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Author Frank Schaarschmidt, Daniel Gerhard, Martin Sill

Maintainer Frank Schaarschmidt <schaarschmidt@biostat.uni-hannover.de>

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Description Multiple contrast tests and simultaneous confidence intervals based on normal approximation. With implementations for binomial proportions in a 2xk setting (risk difference and odds ratio), poly-3-adjusted tumour rates, biodiversity indices (multinomial data) and expected values under lognormal assumption. Approximative power calculation for multiple contrast tests of binomial and Gaussian data.

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MCPAN-package

Multiple comparison procedures based on normal approximation.

Description

Multiple contrast tests and simultaneous confidence intervals using normal approximation, if individuals are randomly assigned to treatments in a oneway layout. For dichotomous variables, approximate confidence intervals for the risk difference, risk ratio and odds ratio are available. If the variable of interest is the rate of tumours in long-term rodent carcinogenicity trials (without cause of death information), confidence intervals for poly-k-adjusted tumour rates are available. For abundance data of multiple species, approximate simultaneous confidence intervals for differences of Simpson and Shannon-indices are implemented.

Please note that not all methods in this package are published.

Details

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Version: 1.1-14
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Author(s)

Frank Schaarschmidt, Daniel Gerhard, Martin Sill Maintainer: Frank Schaarschmidt <schaarschmidt@biostat.uni-hannover.de>

References

Reference for multiple differences of proportions:

Schaarschmidt, F., Sill, M., and Hothorn, L.A. (2008): Approximate Simultaneous Confidence Intervals for Multiple Contrasts of Binomial Proportions. Biometrical Journal 50, 782-792.

See [binomRDci](#) for the code that reproduces the examples in this reference.

Reference for multiple differences of poly-k-rates:

Schaarschmidt, F., Sill, M., and Hothorn, L.A. (2008): Poly-k-trend tests for survival adjusted analysis of tumor rates formulated as approximate multiple contrast test. Journal of Biopharmaceutical Statistics 18, 934-948.

See [poly3ci](#) for the code that reproduces the example in this reference.

Simultaneous confidence intervals for Dunnett-type comparisons to control using odds ratios:

Holford, T.R., Walter, S.D. and Dunnett, C.W. (1989): Simultaneous interval estimates of the odds ratio in studies with two or more comparisons. Journal of Clinical Epidemiology 42, 427-434.

Background references:

The Add-2 Confidence interval for the difference of two binomial proportions:

Agresti, A. and Caffo, B.(2000): Simple and effective confidence intervals for proportions and differences of proportions result from adding two successes and two failures. *American Statistician* 54, p. 280-288.

and its generalization to a single contrast of several proportions in:

Price, R.M. and Bonett, D.G. (2004): An improved confidence interval for a linear function of binomial proportions. *Computational Statistics and Data Analysis* 45, 449-456.

For long-term rodent carcinogenicity data: The assumptions of poly-3-adjustment are outlined in:

Bailer, J.A. and Portier, C.J. (1988): Effects of treatment-induced mortality and tumor-induced mortality on tests for carcinogenicity in small samples. *Biometrics* 44, 417-431.

Peddada, S.D., Dinse, G.E., and Haseman, J.K. (2005): A survival-adjusted quantal response test for comparing tumor incidence rates. *Applied Statistics* 54, 51-61.

For correlation of multiple contrasts of binomial proportions, see: *Bretz F, Hothorn L.:* Detecting dose-response using contrasts: asymptotic power and sample size determination for binomial data. *Statistics in Medicine* 2002; 21: 3325-3335.

Examples

```
## # 1)
# Adjusted p-values and simultaneous confidence intervals
# for 2xk tables of binomial data:
# binomRDtest, binomRDci

# Difference of proportions

binomRDtest(x=c(2,6,4,13), n=c(34,33,36,34),
  names=c("Placebo", "50", "75", "150"),
  type="Dunnett", method="ADD1")

binomRDci(x=c(2,6,4,13), n=c(34,33,36,34),
  names=c("Placebo", "50", "75", "150"),
  type="Dunnett", method="ADD1")

# Odds ratios:

binomORci(x=c(2,6,4,13), n=c(34,33,36,34),
  names=c("Placebo", "50", "75", "150"),
  type="Dunnett")

# For more details on evaluation,
# see:
# ?liarozole

data(liarozole)
```

```

# # # 2)
# Adjusted p-values and simultaneous confidence intervals
# for poly-3-adjusted tumour rates:
# poly3test, poly3ci

data(methyl)
methyl

# poly-3-adjusted sample estimates:

poly3estf(time=methyl$death,
           status=methyl$tumour,
           f=methyl$group)

# Simultaneous Add-1-confidence intervals
# for difference to the control group:

poly3ci(time=methyl$death, status=methyl$tumour,
         f=methyl$group, method="ADD1",
         type="Dunnett", alternative="greater")

# Test for trend, based on Changepoint contrasts:

poly3test(time=methyl$death, status=methyl$tumour,
          f=methyl$group, method="ADD1",
          type="Changepoint", alternative="greater")

# # # 3) Plot of confidence intervals
# created by binomRDci and poly3ci:

MethylCI <- poly3ci(time=methyl$death, status=methyl$tumour,
                   f=methyl$group, method="ADD1",
                   type="Dunnett", alternative="greater")

plot(MethylCI)

```

binomest

Only for internal use.

Description

Groupwise point and variance estimates for binomials, if data are given as numeric vectors of successes x and trials n , with I the number of levels in the one-way layout

Usage

```
binomest(x, ...)
```

```
## Default S3 method:
binomest(x, n, names=NULL,
         method="Wald", success=NULL, ...)

## S3 method for class 'formula'
binomest(formula, data,
         method="Wald", success=NULL, ...)

## S3 method for class 'table'
binomest(x, method="Wald",
         success=NULL, ...)
```

Arguments

x	either a numeric vector of the number of successes, then n must be specified, or an object of class "table" as can be obtained by using table on a data.frame with to categorical variables
n	a numeric vector of the number of trials, only needed, if x is a numeric vector
names	optional character vector of group names (labels for later output), must have the same length as x and n
formula	a two-sided formula of the style 'response ~ treatment', where 'response' should be a categorical variable with two levels, while treatment should be a factor specifying the treatment levels
data	a data.frame, containing the variables specified in formula
method	a character string, specifying the method for variance estimator, with options "Wald", "ADD1", "ADD2"
success	an optional character string, to indicate the level considered as success if data are given as table, or data.frame, or, serving as label for the event considered as success in later functions
...	currently not used

Details

Only for internal use.

