Package 'opa'

March 11, 2024

```
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      and longitudinal data, as described by Thorngate (1987)
      <a href="https://doi.org/10.1016/S0166-4115(08)60083-7">doi:10.1016/S0166-4115(08)60083-7</a>> and Grice et al., (2015)
      <doi:10.1177/2158244015604192>. Hypothesis and data are encoded as
      pairwise relative orderings which are then compared to determine the
      percentage of orderings in the data that are matched by the hypothesis.
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```

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Description

A data frame with 20 rows and 14 columns containing times between visits to a mechanical flower by bees in two experimental conditions.

Usage

bees

compare_conditions 3

Format

bee Unique identifier for each individual bee.

condition Factor identifying the two experimental conditions. In the frustrated condition bees were temporarily restricted from returning to the hive after collecting nectar, in the free condition bees were able to return to the hive without delay.

t1-t12 Time between visits to the mechanical flower (in seconds) in each of 12 consecutive trials.

Source

Grice, J. W., Craig, D. P. A., & Abramson, C. I. (2015). A Simple and Transparent Alternative to Repeated Measures ANOVA. SAGE Open, 5(3), 215824401560419. https://doi.org/10.1177/2158244015604192

compare_conditions Calculates PCCs and c-values based on pairwise comparison of conditions.

Description

Calculates PCCs and c-values based on pairwise comparison of conditions.

Usage

```
compare_conditions(result, nreps = 1000L)
```

Arguments

```
result an object of class "opafit" produced by a call to opa().

nreps an integer
```

Value

compare_conditions returns a list with the following elements

pcc_mat A lower triangle matrix containing PCCs calculated from each pairing of data columns.

cval_mat A lower triangle matrix containing c-values calculated from each pairing of data columns.

pccs A vector containing PCCs calculated from each pairing of data.

cvals A vector containing c-values calculated from each pairing of data.

nreps The number of permutations used to calculate the c-values.

compare_groups	Calculate the c-value of the difference in PCCs produced by two
	groups

Description

Calculate the c-value of the difference in PCCs produced by two groups

Usage

```
compare_groups(m, group1, group2, two_tailed)
```

Arguments

```
m an object of class "opafit" produced by a call to opa().

group1 a character string which matches a group level passed to opa().

group2 a character string which matches a group level passed to opa().

two_tailed a boolean indicating whether the comparison is two-tailed.
```

Value

an object of class "opaGroupComparison".

Examples

compare_hypotheses

Calculate the c-value of the difference in PCCs produced by two hypotheses

Description

Calculate the c-value of the difference in PCCs produced by two hypotheses

```
compare_hypotheses(m1, m2, two_tailed)
```

correct_pairs 5

Arguments

```
m1 an object of class "opafit" produced by a call to opa().
m2 an object of class "opafit" produced by a call to opa().
two_tailed a boolean indicating whether the comparison is two-tailed.
```

Value

an object of class "opaHypothesisComparison".

Examples

correct_pairs

Return the number of pairs of observations matched by the hypothesis

Description

Return the number of pairs of observations matched by the hypothesis

Usage

```
correct_pairs(m)
```

Arguments

m an object of class "opafit"

Value

a non-negative integer

group_cvals

cval_plot

Plot individual chance values

Description

Plot individual chance values

Usage

```
cval_plot(m)
```

Arguments

m

an object of class "opafit"

Value

No return value, called for side effects.

Examples

group_cvals

Return the group chance values of the specified model

Description

Return the group chance values of the specified model

Usage

```
group_cvals(m)
```

Arguments

m

an object of class "opafit"

Value

a numeric vector

group_pccs 7

Examples

```
\begin{array}{lll} \text{dat} \leftarrow & \text{data.frame}(\text{t1 = c(9, 4, 8, 10),} \\ & & \text{t2 = c(8, 8, 12, 10),} \\ & & \text{t3 = c(8, 5, 10, 11))} \\ \text{h} \leftarrow & \text{hypothesis(1:3)} \\ \text{opamod} \leftarrow & \text{opa(dat, h)} \\ \text{group\_cvals(opamod)} \end{array}
```

group_pccs

Return the group PCCs of the specified model

Description

Return the group PCCs of the specified model

Usage

```
group_pccs(m)
```

Arguments

m

an object of class "opafit"

Value

a numeric vector

Examples

 $group_results$

Group-level PCC and chance values.

Description

Group-level PCC and chance values.

