

Package ‘clubpro’

June 7, 2024

Type Package

Title Classification Using Binary Procrustes Rotation

Version 0.6.2

Description

Implements a classification method described by Grice (2011, ISBN:978-0-12-385194-9) using binary procrustes rotation; a simplified version of procrustes rotation.

License GPL (>= 3)

URL <https://timbeechey.github.io/clubpro/>

BugReports <https://github.com/timbeechey/clubpro/issues>

Encoding UTF-8

LazyData true

RoxygenNote 7.3.1

LinkingTo Rcpp, RcppArmadillo, RcppProgress

Imports Rcpp, lattice, stats, graphics

Suggests knitr, rmarkdown, tinytest

SystemRequirements C++17

Depends R (>= 2.10)

VignetteBuilder knitr

NeedsCompilation yes

Author Timothy Beechey [aut, cre] (<<https://orcid.org/0000-0001-8858-946X>>)

Maintainer Timothy Beechey <tim.beechey@proton.me>

Repository CRAN

Date/Publication 2024-06-07 15:50:02 UTC

Contents

accuracy	2
as.data.frame.clubprocsi	3
as.data.frame.clubprorand	4
caffeine	4
club	5
compare	6
csi	6
cval	7
individual_results	8
jellyfish	8
median_csi	9
n_ambiguous	9
n_correct	10
n_incorrect	11
pcc	11
pcc_replicates	12
plot.clubproaccuracy	13
plot.clubprocomparison	13
plot.clubprocsi	14
plot.clubprofit	15
plot.clubpropredictions	16
plot.clubprorand	16
plot.clubprothreshold	17
predict.clubprofit	18
summary.clubprocomparison	18
summary.clubprofit	19
threshold	20
Index	21

accuracy	<i>Classification accuracy for each observation.</i>
----------	--

Description

Classification accuracy for each observation.

Usage

accuracy(m)

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns a character vector containing a string corresponding to each observation indicating whether classification of that observation was "correct", "incorrect", or "ambiguous".

Value

a table

Examples

```
mod <- club(rate ~ dose, data = caffeine)
accuracy(mod)
```

as.data.frame.clubprocsi

Convert the output of csi() to a data.frame.

Description

Convert the output of csi() to a data.frame.

Usage

```
## S3 method for class 'clubprocsi'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

x	an object of class "clubprocsi"
row.names	ignored
optional	ignored
...	ignored

Details

This function is useful to format pcc replicates data for plotting.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
z <- csi(mod)
as.data.frame(z)
```

```
as.data.frame.clubprorand
```

Convert the output of pcc_replicates() to a data.frame.

Description

Convert the output of pcc_replicates() to a data.frame.

Usage

```
## S3 method for class 'clubprorand'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

x	an object of class "clubprorand"
row.names	ignored
optional	ignored
...	ignored

Details

This function is useful to format pcc replicates data for plotting.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
z <- pcc_replicates(mod)
as.data.frame(z)
```

```
caffeine
```

Caffeine data

Description

Effect of three different doses of caffeine on finger tapping rate.

Usage

```
caffeine
```

Format

A data frame with 30 rows and 2 columns:

dose dose of caffeine in mg

rate finger taps per minute

Source

Hand D.J., Daly F., Lunn A.D., McConway K.J., Ostrowski E. (1994) A Handbook of Small Data Sets. London: Chapman & Hall. Data set 50.

club	<i>Classify observations.</i>
------	-------------------------------

Description

club() is used to classify observations using binary procrustes rotation.

Usage

```
club(
  f,
  data,
  imprecision,
  nreps,
  normalise_cols,
  reorder_obs,
  display_progress
)
```

Arguments

f	a formula.
data	a data.frame.
imprecision	a number indicating the margin of imprecision allowed in classification.
nreps	the number of replicates to use in the randomisation test.
normalise_cols	a boolean indicating whether to normalise matrix columns.
reorder_obs	a string indicating the method for reordering observations to calculate c-values.
display_progress	a boolean indicating whether a progress bar should be displayed.