Value

Y	a numeric vector of the groupwise number of successes
n	a numeric vector of the groupwise number of trials (observations)
estimate	a numeric vector of groupwise point estimates for the probability of success, Y/n
estp	a numeric vector of groupwise interval midpoints
varp	a numeric vector of groupwise variance estimators, to be used for CI construction

varcor a numeric vector of groupwise variance estimators, to be used for correlation structures and test these values are adjusted to be larger than zero also in cases where $Y=0$ or $Y=n$

names a character vector of the level names of f

success a character string naming the category which has been defined as success

Examples

```
# if data are available as counts:

nsuccess<-c(1,2,6,8)
ntrials<-c(20,20,20,20)
binomest(x=nsuccess, n=ntrials)

binomest(x=nsuccess, n=ntrials,
  names=c("Control", "A", "B", "C"))

# if data are available as data.frame
# with categorical response variable
# and factor as grouping variable

data(liarozole)
binomest(Improved ~ Treatment, data=liarozole)
binomest(Improved ~ Treatment, data=liarozole, success="y")

# if data are available as table
# and factor as grouping variable

data(liarozole)
tab<-table(liarozole)

binomest(tab)
binomest(tab, success="y")
```

Description

Approximate simultaneous confidence intervals for (weighted geometric means of) odds ratios are constructed. Estimates are derived from fitting a glm on the logit-link, approximate intervals are constructed on the log-link, and transformed to original scale.

Usage

```
binomORci(x, ...)

## Default S3 method:
binomORci(x, n, names = NULL,
  type = "Dunnett", method="GLM", cmat = NULL,
  alternative = "two.sided", conf.level = 0.95,
  dist="MVN", ...)

## S3 method for class 'formula'
binomORci(formula, data,
  type = "Dunnett", method="GLM", cmat = NULL,
  alternative = "two.sided", conf.level = 0.95,
  dist="MVN", ...)

## S3 method for class 'table'
binomORci(x,
  type = "Dunnett",method="GLM", cmat = NULL,
  alternative = "two.sided", conf.level = 0.95,
  dist="MVN", ...)

## S3 method for class 'matrix'
binomORci(x,
  type = "Dunnett", method="GLM", cmat = NULL,
  alternative = "two.sided", conf.level = 0.95,
  dist="MVN", ...)
```

Arguments

x	a numeric vector, giving the number of successes in I independent samples, or an object of class "table", representing the 2xk-table, or an object of class "matrix", representing the 2xk-table
n	numeric vector, giving the number of trials (i.e. the sample size) in each of the I groups (only required if x is a numeric vector, ignored otherwise)
names	an optional character string, giving the names of the groups/ sample in x, n; if not specified the possible names of x are taken as group names (ignored if x is a table or matrix)
formula	a two-sided formula of the style 'response ~ treatment', where 'response' should be a categorical variable with two levels, while treatment should be a factor specifying the treatment levels
data	a data.frame, containing the variables specified in formula
type	a character string, giving the name of a contrast method, as defined in <code>contrMat(multcomp)</code> ; ignored if cmat is sepcified
method	a single character string, specifying the method for confidence interval computation; Options are "GLM" and "Woolf". "GLM" takes the maximum likelihood estimates and the their standard errors; this yields a conservative confidence intervals with uninformative limits if x=0 and x=n occures. "Woolf" adds 0.5 to

	the cell counts, resulting in less conservative bounds. These can be liberal when extreme proportions are compared.
<code>cmat</code>	a optional contrast matrix
<code>alternative</code>	a single character string, one of "two.sided", "less", "greater"
<code>conf.level</code>	a single numeric value, simultaneous confidence level
<code>dist</code>	a character string, "MVN" invokes multiplicity adjustment via the multivariate normal distribution, "N" invokes use of quantiles of the univariate normal distribution
<code>...</code>	arguments to be passed to <code>binomest</code> , currently only success labelling the event which should be considered as success

Details

This function calls `glm` and fits a one-way-model with family binomial on the logit-link. Then, the point estimates and variances estimates from the fit are taken to construct simultaneous confidence intervals for differences (of weighted arithmetic means) of log-odds. Applying the exponential function to these intervals on the logit scale yields intervals for ratios (of weighted geometric) of odds. For simple groupwise comparisons, one yields intervals for oddsratios. For the case of Dunnett-type contrasts, the calculated simultaneous confidence intervals are those described in Holford et al. (1989).

Specifying `method="GLM"` takes maximum likelihood estimates for the log-odds and their standard errors evaluated at the estimate.

Specifying `method="Woolf"` takes adds 0.5 to each cell count and computes point estimates and standard errors for these continuity corrected values. For the two-sample comparison this method is referred to as "adjusted Woolf" (Lawson, 2005). In this implementation, the lower bounds yielded by this method are additionally expanded to 0, if all values in the denominator are $x=n$ or all values in the numerator are $x=0$, and the upper bounds are expanded to Inf , if all values in the denominator are $x=0$ or all values in the numerator are $x=n$.

Note, that for the case of general contrasts, the methods are not described explicitly so far.

Value

A object of class "binomORci", a list containing:

<code>conf.int</code>	a matrix with 2 columns: lower and upper confidence bounds, and M rows
<code>alternative</code>	character string, as input
<code>conf.level</code>	single numeric value, as input
<code>estimate</code>	a matrix with 1 column: containing the estimates of the contrasts
<code>x</code>	the observed number of successes
<code>n</code>	the number of trials
<code>p</code>	the estimated proportions
<code>success</code>	a character string labelling the event considered as success
<code>names</code>	the group names
<code>method</code>	a character string, specifying the method of interval construction
<code>cmat</code>	the contrast matrix used

Author(s)

Frank Schaarschmidt, Daniel Gerhard

References

Holford, TR, Walter, SD and Dunnett, CW (1989). Simultaneous interval estimates of the odds ratio in studies with two or more comparisons. *Journal of Clinical Epidemiology* 42, 427-434.

See Also

Intervals for the risk difference [binomRDci](#), summary for odds ratio confidence intervals [summary.binomORci](#) plot for confidence intervals [plot.sci](#)

Examples

```

data(liarozole)

table(liarozole)

# Comparison to the control group "Placebo",
# which is the fourth group in alpha-numeric
# order:

ORlia<-binomORci(Improved ~ Treatment,
  data=liarozole, success="y", type="Dunnett", base=4)
ORlia
summary(ORlia)
plot(ORlia)

# if data are available as table:

tab<-table(liarozole)
tab
ORlia2<-binomORci(tab, success="y", type="Dunnett", base=4)
ORlia2

plot(ORlia2, lines=1, lineslty=3)

#####

# Performance for extreme cases

# method="GLM" (the default)

test1<-binomORci(x=c(0,1,5,20), n=c(20,20,20,20), names=c("A", "B", "C", "D"))
test1
plot(test1)

# adjusted Woolf interval

test2<-binomORci(x=c(0,1,5,20), n=c(20,20,20,20), names=c("A", "B", "C", "D"), method="Woolf")

```

```
test2
plot(test2)
```

binomRDci	<i>Simultaneous confidence intervals for contrasts of independent binomial proportions (in a oneway layout)</i>
-----------	---

Description

Simultaneous asymptotic CI for contrasts of binomial proportions, assuming that standard normal approximation holds. The contrasts can be interpreted as differences of (weighted averages) of proportions (risk ratios).