```
group_results(m, digits)
```

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Arguments

```
m an object of class "opafit" produced by opa().
digits a positive integer.
```

Details

If the model was fitted with no grouping variable, a single PCC and c-value are returned. If a grouping variable was specified in the call to opa then PCCs and c-values are returned for each factor level of the grouping variable.

Value

a matrix with 1 row per group.

Examples

hypothesis

Create a hypothesis object

Description

Create a hypothesis object

Usage

```
hypothesis(xs, type = "pairwise")
```

Arguments

```
xs a numeric vector type a string
```

Value

a list containing the following elements

```
h1 <- hypothesis(c(2, 1, 3, 4), type = "pairwise")
h2 <- hypothesis(c(2, 1, 3, 4), type = "adjacent")
```

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incorrect_pairs

Return the number of pairs of observations not matched by the hypothesis

Description

Return the number of pairs of observations not matched by the hypothesis

Usage

```
incorrect_pairs(m)
```

Arguments

m

an object of class "opafit"

Value

a non-negative integer

Examples

individual_cvals

Return the individual chance values of the specified model

Description

Return the individual chance values of the specified model

Usage

```
individual_cvals(m)
```

Arguments

m

an object of class "opafit"

Value

a numeric vector

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Examples

individual_pccs

Return the individual PCCs of the specified model

Description

Return the individual PCCs of the specified model

Usage

```
individual_pccs(m)
```

Arguments

m

an object of class "opafit"

Value

a numeric vector

Examples

 $individual_results$

Individual-level PCC and chance values.

Description

Individual-level PCC and chance values.

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

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Arguments

```
m an object of class "opafit" produced by opa()
digits an integer
```

Details

If the opa model was fitted with no grouping variable, a matrix of PCCs and c-values are returned corresponding to the order of rows in the data. If the opa model was fitted with a grouping variable specified, a table of PCCs and c-values is returned ordered by factor level of the grouping variable.

Value

a matrix containing a column of PCC values and a column of c-values with 1 row per row of data.

Examples

ора

Fit an ordinal pattern analysis model

Description

opa is used to fit ordinal pattern analysis models by computing the percentage of pair orderings in each row of data which are matched by corresponding pair orderings in an hypothesis, in addition the chance of a permutation of the data producing a percentage match as great.

```
opa(
  dat,
  hypothesis,
  group = NULL,
  pairing_type = "pairwise",
  diff_threshold = 0,
  nreps = 1000L,
  shuffle_across_individuals = FALSE
)
```

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Arguments

dat a data frame
hypothesis a numeric vector

group an optional factor vector

pairing_type a string

diff_threshold a positive integer or floating point number
nreps an integer, ignored if cval_method = "exact"

shuffle_across_individuals

a boolean indicating whether to randomize data across individuals in c-value

computation.

Details

Data is expected in **wide** format with 1 row per individual and 1 column per measurement condition. Data must contain only columns consisting of numerical values of the *dependent* variable.

The length of the hypothesis must be equal to the number of columns in the dependent variable data.frame dat.

Any *independent* variable must be passed separately as a vector with the group keyword. The grouping vector must be a *factor*.

pairing_type must be either "pairwise" or "adjacent". The "pairwise" option considered the relative ordering of every pair of observations in the data and every pair of elements of the hypothesis. The "adjacent" option considers the ordering of adjacent pairs only. If unspecified, the default is "pairwise".

diff_threshold may be a positive integer or double. If unspecified a default zero threshold is used. The diff_threshold is never applied to the hypothesis.

nreps specifies the number of random reorderigs to use in the calculation of chance-values.

Value

opa returns an object of class "opafit".

An object of class "opafit" is a list containing the following components:

group_pcc the percentage of pairwise orderings from all pooled data rows which were correctly classified by the hypothesis.

individual_pccs a vector containing the percentage of pairwise orderings that were correctly classified by the hypothesis for each data row.

correct_pairs an integer representing the number of pairwise orderings pooled across all data rows that were correctly classified by the hypothesis.

total_pairs an integer, the number of pair orderings contained in the data.

group_cval the group-level chance value.

individual_cvals a vector containing chance values for each data row

rand_pccs A vector of PCCS calculated from each random ordering with length equal to nreps, a list of vectors if a group vector was passed to opa().

pcc_plot 13

call The matched call

hypothesis The hypothesis vector passed to opa()

pairing_type A string indicating the method of pairing passed to opa().

diff_threshold The numeric difference threshold used to calculate PCCs. If no value was passed in the diff_threshold, the default of 0 is used.

data The data.frame passed to opa().

groups The vector of groups passed to opa. If no group vector was passed to opa() the default of NULL is used.

nreps an integer, the number of random re-orderings of the data used to compute chance values.

References

Grice, J. W., Craig, D. P. A., & Abramson, C. I. (2015). A Simple and Transparent Alternative to Repeated Measures ANOVA. SAGE Open, 5(3), 215824401560419. https://doi.org/10.1177/2158244015604192

Thorngate, W. (1987). Ordinal Pattern Analysis: A Method for Assessing Theory-Data Fit. Advances in Psychology, 40, 345–364. https://doi.org/10.1016/S0166-4115(08)60083-7>

Examples

pcc_plot

Plot individual PCCs.

Description

Plot individual PCCs.

Usage