Value

an object of class "clubprofit" is a list containing the following components:

prediction a character vector of predicted classifications.

accuracy a character vector indicating whether each classification is "correct", "incorrect", or "ambiguous".

pcc the percentage of correct classifications.

cval the chance of randomly reordered data producing a PCC \geq the observed PCC.

pcc_replicates a vector of PCCs generated from randomly reordered data used to calculate cval.

call the matched call.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
```

compare	<i>Compare models.</i>
---------	------------------------

Description

Compare models.

Usage

```
compare(m1, m2)
```

Arguments

m1	an object of class "clubprofit" produced by club()
m2	an object of class "clubprofit" produced by club()

Details

Compare the PCC of two clubprofit models and compute the chance-value of the difference.

Value

an object of type "clubprocomparison"

Examples

```
m1 <- club(width ~ location, jellyfish)
m2 <- club(length ~ location, jellyfish)
compare(m1, m2)
```

csi	<i>Classification strength indices.</i>
-----	---

Description

Classification strength indices.

Usage

```
csi(m)
```

Arguments

m	an object of class "clubprofit" produced by club()
---	--

Details

Returns a vector containing the classification strength index for each observation.

Value

a numeric vector.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
csi(mod)
```

cval

Chance value.

Description

Chance value.

Usage

```
cval(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Compute the chance that randomly reordered data results in a percentage of correctly classified observations at least as high as the observed data.

Value

a numeric value.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
cval(mod)
```

`individual_results` *Individual level classification results.*

Description

Individual level classification results.

Usage

```
individual_results(m, digits)
```

Arguments

`m` an object of class "clubprofit" produced by `club()`
`digits` an integer

Details

Returns a data.frame containing predicted classifications and classification accuracy for each individual observation.

Value

a data.frame containing a columns of predictions and prediction accuracy

Examples

```
mod <- club(rate ~ dose, data = caffeine)
individual_results(mod)
```

`jellyfish` *Jellyfish dimension data*

Description

Sizes of jellyfish from two locations in the Hawkesbury River, New South Wales, Australia.

Usage

```
jellyfish
```

Format

A data frame with 46 rows and 3 columns:

location location where jellyfish was caught

width jellyfish width in mm

length jellyfish length in mm

Source

Hand D.J., Daly F., Lunn A.D., McConway K.J., Ostrowski E. (1994) A Handbook of Small Data Sets. London: Chapman & Hall. Data set 225.

median_csi	<i>Median classification strength index.</i>
------------	--

Description

Median classification strength index.

Usage

```
median_csi(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns the median classification strength index.

Value

a numeric vector.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
median_csi(mod)
```

n_ambiguous	<i>Number of ambiguous classifications.</i>
-------------	---

Description

Number of ambiguous classifications.

Usage

```
n_ambiguous(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns the number of observations which were classified ambiguously by the model.

Value

an integer.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
n_ambiguous(mod)
```

n_correct	<i>Number of correct classifications.</i>
-----------	---

Description

Number of correct classifications.

Usage

```
n_correct(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns the number of observations which were classified correctly by the model.

Value

an integer.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
n_correct(mod)
```

n_incorrect	<i>Number of incorrect classifications.</i>
-------------	---

Description

Number of incorrect classifications.

Usage

```
n_incorrect(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns the number of observations which were classified incorrectly by the model.

Value

an integer.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
n_incorrect(mod)
```

pcc	<i>Percentage of correct classifications.</i>
-----	---

Description

Percentage of correct classifications.

Usage

```
pcc(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns the percentage of correctly classified observations.

Value

a numeric value.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
pcc(mod)
```

pcc_replicates	<i>PCC replicates.</i>
----------------	------------------------

Description

PCC replicates.

Usage

```
pcc_replicates(m)
```

Arguments

`m` an object of class "clubprofit" produced by `club()`

Details

Returns an object containing a vector of PCC replicates used to calculate the chance-value.

Value

an object of class clubprorand.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
head(pcc_replicates(mod))
```

`plot.clubproaccuracy` *Plot accuracy.*

Description

Plot accuracy.

Usage

```
## S3 method for class 'clubproaccuracy'  
plot(x, ...)
```

Arguments

`x` an object of class "clubproaccuracy"
`...` ignored

Details

Produces a mosaic plot of prediction accuracy by category

Value

called for side-effects only

Examples

```
mod <- club(rate ~ dose, data = caffeine)  
z <- accuracy(mod)  
plot(z)
```

`plot.clubprocomparison`
Plot model comparison.

Description

Plot model comparison.

Usage

```
## S3 method for class 'clubprocomparison'  
plot(x, ...)
```

Arguments

x an object of class "clubprocomparison".
... ignored

Details

Plot a distribution of PCCs computed from randomly reordered data used to calculate the chance-value for a model comparison.

Value

no return value, called for side effects only.

Examples

```
m1 <- club(width ~ location, jellyfish)
m2 <- club(length ~ location, jellyfish)
z <- compare(m1, m2)
plot(z)
```

plot.clubprocsi *Plot classification strength indices.*

Description

Plot classification strength indices.

Usage