Usage

```
binomRDci(x,...)

## Default S3 method:
binomRDci(x, n, names=NULL,
  type="Dunnett", cmat=NULL, method="Wald",
  alternative="two.sided", conf.level=0.95,
  dist="MVN", ...)

## S3 method for class 'formula'
binomRDci(formula, data,
  type="Dunnett", cmat=NULL, method="Wald",
  alternative="two.sided", conf.level=0.95,
  dist="MVN",...)

## S3 method for class 'table'
binomRDci(x, type="Dunnett",
  cmat=NULL, method="Wald", alternative="two.sided",
  conf.level=0.95, dist="MVN",...)

## S3 method for class 'matrix'
binomRDci(x, type="Dunnett",
  cmat=NULL, method="Wald", alternative="two.sided",
  conf.level=0.95, dist="MVN",...)
```

Arguments

x	a numeric vector, giving the number of successes in I independent samples, or an object of class "table", representing the 2xk-table, or an object of class "matrix", representing the 2xk-table
---	--

n	a numeric vector, giving the number of trials (i.e. the sample size) in each of the I groups (only required if x is a numeric vector, ignored otherwise)
names	an optional character string, giving the names of the groups/ sample in x, n; if not specified the possible names of x are taken as group names (ignored if x is a table or matrix)
formula	a two-sided formula of the style 'response ~ treatment', where 'response' should be a categorical variable with two levels, while treatment should be a factor specifying the treatment levels
data	a data.frame, containing the variables specified in formula
type	a character string, giving the name of a contrast method, as defined in <code>contrMat(multcomp)</code> ; ignored if cmat is specified
cmat	a optional contrast matrix
method	a single character string, specifying the method for confidence interval construction; options are: "Wald", "ADD1", or "ADD2"
alternative	a single character string, one of "two.sided", "less", "greater"
conf.level	a single numeric value, simultaneous confidence level
dist	a character string, "MVN" invokes multiplicity adjustment via the multivariate normal distribution, "N" invokes use of quantiles of the univariate normal distribution
...	arguments to be passed to <code>binomest</code> , currently only success labelling the event which should be considered as success

Details

See the examples for different usages.

Value

A object of class "binomRDci", a list containing:

conf.int	a matrix with 2 columns: lower and upper confidence bounds, and M rows
alternative	character string, as input
conf.level	single numeric value, as input
quantile	the quantile used to construct the confidence intervals
estimate	a matrix with 1 column: containing the estimates of the contrasts
x	the observed number of successes in the treatment groups
n	the number of trials in the treatment groups
p	the estimated proportions in the treatment groups
success	a character string labelling the event considered as success
names	the group names
method	a character string, specifying the method of interval construction
cmat	the contrast matrix used

Note

Note, that all implemented methods are approximate only. The coverage probability of the intervals might seriously deviate from the nominal level for small sample sizes and extreme success probabilities. See the simulation results in Sill (2007) for details.

References

Schaarschmidt, F., Sill, M. and Hothorn, L.A. (2008): Approximate simultaneous confidence intervals for multiple contrasts of binomial proportions. *Biometrical Journal* 50, 782-792.

Background references:

The ideas underlying the "ADD1" and "ADD2" adjustment are described in:

Agresti, A. and Caffo, B.(2000): Simple and effective confidence intervals for proportions and differences of proportions result from adding two successes and two failures. *American Statistician* 54, p. 280-288.

And have been generalized for a single contrast of several proportions in:

Price, R.M. and Bonett, D.G. (2004): An improved confidence interval for a linear function of binomial proportions. *Computational Statistics and Data Analysis* 45, 449-456.

More detailed simulation results are available in:

Sill, M. (2007): Approximate simultaneous confidence intervals for multiple comparisons of binomial proportions. Master thesis, Institute of Biostatistics, Leibniz University Hannover.

See Also

[summary.binomRDci](#), [plot.sci](#)

Examples

```
#####
### Example 1 Tables 1,7,8 in Schaarschmidt et al. (2008): ###
#####

# Number of patients under observation:
n <- c(29, 24, 25, 24, 46)

# Number of patients with complete response:
cr <- c(7, 11, 10, 12, 21)

# (Optional) names for the treatments
dn <- c("0.3_1.0", "3", "10", "30", "90")

# Assume we aim to infer an increasing trend with increasing dosage,
# Using the changepoint contrasts (Table 7, Schaarschmidt et al., 2008)

contrMat(n=n, type="Changepoint")
```

```

# The results in Table 8 can be reproduced by calling:

binomRDci(n=n, x=cr, names=dn, alternative="greater",
  method="ADD2", type="Changepoint")

binomRDci(n=n, x=cr, names=dn, alternative="greater",
  method="ADD1", type="Changepoint")

binomRDci(n=n, x=cr, names=dn, alternative="greater",
  method="Wald", type="Changepoint")

#####

### Example 2, Tables 2,9,10 in Schaarschmidt et al. 2008 ###

#####

# Data (Table 2)

# animals under risk
n<-c(30,30,30,30)

# animals showing cancer
cancer<-c(20,14,27,19)

# short names for the treatments
trtn<-c("HFaFi","LFaFi","HFaNFi","LFaNFi")

# User-defined contrast matrix (Table 9),
# columns of the contrast matrix

cmat<-rbind(
  "Fiber - No Fiber"=c( 0.5, 0.5,-0.5,-0.5),
  "Low Fat - High Fat"=c(-0.5, 0.5,-0.5, 0.5),
  "Interaction Fat:Fiber"=c( 1, -1, -1, 1))

cmat

# The results in Table 10 can be reproduced by calling:

# simultaneous CI using the add-2 adjustment

sci<-binomRDci(x=cancer, n=n, names=trtn, method="ADD2",
  cmat=cmat, dist="MVN")

sci

# marginal CI using the basic Wald formula

ci<-binomRDci(x=cancer, n=n, names=trtn, method="Wald",
  cmat=cmat, dist="N")