```
pcc_plot(m)
```

Arguments

m

an object of class "opafit"

Value

No return value, called for side effects.

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Examples

pituitary

Childhood growth data

Description

A data frame with 108 rows and 4 columns containing data on the distance from the pituitary to the pteryo-maxillary fissure in children.

Usage

pituitary

Format

distance Distance in mm from the pituitary to the pteryo-maxillary fissure.

age Age in years.

individual Unique identifier for each individual.

sex Biological sex of each individual.

Source

Potthoff, R. F., & Roy, S. N. (1964). A Generalized Multivariate Analysis of Variance Model Useful Especially for Growth Curve Problems. Biometrika, 51(3/4), 313–326. https://doi.org/10.2307/2334137

plot.opafit

Plots individual-level PCCs and chance-values.

Description

Plots individual-level PCCs and chance-values.

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

Arguments

```
x an object of class "opafit" produced by opa()
... ignored
```

Value

No return value, called for side effects.

Examples

```
\begin{array}{lll} \text{dat} <& -\text{ data.frame}(\text{t1 = c(9, 4, 8, 10),} \\ && \text{t2 = c(8, 8, 12, 10),} \\ && \text{t3 = c(8, 5, 10, 11))} \\ \text{h} <& -\text{ hypothesis(1:3)} \\ \text{opamod} <& -\text{ opa(dat, h)} \\ \text{plot(opamod)} \end{array}
```

plot.opaGroupComparison

Plot group comparison PCC replicates.

Description

Plot group comparison PCC replicates.

Usage

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

Arguments

```
x an object of class "oparandpccs" produced by random_pccs()
... ignored
```

Details

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

Value

no return value, called for side effects only.

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Examples

```
\label{eq:data-frame} \begin{array}{lll} \mbox{data-frame}(\mbox{group} = \mbox{c("a", "b", "a", "b")}, \\ & \mbox{t1} = \mbox{c(9, 4, 8, 10)}, \\ & \mbox{t2} = \mbox{c(8, 8, 12, 10)}, \\ & \mbox{t3} = \mbox{c(8, 5, 10, 11)}) \\ \mbox{dat$group} <- \mbox{factor}(\mbox{dat$group}, \mbox{levels} = \mbox{c("a", "b")}) \\ \mbox{h} <- \mbox{hypothesis(1:3)} \\ \mbox{opamod} <- \mbox{opa}(\mbox{dat$[,2:4], h, group} = \mbox{dat$group}) \\ \mbox{z} <- \mbox{compare\_groups}(\mbox{opamod}, "a", "b") \\ \mbox{plot(z)} \end{array}
```

plot.opahypothesis

Plot a hypothesis.

Description

Plot a hypothesis.

Usage

```
## S3 method for class 'opahypothesis'
plot(x, title = TRUE, ...)
```

Arguments

```
x an object of class "opaHypothesis"title a boolean indicating whether to include a plot title... ignored
```

Value

No return value, called for side effects.