```
## S3 method for class 'clubprocsi'
plot(x, ...)
```

Arguments

x an object of class "clubprocsi"
... ignored

Details

Produces dotplot showing classification strength for each individual.

Value

called for side-effects only

Examples

```
mod <- club(rate ~ dose, data = caffeine)
z <- csi(mod)
plot(z)
```

plot.clubprofit *Plot classification accuracy.*

Description

Plot classification accuracy.

Usage

```
## S3 method for class 'clubprofit'
plot(x, ...)
```

Arguments

x	an object of class "clubprofit" produced by club()
...	ignored

Details

Produces bar plot showing counts of individuals against observed values within each target grouping. Fill colours indicate whether each individual was classified correctly, incorrectly or ambiguously.

Value

called for side-effects only

Examples

```
mod <- club(rate ~ dose, data = caffeine)
plot(mod)
```

plot.clubpropredictions

Plot predictions.

Description

Plot predictions.

Usage

```
## S3 method for class 'clubpropredictions'  
plot(x, ...)
```

Arguments

x	an object of class "clubpropredictions"
...	ignored

Details

Produces a mosaic plot of observed versus predicted categories

Value

called for side-effects only

Examples

```
mod <- club(rate ~ dose, data = caffeine)  
z <- predict(mod)  
plot(z)
```

plot.clubprorand

Plot PCC replicates.

Description

Plot PCC replicates.

Usage

```
## S3 method for class 'clubprorand'  
plot(x, ...)
```

Arguments

x an object of class "clubprofit" produced by club()
... ignored

Details

Plot the distribution of PCCs computed from randomly reordered data used to calculate the chance-value.

Value

no return value, called for side effects only.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
plot(pcc_replicates(mod))
```

`plot.clubprothreshold` *Plot PCC as a function of binary category boundary location.*

Description

Plot PCC as a function of binary category boundary location.

Usage

```
## S3 method for class 'clubprothreshold'
plot(x, ...)
```

Arguments

x an object of class "clubprothreshold"
... ignored

Details

Produces an xyplot showing the PCC returned for each possible category boundary.

Value

called for side-effects only

Examples

```
mod <- club(rate ~ dose, data = caffeine)
z <- threshold(mod)
plot(z)
```

predict.clubprofit *Predicted category for each observation.*

Description

Predicted category for each observation.

Usage

```
## S3 method for class 'clubprofit'
predict(object, ...)
```

Arguments

object	an object of class "clubprofit" produced by club()
...	ignored

Details

Returns a character vector containing the name of the predicted category for each observed value.

Value

a table

Examples

```
mod <- club(rate ~ dose, data = caffeine)
predict(mod)
```

summary.clubprocomparison

Generate a summary of a comparison of clubprofit models.

Description

Generate a summary of a comparison of clubprofit models.

Usage

```
## S3 method for class 'clubprocomparison'
summary(object, ...)
```

Arguments

object	an object of class "clubprocomparison".
...	ignored

Value

No return value, called for side effects.

Examples

```
m1 <- club(width ~ location, jellyfish)
m2 <- club(length ~ location, jellyfish)
z <- compare(m1, m2)
summary(z)
```

summary.clubprofit	<i>Generate a summary of results from a fitted clubpro model.</i>
--------------------	---

Description

Generate a summary of results from a fitted clubpro model.

Usage

```
## S3 method for class 'clubprofit'
summary(object, ...)
```

Arguments

object	an object of class "clubprofit".
...	ignored

Value

No return value, called for side effects.

Examples

```
mod <- club(rate ~ dose, data = caffeine)
summary(mod)
```

threshold	<i>Classification strength indices.</i>
-----------	---

Description

Classification strength indices.

Usage

```
threshold(m)
```

Arguments

m an object of class "clubprofit" produced by club()

Details

Returns a vector containing the classification strength index for each observation.

Value

an object of class clubprothreshold

Examples

```
mod <- club(width ~ location, data = jellyfish)
threshold(mod)
```

Index

* datasets

- caffeine, 4
- jellyfish, 8

accuracy, 2

as.data.frame.clubprocsi, 3

as.data.frame.clubprorand, 4

caffeine, 4

club, 5

compare, 6

csi, 6

cval, 7

individual_results, 8

jellyfish, 8

median_csi, 9

n_ambiguous, 9

n_correct, 10

n_incorrect, 11

pcc, 11

pcc_replicates, 12

plot.clubproaccuracy, 13

plot.clubprocomparison, 13

plot.clubprocsi, 14

plot.clubprofit, 15

plot.clubpropredictions, 16

plot.clubprorand, 16

plot.clubprothreshold, 17

predict.clubprofit, 18

summary.clubprocomparison, 18

summary.clubprofit, 19

threshold, 20