```

```

ci

# check, whether the intended contrasts have been defined:

summary(sci)

# plot the result:

plot(sci, lines=0, lineslty=3)

#####

# In simple cases, counts of successes
# and number of trials can be just typed:

ntrials <- c(40,20,20,20)
xsuccesses <- c(1,2,2,4)
names(xsuccesses) <- LETTERS[1:4]
ex1D<-binomRDci(x=xsuccesses, n=ntrials, method="ADD1",
  type="Dunnett")
ex1D

ex1W<-binomRDci(x=xsuccesses, n=ntrials, method="ADD1",
  type="Williams", alternative="greater")
ex1W

# results can be plotted:
plot(ex1D, main="Comparisons to control group A", lines=0, linescol="red", lineslwd=2)

# summary gives a more detailed print out:
summary(ex1W)

# if data are represented as dichotomous variable
# in a data.frame one can make use of table:

#####

data(liarozole)

head(liarozole)

binomRDci(Improved ~ Treatment, data=liarozole,
  type="Tukey")

# here, it might be important to define which level of the
# variable 'Improved' is to be considered as success

binomRDci(Improved ~ Treatment, data=liarozole,

```

```

type="Dunnett", success="y", base=4)

# If data are available as a named kx2-contingency table:

tab<-table(liarozole)
tab

# Comparison to the control group "Placebo",
# which is the fourth group in alpha-numeric order:

CIs<-binomRDci(tab, type="Dunnett", success="y", base=4)

plot(CIs, lines=0)

```

binomRDtest	<i>Simultaneous test for contrasts of independent binomial proportions (in a oneway layout)</i>
-------------	---

Description

P-value of maximum test and adjusted p-values for M contrasts of I groups in a one-way layout. Tests are performed for contrasts of proportions, which can be interpreted as differences of (weighted averages of) proportions.

Usage

```

binomRDtest(x, ...)

## Default S3 method:
binomRDtest(x, n, names=NULL,
  type="Dunnett", cmat=NULL, method="Wald",
  alternative="two.sided", dist="MVN", ...)

## S3 method for class 'formula'
binomRDtest(formula, data,
  type="Dunnett", cmat=NULL, method="Wald",
  alternative="two.sided", dist="MVN", ...)

## S3 method for class 'table'
binomRDtest(x, type="Dunnett",
  cmat=NULL, method="Wald", alternative="two.sided",
  dist="MVN", ...)

## S3 method for class 'matrix'
binomRDtest(x, type="Dunnett",
  cmat=NULL, method="Wald", alternative="two.sided",
  dist="MVN", ...)

```

Arguments

x	a numeric vector, giving the number of successes in I independent samples, or an object of class "table", representing the 2xk-table, or an object of class "matrix", representing the 2xk-table
n	a numeric vector, giving the number of trials (i.e. the sample size) in each of the I groups
names	an optional character vector, giving the names of the groups in x, n; if not specified, possibly available names of x are taken as group names
formula	a two-sided formula of the style 'response ~ treatment', where 'response' should be a categorical variable with two levels, while treatment should be a factor specifying the treatment levels
data	a data.frame, containing the variables specified in formula
type	a character string specifying the contrast type
cmat	an optional user defined contrast matrix of dimension MxI
method	a single character string, specifying the method for adjustment, with options: "Wald" (Maximum likelihood estimators), "ADD1" (add1-adjustment on the raw proportion estimates) "ADD2" (add2-adjustment on proportion estimates following Agresti Caffo (2000))
alternative	a character string specifying the direction of the alternative hypothesis
dist	a character string, where "MVN" invokes the computation of p-values using the multivariate normal distribution, and "N" invokes use p-value computation using the univariate normal distribution
...	arguments to be passed to binomest , currently only success labelling the event which should be considered as success

Details

For usage, see the examples.

Value

An object of class "binomRDtest", a list containing:

teststat	a numeric vector of teststatistics of length M
pval	a single numeric p-value, the p-value of the maximum test (minimum p-value)
p.val.adj	a vector of length M, the adjusted p-values of the single contrasts
dist	character string indicating whether the multivariate normal or normal distribution was used for computation of p-values
alternative	a single character vector, as the input
x	the observed number of successes in the treatment groups
n	the number of trials in the treatment groups
p	the estimated proportions in the treatment groups
success	a character string labelling the event considered as success
method	as input, a character string
cmat	used contrast matrix

Note

Note, that all implemented methods are approximate only. The size of the test might seriously deviate from the nominal level for small sample sizes and extreme success probabilities. See the simulation results in Sill (2007) for details.

References

Statistical procedures and characterization of coverage probabilities are described in: Sill, M. (2007): Approximate simultaneous confidence intervals for multiple comparisons of binomial proportions. Master thesis, Institute of Biostatistics, Leibniz University Hannover.

See Also

[summary.binomRDtest](#)

Examples

```
ntrials <- c(40,20,20,20)
xsucceses <- c(1,2,2,4)
names(xsucceses) <- LETTERS[1:4]
binomRDtest(x=xsucceses, n=ntrials, method="ADD1",
  type="Dunnett")

binomRDtest(x=xsucceses, n=ntrials, method="ADD1",
  type="Williams", alternative="greater")

binomRDtest(x=xsucceses, n=ntrials, method="ADD2",
  type="Williams", alternative="greater")
```

binomRRci	<i>Simultaneous confidence intervals for ratios of independent binomial proportions</i>
-----------	---

Description

Simultaneous asymptotic CI for contrasts of binomial proportions, assuming that standard normal approximation holds on the log scale. Confidence intervals for ratios of (weighted geometric means) of proportions are calculated based on differences of log-proportions, and normal approximation on the log-scale.

Usage

```
binomRRci(x,...)

## Default S3 method:
binomRRci(x, n, names=NULL,
```

```

type="Dunnett", cmat=NULL,
alternative="two.sided", conf.level=0.95,
dist="MVN", ...)

## S3 method for class 'formula'
binomRRci(formula, data,
  type="Dunnett", cmat=NULL,
  alternative="two.sided", conf.level=0.95,
  dist="MVN",...)

## S3 method for class 'table'
binomRRci(x, type="Dunnett",
  cmat=NULL, alternative="two.sided",
  conf.level=0.95, dist="MVN",...)

## S3 method for class 'matrix'
binomRRci(x, type="Dunnett",
  cmat=NULL, alternative="two.sided",
  conf.level=0.95, dist="MVN",...)

