome) is a criterion that takes a conservative approach to risk analysis in diversification strategies.

#### Usage

```
regret(object, ..., bootstrap = TRUE, normalize = TRUE)
## Default S3 method:
regret(object, ..., values, items, group, bootstrap = TRUE, normalize = TRUE)
## S3 method for class 'pltree'
regret(object, bootstrap = TRUE, normalize = TRUE, ...)
```

regret 29

```
## S3 method for class 'list'
regret(object, bootstrap = TRUE, normalize = TRUE, ...)
```

#### **Arguments**

object a data.frame, an object of class pltree, or a list with PlackettLuce models

... further arguments passed to methods

bootstrap logical, to run a Bayesian bootstrap on object

normalize logical, to normalize values to sum to 1

values an index in *object* with the values to compute regret

items an index in *object* for the different items group an index in *object* for the different scenarios

#### **Details**

Additional details for Bayesian bootstrap: statistic A function that accepts data as its first argument and possibly, the weights as its second, if use\_weights is TRUE; n1 The size of the bootstrap sample; n2 The sample size used to calculate the statistic each bootstrap draw

#### Value

A data frame with regret estimates

items the item names
worth the worth parameters
regret the squared regret
worst\_regret the worst regret

## Author(s)

Jacob van Etten and Kauê de Sousa

#### References

```
Loomes G. & Sugden R. (1982). The Economic Journal, 92(368), 805. doi:10.2307/2232669
Bleichrodt H. & Wakker P. P. (2015). The Economic Journal, 125(583), 493–532. doi:10.1111/ecoj.12200
```

```
# Case 1 ####
library("PlackettLuce")
data("breadwheat", package = "gosset")
# convert the tricot rankings from breadwheat data
# into a object of class 'grouped_rankings'
```

30 reliability

```
G = rank_tricot(breadwheat,
                 items = c("variety_a", "variety_b", "variety_c"),
                 input = c("overall_best", "overall_worst"),
                 group = TRUE)
# combine grouped rankings with temperature indices
mydata = cbind(G, breadwheat[c("lon","lat")])
# fit a pltree model using geographic data
mod = pltree(G ~ ., data = mydata)
regret(mod)
# Case 2 ####
# list of PlackettLuce models
R = matrix(c(1, 2, 3, 0,
              4, 1, 2, 3,
              2, 1, 3, 4,
              1, 2, 3, 0,
              2, 1, 3, 0,
              1, 0, 3, 2), nrow = 6, byrow = TRUE)
colnames(R) = c("apple", "banana", "orange", "pear")
mod1 = PlackettLuce(R)
R2 = matrix(c(1, 2, 0, 3,
               2, 1, 0, 3,
               2, 1, 0, 3,
               1, 2, 0, 3,
               2, 1, 0, 3,
               1, 3, 4, 2), nrow = 6, byrow = TRUE)
colnames(R2) = c("apple", "banana", "orange", "pear")
mod2 = PlackettLuce(R2)
mod = list(mod1, mod2)
regret(mod, n1 = 500)
```

reliability

Probability of outperforming a check

# Description

Measures the precision of estimated values, and the potential response to selection on those estimated values compared to a check

reliability 31

## Usage

```
reliability(x, ...)
## Default S3 method:
reliability(x, y = NULL, ...)
## S3 method for class 'PlackettLuce'
reliability(x, ref, ...)
## S3 method for class 'pltree'
reliability(x, ref, ...)
```

## **Arguments**

```
    a numeric vector, or an object of class PlackettLuce or pltree
    additional arguments passed to methods
    numeric, the reference value
    a character or integer for indexing the element containing reference values in x
```

#### Value

the reliability based on the worth parameters

## Author(s)

Kauê de Sousa, David Brown, Jacob van Etten

# References

Eskridge and Mumm (1992). Theoret. Appl. Genetics 84, 494–500 doi:10.1007/BF00229512.

32 resample

resample

Re-sample model estimates

# Description

Applies a k-fold approach to re-sample estimates from PlackettLuce model. The function will subset the data into 'k' number folds and re-calculate the model estimates. Optionally, a Bayesian bootstrapping technique can be used to increase output size and normalize the distribution of estimates

## Usage

```
resample(object, k = 5, bootstrap = FALSE, seed = NULL, ...)
```

# **Arguments**

object a PlackettLuce model object

k an integer for the number of bins to subset the data bootstrap logical, to run a Bayesian bootstrapping on object

seed integer, the seed for random number generation. If NULL (the default), gosset

will set the seed randomly

... additional arguments passed to methods, see details

rowbind 33

# **Details**

Additional details for Bayesian bootstrapping: statistic A function that accepts data as its first argument and possibly, the weights as its second, if use\_weights is TRUE; n1 The size of the bootstrap sample; n2 The sample size used to calculate the statistic each bootstrap draw

## Value

A data frame with re-sampled estimates

#### Author(s)

Kauê de Sousa

#### **Examples**

rowbind

Combine R objects by rows

#### **Description**

Combine R objects when number and names of columns do not match

34 set\_binomialfreq

#### Usage

```
rowbind(x, ...)
## Default S3 method:
rowbind(x, y, ...)
## S3 method for class 'list'
rowbind(x, ...)
```

#### **Arguments**

x a R object, typically a data.frame, matrix or list
 ... additional arguments passed to methods
 y a matrix, a data.frame (or any other object that can be coerced to data.frame)

#### Value

a data.frame with the combined data

## **Examples**

```
# two data frames
rowbind(airquality, mtcars)
# a list of data frames
1 = list(airquality, mtcars)
rowbind(1)
```

set\_binomialfreq

Binomial frequency rankings from pairwise contests

# Description

Binary comparisons from a ranking object. Ties are not taken into account, then they are added as NA's.