```
h \leftarrow hypothesis(c(1,2,3,3,3))
plot(h)
```

```
plot.opaHypothesisComparison
```

Plot hypothesis comparison PCC replicates.

Description

Plot hypothesis comparison PCC replicates.

Usage

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

Arguments

```
x an object of class "oparandpccs" produced by random_pccs()... ignored
```

Details

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

Value

no return value, called for side effects only.

print.opafit

plot.oparandpccs

Plot PCC replicates.

Description

Plot PCC replicates.

Usage

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

Arguments

```
x an object of class "oparandpccs" produced by random_pccs()... ignored
```

Details

Plot a histogram of PCCs computed from randomly reordered data used to calculate the chance-value

Value

no return value, called for side effects only.

Examples

```
\label{eq:data-frame} \begin{array}{lll} \mbox{dat} <- \mbox{ data.frame}(\mbox{t1 = c(9, 4, 8, 10),} \\ & \mbox{t2 = c(8, 8, 12, 10),} \\ & \mbox{t3 = c(8, 5, 10, 11))} \\ \mbox{h <- hypothesis(1:3)} \\ \mbox{opamod <- opa(dat, h)} \\ \mbox{plot(random_pccs(opamod))} \end{array}
```

print.opafit

Displays the call used to fit an ordinal pattern analysis model.

Description

Displays the call used to fit an ordinal pattern analysis model.

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

Arguments

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

Value

No return value, called for side effects.

Examples

```
\begin{array}{lll} \text{dat} \leftarrow & \text{data.frame}(\texttt{t1} = \texttt{c(9, 4, 8, 10)}, \\ & & \texttt{t2} = \texttt{c(8, 8, 12, 10)}, \\ & & & \texttt{t3} = \texttt{c(8, 5, 10, 11)}) \\ \text{h} \leftarrow & \text{hypothesis(1:3)} \\ \text{opamod} \leftarrow & \text{opa(dat, h)} \\ \text{print(opamod)} \end{array}
```

print.opaGroupComparison

Prints a summary of results from hypothesis comparison.

Description

Prints a summary of results from hypothesis comparison.

Usage

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

Arguments

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

Value

No return value, called for side effects.

```
z <- compare_groups(opamod, "a", "b")
print(z)</pre>
```

print.opahypothesis

Print details of a hypothesis

Description

Print details of a hypothesis

Usage

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

Arguments

```
x an object of type "opaHypothesis"
... ignored
```

Value

No return value, called for side-effects.

Examples

```
h1 <- hypothesis(c(2, 1, 3, 4), type = "pairwise")
print(h1)
h2 <- hypothesis(c(2, 1, 3, 4), type = "adjacent")
print(h2)</pre>
```

 $\verb"print.opaHypothesisComparison"$

Prints a summary of results from hypothesis comparison.

Description

Prints a summary of results from hypothesis comparison.

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

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Arguments

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

Value

No return value, called for side effects.

Examples

```
\label{eq:data-frame} \begin{array}{llll} \text{dat} &<- \text{ data.frame}(\text{t1} = \text{c(9, 4, 8, 10)}, \\ & \text{t2} = \text{c(8, 8, 12, 10)}, \\ & \text{t3} = \text{c(8, 5, 10, 11)}, \\ & \text{t4} = \text{c(10, 5, 11, 12)}) \\ \text{h1} &<- \text{ hypothesis}(\text{c(1, 2, 3, 4)}) \\ \text{h2} &<- \text{ hypothesis}(\text{c(1, 4, 2, 3)}) \\ \text{opamod1} &<- \text{ opa}(\text{dat, h1}) \\ \text{opamod2} &<- \text{ opa}(\text{dat, h2}) \\ \text{z} &<- \text{ compare\_hypotheses}(\text{opamod1, opamod2}) \\ \text{print}(\text{z}) \end{array}
```

Description

Displays the results of a pairwise ordinal pattern analysis.

Usage

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

Arguments

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

Value

No return value, called for side effects.

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Examples