```

Arguments

x	a numeric vector, giving the number of successes in I independent samples, or an object of class "table", representing the 2xk-table, or an object of class "matrix", representing the 2xk-table
n	a numeric vector, giving the number of trials (i.e. the sample size) in each of the I groups (only required if x is a numeric vector, ignored otherwise)
names	an optional character string, giving the names of the groups/ sample in x, n; if not specified the possible names of x are taken as group names (ignored if x is a table or matrix)
formula	a two-sided formula of the style 'response ~ treatment', where 'response' should be a categorical variable with two levels, while treatment should be a factor specifying the treatment levels
data	a data.frame, containing the variables specified in formula
type	a character string, giving the name of a contrast method, as defined in <code>contrMat(multcomp)</code> ; ignored if cmat is specified
cmat	a optional contrast matrix
alternative	a single character string, one of "two.sided", "less", "greater"
conf.level	a single numeric value, simultaneous confidence level
dist	a character string, "MVN" invokes multiplicity adjustment via the multivariate normal distribution, "N" invokes use of quantiles of the univariate normal distribution
...	arguments to be passed to <code>binomest</code> , currently only success labelling the event which should be considered as success

Details

The interval for the ratio of two independent proportions, described in section "Crude Methods using first-order variance estimation" in Gart and Nam (1988) are extended to multiple contrasts. Confidence intervals are constructed based on contrasts for differences of $lp = \log(x+0.5)/(n+0.5)$, using quantiles of the multivariate normal or normal approximation. Applying the exponential functions to the bounds results in intervals for the risk ratio. In case that 0 occur in both, the numerator and denominator of the ratio, the interval is expanded to $[0, \text{Inf}]$, in case that only 0s numerator go to the numerator, the lower bound is expanded to 0, in case that only 0s go to the denominator, the upper bound is expanded to Inf .

See the examples for different usages.

Value

A object of class "binomRDci", a list containing:

conf.int	a matrix with 2 columns: lower and upper confidence bounds, and M rows
alternative	character string, as input
conf.level	single numeric value, as input
quantile	the quantile used to construct the confidence intervals
estimate	a matrix with 1 column: containing the estimates of the contrasts
x	the observed number of successes in the treatment groups
n	the number of trials in the treatment groups
p	the estimated proportions in the treatment groups
success	a character string labelling the event considered as success
names	the group names
method	a character string, specifying the method of interval construction
cmat	the contrast matrix used

Note

Note, that all implemented methods are approximate only. The coverage probability of the intervals might seriously deviate from the nominal level for small sample sizes and extreme success probabilities.

References

Gart, JJ and Nam, J-m (1988): Approximate interval estimation of the ratio of binomial parameters: a review and corrections for skewness. *Biometrics* 44, 323-338.

See Also

[summary.binomRDci](#) for the risk difference, [summary.binomORci](#) for the odds ratio, [plot.sci](#) for plotting

Examples

```
# In simple cases, counts of successes
# and number of trials can be just typed:

ntrials <- c(40,20,20,20)
xsuccesses <- c(1,2,2,4)
names(xsuccesses) <- LETTERS[1:4]
ex1D<-binomRRci(x=xsuccesses, n=ntrials,
  type="Dunnett")
ex1D

ex1W<-binomRRci(x=xsuccesses, n=ntrials,
  type="Williams", alternative="greater")
ex1W

# results can be plotted:
plot(ex1D, main="Comparisons to control group A")

# summary gives a more detailed print out:
summary(ex1W)

# if data are represented as dichotomous variable
# in a data.frame one can make use of table:

data(liarozole)

head(liarozole)

# here, it might be important to define which level of the
# variable 'Improved' is to be considered as success

binomRRci(Improved ~ Treatment, data=liarozole,
  type="Dunnett", success="y", base=4, alternative="greater")

# If data are available as a named kx2-contingency table:

tab<-table(liarozole)
tab

binomRRci(tab, type="Dunnett", success="y", base=4, alternative="greater")

# Performance for extreme cases:

binomRRci(x=c(0,0,20,5),n=c(20,20,20,20),names=c("A","B","C","D"),
  type="Dunnett", alternative="greater")
```

bronch

*Rodent bronchial carcinoma data***Description**

In a long term rodent carcinogenicity study on female B6C3F1 mice, the effect of vinylcyclohexene diepoxide on the incidence of murine alveolar/bronchiolar tumors was assessed. The mice were exposed to 0 mg/ml, (group 0), 25 mg/ml (group 1), 50 mg/ml (group 2), and 100 (group 3) mg/ml, with 50 animals per group.

Usage

```
data(bronch)
```

Format

A data frame with 200 observations on the following 3 variables.

group a factor with levels 0, 1, 2, 3, labelling the control and the three dose groups

Y a logical vector, indicating whether a tumour was present at time of death (if TRUE), or not (if FALSE)

time a numeric vector, the time of death, counted in days? from begin of the study

Details

Not yet checked for consistency with the source!

Source

Piegorsch WW and Bailer AJ (1997): Statistics for environmental biology and toxicology. Chapman and Hall, London. Table 6.5, page 238.

References

Portier cJ and Bailer AJ (1989): Testing for increased carcinogenicity using survival-adjusted quantal response tests. *Fundamental and Applied Toxicology* 12, 731.

Examples

```
data(bronch)
# raw tumour counts:

table(bronch[c("group", "Y")])

# groupwise times of death:

boxplot(time ~ group, data=bronch, horizontal=TRUE)
```

```
# Using poly3estf, we can produce the
# summary statistics as presented in
# Table 6.6, page 239, of Piegorsch and Bailer (1997):

poly3estf(status=bronch$Y, time=bronch$time, f=bronch$group)
```

censsample

Random data for Poly-k

Description

Random numbers from two independent Weibull distributions for Mortality and tumour induction.

Usage

```
censsample(n, scale.m, shape.m,
           scale.t, shape.t = 3, tmax)
```

Arguments

n	a single numeric value, the number of individuals
scale.m	a single numeric value, scale parameter of the Weibull distribution for mortality
shape.m	a single numeric value, shape parameter of the Weibull distribution for mortality
scale.t	a single numeric value, scale parameter of the Weibull distribution for tumour induction
shape.t	a single numeric value, shape parameter of the Weibull distribution for tumour induction
tmax	a single numeric value, maximum time in the trial

Value

A data.frame with columns

time	a numeric vector of length n, the time of death of an individual
status	a logical vector of length n, the tumour status at time of death (TRUE: tumour present, FALSE: no tumour present)
T.t	time of tumour induction (unobservable)
T.m	time of death
tmax	maximum time of death

censsamplef	<i>Random data for Poly-k</i>
-------------	-------------------------------

Description

Random data for Poly-k for a one-way layout, with I groups.