# Usage

```
set_binomialfreq(object, drop.null = FALSE, disaggregate = FALSE)
```

# Arguments

object an object of class rankings, grouped\_rankings or paircomp

drop.null logical, an optional argument to remove null contests

disaggregate logical, if TRUE binaries are disaggregated by individual contests

set\_paircomp 35

#### Value

A data.frame with binary rank of pairwise contests:

```
player1 a factor with n levels for the first player in the contests
player2 a factor with n levels (same as player1) for the second player in the contests
win1 number of times player1 wins against player2
win2 number of times player2 wins against player1
```

#### Author(s)

Kauê de Sousa

#### References

Turner H. & Firth D. (2012). Journal of Statistical Software, 48(9), 1–21. doi:10.18637/jss.v048.i09

#### See Also

```
Other rank functions: rank_numeric(), rank_tricot(), set_paircomp()
```

## **Examples**

set\_paircomp

Pairwise comparison rankings

## **Description**

Pairwise comparisons from a PlackettLuce ranking object. Ties are not taken into account, then coerced to NA's.

36 set\_paircomp

#### Usage

```
set_paircomp(object)
```

#### **Arguments**

object an object of class "rankings" or "grouped\_rankings" which is a matrix of dense rankings.

#### Value

an object of class "paircomp" which is a matrix of pairwise comparisons

#### Author(s)

Kauê de Sousa and Jacob van Etten

#### References

Strobl C., Wickelmaier F. & Zeileis A. (2011). Journal of Educational and Behavioral Statistics, 36(2), 135–153. doi:10.3102/1076998609359791

#### See Also

```
rankings, paircomp
Other rank functions: rank_numeric(), rank_tricot(), set_binomialfreq()
```

```
library("PlackettLuce")
R = matrix(c(1, 2, 0, 0,
             4, 1, 2, 3,
             2, 4, 3, 1,
             1, 2, 3, 0,
             2, 1, 1, 0,
             1, 0, 3, 2), nrow = 6, byrow = TRUE)
colnames(R) = c("apple", "banana", "orange", "pear")
R = as.rankings(R)
PC = set_paircomp(R)
# using breadwheat data
data("breadwheat", package = "gosset")
# convert the tricot rankings from breadwheat data
# into a object of class 'rankings' from PlackettLuce
R = rank_tricot(breadwheat,
               items = c("variety_a","variety_b","variety_c"),
               input = c("overall_best", "overall_worst"))
```

worth\_map 37

```
PC = set_paircomp(R)
```

worth\_map

Plot worth parameters

#### **Description**

Produces plots to highlight worth coefficients of items in a party tree of a list of PlackettLuce models

# Usage

```
worth_map(object, ...)
## Default S3 method:
worth_map(object, ...)
## S3 method for class 'list'
worth_map(object, labels, labels.order = NULL, items.order = NULL, ...)
worth_bar(object, ...)
```

# Arguments

object an object of class party or an object of class PlackettLuce or a list objects of class PlackettLuce
... additional arguments passed to methods
labels a vector with the name of models in object
labels.order optional, a vector to determine the order of labels
optional, a vector to determine the order of items

38 worth\_map

```
# Ranking of preference on four fruits
# based on traits taste, texture,
# price and storability
# taste
R1 = matrix(c(1, 2, 3, 4,
               4, 1, 3, 2,
               4, 1, 2, 3,
               1, 2, 0, 3), nrow = 4, byrow = TRUE)
colnames(R1) = c("apple", "banana", "orange", "pear")
mod1 = PlackettLuce(R1)
# texture
R2 = matrix(c(1, 4, 2, 3,
               1, 4, 3, 2,
               1, 4, 2, 3,
               1, 4, 2, 3), nrow = 4, byrow = TRUE)
colnames(R2) = c("apple", "banana", "orange", "pear")
mod2 = PlackettLuce(R2)
# price
R3 = matrix(c(2, 4, 3, 1,
               4, 1, 2, 3,
               3, 4, 2, 1,
               4, 3, 1, 2), nrow = 4, byrow = TRUE)
colnames(R3) = c("apple", "banana", "orange", "pear")
mod3 = PlackettLuce(R3)
# storability
R4 = matrix(c(1, 4, 3, 2,
               3, 4, 1, 2,
               1, 3, 2, 4,
               2, 3, 4, 1), nrow = 4, byrow = TRUE)
colnames(R4) = c("apple", "banana", "orange", "pear")
mod4 = PlackettLuce(R4)
# models in a list
mods = list(mod1, mod2, mod3, mod4)
# name for each model
labels = c("Taste", "Texture", "Price", "Storability")
worth_map(mods, labels)
# plot only one model as bar
worth_bar(mod1)
```

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