```
\label{eq:data-frame} \begin{array}{lll} \mbox{dat} & -\mbox{data.frame}(\mbox{t1 = c(9, 4, 8, 10),} \\ & \mbox{t2 = c(8, 8, 12, 10),} \\ & \mbox{t3 = c(8, 5, 10, 11))} \\ \mbox{h <- hypothesis(1:3)} \\ \mbox{opamod <- opa(dat, h)} \\ \mbox{pw <- compare_conditions(opamod)} \\ \mbox{print(pw, digits = 2)} \end{array}
```

random_pccs

Return the random order generated PCCs used to calculate the group chance value

Description

Return the random order generated PCCs used to calculate the group chance value

Usage

```
random_pccs(m)
```

Arguments

m

an object of class "opafit"

Value

a numeric vector

```
\label{eq:data-frame} \begin{array}{lll} \mbox{data.frame}(\mbox{t1 = c(9, 4, 8, 10),} \\ & \mbox{t2 = c(8, 8, 12, 10),} \\ & \mbox{t3 = c(8, 5, 10, 11))} \\ \mbox{h <- hypothesis(1:3)} \\ \mbox{opamod <- opa(dat, h)} \\ \mbox{random\_pccs(opamod)} \end{array}
```

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summary.opafit	Prints a summary of results from a fitted ordinal pattern analysis
	model.

Description

Prints a summary of results from a fitted ordinal pattern analysis model.

Usage

```
## S3 method for class 'opafit'
summary(object, ..., digits = 2L)
```

Arguments

```
object an object of class "opafit".
... ignored
digits an integer used for rounding values in the output.
```

Value

No return value, called for side effects.

Examples

summary.opaGroupComparison

Prints a summary of results from hypothesis comparison.

Description

Prints a summary of results from hypothesis comparison.

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

Arguments

```
object an object of class "opaHypothesisComparison". . . . ignored
```

Value

No return value, called for side effects.

Examples

```
\label{eq:data-frame} (\text{group} = \text{c("a", "b", "a", "b")}, \\ & \text{t1} = \text{c(9, 4, 8, 10)}, \\ & \text{t2} = \text{c(8, 8, 12, 10)}, \\ & \text{t3} = \text{c(8, 5, 10, 11)}) \\ \text{dat\$group} \leftarrow \text{factor(dat\$group, levels} = \text{c("a", "b")}) \\ \text{h} \leftarrow \text{hypothesis(1:3)} \\ \text{opamod} \leftarrow \text{opa(dat[,2:4], h, group} = \text{dat\$group}) \\ \text{z} \leftarrow \text{compare\_groups(opamod, "a", "b")} \\ \text{summary(z)} \\ \end{cases}
```

summary.opaHypothesisComparison

Prints a summary of results from hypothesis comparison.

Description

Prints a summary of results from hypothesis comparison.

Usage

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

Arguments

```
object an object of class "opaHypothesisComparison".
... ignored
```

Value

No return value, called for side effects.

```
\label{eq:data-frame} \begin{array}{lll} \text{dat} <& -\text{ data.frame}(\text{t1} = \text{c(9, 4, 8, 10)}, \\ & & \text{t2} = \text{c(8, 8, 12, 10)}, \\ & & \text{t3} = \text{c(8, 5, 10, 11)}, \\ & & \text{t4} = \text{c(10, 5, 11, 12)}) \\ \text{h1} <& -\text{ hypothesis}(\text{c(1, 2, 3, 4)}) \\ \text{h2} <& -\text{ hypothesis}(\text{c(1, 4, 2, 3)}) \\ \text{opamod1} <& -\text{ opa}(\text{dat, h1}) \\ \text{opamod2} <& -\text{ opa}(\text{dat, h2}) \\ \text{z} <& -\text{ compare\_hypotheses}(\text{opamod1, opamod2}) \\ \text{summary}(\text{z}) \end{array}
```

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