Usage

```
censsamplef(n, scale.m, shape.m, scale.t, shape.t = 3, tmax)
```

Arguments

n	a numeric vector, the numbers of individuals of length I
scale.m	a numeric vector, scale parameters of the Weibull distribution for mortality
shape.m	a numeric vector, shape parameters of the Weibull distribution for mortality
scale.t	a numeric vector, scale parameters of the Weibull distribution for tumour induction
shape.t	a numeric vector, shape parameters of the Weibull distribution for tumour induction
tmax	a single numeric value, maximum time in the trial

Value

A data.frame with columns

time	a numeric vector of length n, the time of death of an individual
status	a logical vector of length n, the tumour status at time of death (TRUE: tumour present, FALSE: no tumour present)
T.t	time of tumour induction (unobservable)
T.m	time of death
tmax	maximum time of death
f	a factor of containing an appropriate grouping variable

corrMatgen *A function to calculate the correlation matrix*

Description

Correlation matrix for test statistics and confidence intervals assuming multivariate standard normal distribution

Usage

```
corrMatgen(CM, varp)
```

Arguments

CM	a matrix of contrast coefficients, dimension MxI, where M=number of contrasts, and I=number of groups in a oneway layout
varp	a numeric vector of groupwise variance estimates (length = I)

Value

A matrix of dimension MxM.

References

For correlation of contrasts of binomial proportion, see: Bretz F, Hothorn L.: Detecting dose-response using contrasts: asymptotic power and sample size determination for binomial data. *Statistics in Medicine* 2002; 21: 3325-3335.

estShannon *Estimate the Shannon Index*

Description

Calculates estimates of the Shannon-Wiener from count data.

Usage

```
estShannon(x, Nspec = NULL)
```

Arguments

x	a integer(numeric) vector of species counts
Nspec	a single integer value, fixing the number of species to a certain value

Value

A list, containing the elements:

estimate	a single numeric value, the estimate with bias correction according to Fritsch and Hsu (1999)
estraw	a single numeric value, the raw estimate
varest	a single numeric value, the variance estimate according to Fritsch and Hsu (1999)

References

Fritsch, KS, and Hsu, JC (1999): Multiple Comparison of Entropies with Application to Dinosaur Biodiversity. *Biometrics* 55, 1300-1305.

See Also

[estSimpsonf](#) for estimating Shannon indices pooled over several samples, grouped by a factor

estShannonf	<i>Estimate the Shannon-Wiener index</i>
-------------	--

Description

Calculate estimates of the Shannon-Wiener index after pooling over several samples, grouped by a factor variable.

Usage

```
estShannonf(X, f)
```

Arguments

X	a data.frame of dimension n times p with integer entries, where n is the number of samples and p is the number of species
f	a factor variable of length n, grouping the observations in X

Details

The function splits X according to the levels of the grouping variable f, builds the sum over each column and calculates the Shannon index over the resulting counts.

Value

A list, containing the elements:

estimae	a named numeric vector, the groupwise Shannon indices with bias correction according to Fritsch and Hsu (1999)
estraw	a named numeric vector, the groupwise Shannon indices, without bias correction
varest	a named numeric vector, the groupwise variance estimates of the Shannon indices
table	a matrix, giving the summarized counts of the groups in the rows

References

Fritsch, KS, and Hsu, JC (1999): Multiple Comparison of Entropies with Application to Dinosaur Biodiversity. *Biometrics* 55, 1300-1305.

Examples

```
data(HCD)
HCD

# Groupwise point estimates:
est<-estShannonf(X=HCD[,-1], f=HCD[,1])

est
```

estSimpson	<i>Simpson index</i>
------------	----------------------

Description

Calculates the estimate of the Simpson index from a vector of counts.

Usage

```
estSimpson(x)
```

Arguments

x a integer (numeric) vector, giving the counts of p species in a community

`estSimpsonf`*Estimate the Simpson index from several samples*

Description

Calculate estimates of the Simpson index after pooling over several samples, grouped by a factor variable.

Usage

```
estSimpsonf(X, f)
```

Arguments

<code>X</code>	a data.frame of dimension n times p with integer entries, where n is the number of samples and p is the number of species
<code>f</code>	a factor variable of length n, grouping the observations in X

Details

The function splits X according to the levels of the grouping variable f, builds the sum over each column and calculates the Shannon index over the resulting counts.

Value

A list containing the items:

<code>estimate</code>	the groupwise point estimates of the Simpson index
<code>varest</code>	the groupwise variance estimates of the Simpson index
<code>table</code>	a matrix of counts, containing the summed observations for each level of f in its rows

References

Rogers, JA and Hsu, JC (2001): Multiple Comparisons of Biodiversity. Biometrical Journal 43, 617-625.

See Also

[estShannonf](#)

Examples

```

# Here, the estimates for the Hell Creek Dinosaur
# example are compared to the estimates in
# Tables 2 and 3 of Rogers and Hsu (2001).

data(HCD)
HCD

# Groupwise point estimates:

est<-estSimpsonf(X=HCD[,-1], f=HCD[,1])

est

# Table 2:

cmat<-rbind(
"lower-middle"=c(1,-1,0),
"lower-upper"=c(1, 0,-1),
"middle-upper"=c(0,1,-1))

# the point estimates:

# cmat %*% est$estimate
crossprod(t(cmat), est$estimate)

# the standard errors:
# sqrt(diag(cmat %*% diag(est$varest) %*% t(cmat)))

sqrt(diag(crossprod(t(cmat), crossprod(diag(est$varest), t(cmat)) ) ) ) )

# Table 3:

cmat<-rbind(
"middle-lower"=c(-1,1,0),
"upper-lower"=c(-1,0,1))

# cmat %*% est$estimate
crossprod(t(cmat), est$estimate)

# sqrt(diag(cmat %*% diag(est$varest) %*% t(cmat)))

sqrt(diag(crossprod(t(cmat), crossprod(diag(est$varest), t(cmat)) ) ) ) )

# Note, that the point estimates are exactly
# the same as in Rogers and Hsu (2001),

```

```
# but the variance estimates are not, whenever  
# the Upper group is involved.
```

HCD

Hell Creek Dinosaur Data

Description

Counts of dinosaur families found in three stratigraphic levels of the Cretaceous period in the Hell Creek formation in North Dakota. The eight families are the Ceratopsidae (Ce), Hadrosauridae (Ha), Hypsilophodontidae (Hy), Pachycephalosauridae (Pa), Tyrannosauridae (Ty), Ornithomimidae (Or), Sauronithoididae (Sa) and Dromiaeosauridae (Dr).

Usage

```
data(HCD)
```

Format

A data frame with 3 observations on the following 9 variables.

Level a factor with levels Lower, Middle, Upper, specifying the stratigraphic levels

Cr a numeric vector, counts of the Ceratopsidae

Ha a numeric vector, counts of the Hadrosauridae

Hy a numeric vector, counts of the Hypsilophodontidae

Pa a numeric vector, counts of the Pachycephalosauridae

Ty a numeric vector, counts of the Tyrannosauridae

Or a numeric vector, counts of the Ornithomimidae

Sa a numeric vector, counts of the Sauronithoididae

Dr a numeric vector, counts of the Dromiaeosauridae

Source

Table 1 in: Rogers, JA and Hsu, JC (2001): Multiple Comparisons of Biodiversity. Biometrical Journal 43, 617-625.

References

Sheehan, P.M., et al. (1991): Sudden extinction of the Dinosaurs: Latest Cretaceous, Upper Great Plains, U.S.A. Science 254, 835-839.

Examples

```
data(HCD)
str(HCD)
HCD

mat<-as.matrix(HCD[,-c(1)])

rownames(mat)<-HCD$Level

mosaicplot(mat, las=1)

estSimpsonf(X=HCD[,-c(1)], f=HCD$Level)

estShannonf(X=HCD[,-c(1)], f=HCD$Level)
```

liarozole

Marked improvement of psoriasis after application of liarozole

Description

In a placebo controlled clinical trial, patients with psoriasis were randomly assigned to a placebo group and three dose groups (50 mg, 75 mg, and 150 mg). Variable of primary interest was the proportion of patients with marked improvement of psoriasis. This data.frame mimics how raw data could have been represented in a larger data frame.

Usage

```
data(liarozole)
```

Format

A data frame with 137 observations on the following 2 variables.

Improved a factor with levels n, y, for "no" and "yes"

Treatment a factor with levels Dose150, Dose50, Dose75, Placebo

Details

For illustrative purpose only. Number of successes recalculated from proportions presented in the publication, while the number of patients in group Dose50 was not exactly clear.

Source

Berth-Jones J, Todd G, Hutchinson PE, Thestrup-Pedersen K, Vanhoutte FP: Treatment of psoriasis with oral liarozole: a dose-ranging study. *British Journal of Dermatology* 2000; 143: 1170-1176.

Examples

```

data(liarozole)
head(liarozole)
# create a contingency table:

table(liarozole)

# the order of the groups is alpha-numeric,
# and "y" for success is of higher order than
# to change the order:

liarozole$Treatment<-factor(liarozole$Treatment,
  levels=c("Placebo", "Dose50", "Dose75", "Dose150"))

liarozole$Improved<-factor(liarozole$Improved,
  levels=c("y", "n"))

tab<-table(liarozole)
tab

# Approximate simultaneous confidence intervals
# for the differences pDose-pPlacebo:

LCI<-binomRDci(tab, type="Dunnett",
  alternative="greater", method="ADD1")

LCI

plot(LCI, main="Proportion of patients
  with marked improvement")

# Perform a test on increasing trend
# vs. the placebo group:

Ltest<-binomRDtest(tab, type="Williams",
  alternative="greater", method="ADD1")

summary(Ltest)

```

Indest

*Point and variance estimates for expected and log(expected values)
of lognormal samples*

Description

Compute the ML estimates of the expected values and the log of the expected values given several samples of lognormal data, separated by a factor variable. For internal use in `Indci` and `lnrci`.

Usage

```
lndest(x, f)
lnrest(x, f)
```

Arguments

x a numeric vector
f a factor variable (with at least two levels)

Value

A list with elements

estimate a numeric vector, the point estimates for each factor level
varest a numeric vector, the variance estimates (evaluated at the point estimates) for each factor level
n a numeric vector, the sample size of each factor level

References

Chen, Y-H, Zhou, X-H (2006). Interval estimates for the ratio and the difference of two log-normal means. *Statistics in Medicine* 25, 4099-4113.

Inrci	<i>Simultaneous confidence intervals for expected values assuming log-normal distribution.</i>
-------	--

Description

Simultaneous confidence intervals for multiple contrasts of expected values of several (K) groups in a one-way layout, assuming lognormal distribution of the response. Multiple ratios of (weighted geometric means of) expected values can be estimated as well as multiple differences of (weighted arithmetic means of) the expected values in the K groups can be estimated. For both, ratios and differences, a method based on asymptotic normality (not recommended) and a method based on generalized pivotal quantities are available.

Usage

```
lnrci(x, f, type = "Dunnett", cmat = NULL, alternative = c("two.sided", "less", "greater"),
      conf.level = 0.95, method = c("GPQ", "AN"), ...)
lndci(x, f, type = "Dunnett", cmat = NULL, alternative = c("two.sided", "less", "greater"),
      conf.level = 0.95, method = c("GPQ", "AN"), ...)
```

Arguments

<code>x</code>	a numeric vector, the response variable, assumed to be lognormally distributed, should contain only positive values
<code>f</code>	a factor variable, of the same length as <code>x</code> , grouping the observations in <code>x</code>
<code>type</code>	a single character string, naming a contrast type, see contrMat for the options; this argument is ignored if a contrast matrix is specified in <code>cmat</code>
<code>cmat</code>	a matrix with numeric entries, containing contrast coefficients; if there are <code>K</code> levels in <code>f</code> , the matrix should have <code>k</code> columns
<code>alternative</code>	a single character string, with options "two.sided" for two-sided confidence intervals, "less" for upper bounds only and "greater" for lower bounds only
<code>conf.level</code>	a single numeric value between 0 and 1
<code>method</code>	a single character string, naming the method by which to compute the confidence intervals; options are "GPQ" for a method based on generalized pivotal quantities and "AN" for the asymptotic normal method (not recommended).
<code>...</code>	further arguments to be passed to the internal methods, in particular: Argument <code>B</code> specifies the number of samples to be drawn if method "GPQ" (defaults to 10000), it is ignored if method="AN". Second, <code>dist</code> must be a single character string as in Waldci , invoking the use of multivariate normal quantiles <code>dist="MVN"</code> or standard normal quantiles <code>dist="N"</code> when method="AN"; it is ignored if method="GPQ".

Details

This is a testversion. In a setting with `K` treatment groups, assuming a completely randomized design and lognormal distribution of the response variable, multiple (`M`) ratios or differences among expected values of the `K` treatment groups may be of interest. The ratios or differences of interest can be specified via a character string (in argument `type`) or via an `M x K` matrix in argument `cmat`. Intervals for ratios can be computed (via linear contrasts on the log scale) in function `Inrci`, differences can be computed in function `Indci`. By default, the confidence intervals are adjusted for multiple comparisons. The asymptotic normal methods, rely on maximum likelihood estimates for the expected values and the corresponding variance estimates (as given in Chen and Zhou, 2006) and adjust for multiplicity via critical value from the multivariate normal distribution (correlation structure with plug-in of variance estimates). The generalized pivotal quantity methods rely on simulating the distribution of the parameters (number of samples `B=10000` by default, Krishnamoorthy and Mathew, 2003; Chen and Zhou, 2006) and compute simultaneous intervals from this sample using the function `SCSrank`.

Value

A list with elements

<code>estimate</code>	a column vector, containing the point estimates of the contrasts
<code>conf.int</code>	a <code>Mx2</code> matrix of confidence bounds, if <code>M</code> comparisons among the <code>K</code> samples are invoked
<code>alternative</code>	a character string, as input
<code>conf.level</code>	a numeric value, as input

Author(s)

Frank Schaarschmidt

References

Chen, Y-H, Zhou, X-H (2006). Interval estimates for the ratio and the difference of two log-normal means. *Statistics in Medicine* 25, 4099-4113.

Krishnamoorthy K, Mathew T (2003). Inferences on the means of lognormal distributions using generalized p-values and generalized confidence intervals. *Journal of Statistical Planning and Inference* 115, 103-121.

Examples

```
x <- rlnorm(n=100, meanlog=rep(c(0,0.1,1,1), each=25), sdlog=0.5)
f <- as.factor(rep(LETTERS[1:4], each=25))
boxplot(x~f)

lndci(x=x, f=f, type="Dunnett", method="GPQ", B=40000)
lndci(x=x, f=f, type="Dunnett", method="AN")

lnrci(x=x, f=f, type="Tukey", method="GPQ", B=40000)
lnrci(x=x, f=f, type="Tukey", method="AN")
```

methyl

NTP bioassay data: effect of methyleugenol on skin fibroma

Description

NTP bioassay of methyleugenol: 200 male rats were randomly assigned to 4 treatment groups with balanced sample size 50. Individuals in treatment group 0, 1, 2, and 3 received doses of 0, 37, 75, and 150 mg methyleugenol per kg body weight, respectively. The response variable tumour is the presence of skin fibroma at time of death. The variable death gives individual time of death, with a final sacrifice of surviving animals at 730 days after begin of the assay.

Usage

```
data(methyl)
```

Format

A data frame with 200 observations on the following 3 variables.

group a factor with levels 0, 1, 2, 3, specifying dose groups 0, 37, 75, and 150 mg/kg, respectively

tumour a numeric vector, specifying whether a tumour was present at time of death

death a numeric vector, specifying the time of death

Source

National toxicology program (2000).

References

SD Peddada, GE Dinse, JK Haseman (2005): A survival-adjusted quantal response test for comparing tumour incidence rates. *Applied Statistics* 54, 51-61.

Examples

```
data(methyl)
# raw tumour proportions:
table(methyl[c("group", "tumour")])

# time of death:
boxplot(death~group, data=methyl, horizontal=TRUE)
```

mosaicdiv

Mosaicplot for the data in Shannonci and Simpsonci

Description

Create a mosaicplot from objects of class Shannonci or Simpsonci

Usage

```
mosaicdiv(x, decreasing = NULL, ...)
```

Arguments

x	an object of class "Simpsonci" or "Shannonci" as can be obtained from calling Simpsonci or Shannonci
decreasing	a single logical value, indicating whether the species should be plotted in the current order (if decreasing=NULL), in decreasing order (if decreasing=TRUE), or in increasing order (if decreasing=FALSE)
...	further arguments to be passed to mosaicplot, see ?mosaicplot and ?par for details

Details

This function uses the counts in `[["sample.estimate"]][["table"]]` to produce a mosaicplot.

Value

A plot.

Examples

```
data(HCD)

HCDFam <- HCD[,-1]

SCI<-Simpsonci(X=HCDFam, f=HCD[,1])

mosaicdiv(SCI, decreasing=TRUE, col=rainbow(n=8))
```

plot.sci

Plot confidence intervals

Description

Function for convenient plotting of confidence intervals.

Usage

```
## S3 method for class 'sci'
plot(x, ...)
## S3 method for class 'binomRDci'
plot(x, ...)
## S3 method for class 'binomORci'
plot(x, ...)
## S3 method for class 'binomRRci'
plot(x, ...)
## S3 method for class 'poly3ci'
plot(x, ...)
## S3 method for class 'Shannonci'
plot(x, ...)
## S3 method for class 'Simpsonci'
plot(x, ...)
```

Arguments

x an object of class "binomRDci", "binomORci", "binomRRci", "poly3ci", "sci"
... further arguments as described in [plotCI](#)

Details

Extracts some values and calls [plotCI](#).

Value

A plot.

plotCI

Plot confidence intervals

Description

A function for convenient plotting of confidence intervals.

Usage

```
## Default S3 method:
plotCI(x, ...)
## S3 method for class 'sci'
plotCI(x, ...)
## S3 method for class 'sci.ratio'
plotCI(x, ...)
## S3 method for class 'confint.glht'
plotCI(x, ...)
```

Arguments

x	An object of class "sci", "sci.ratio" or "conf.int.glht" or a list with elements estimate, containing a numeric vector, conf.int, containing a matrix with two columns, giving the lower and upper bounds, and a string alternative, one of "two.sided", "less", "greater"
...	additional arguments to be passed to plotCII and plot , see plotCII for details

Details

Plots the estimates, upper and lower limits using points and segments. The names of estimate are passed as labels of the confidence intervals. If infinite bounds occur, the plot region is limited by the most extreme non infinite bound or estimate.

Value

A plot.

See Also

Internally, the function [plotCII](#) is used.

Examples

```
x=c(8,9,9,18,39,44)
n=c(2000,2000,2000,2000,2000,2000)

x<-binomORci(x=x, n=n, names=c("0","120","240","480","600","720"))
```

```
plotCI(x, lines=1)
```

plotCII

Plot confidence intervals

Description

A function for convenient plotting of confidence intervals.

Usage

```
plotCII(estimate, lower = NULL, upper = NULL,
        alternative = c("two.sided", "less", "greater"),
        lines = NULL, lineslty = 2, lineslwd = 1,
        linescol = "black", CIvert = FALSE, CIlty = 1,
        CIlwd = 1, ICex = 1, Cicol = "black", CIlength=NULL,
        HL = TRUE, ...)
```

Arguments

estimate	a (named) numeric vector, the names of the elements are taken as labels of the CI
lower	an optional numeric vector, of the same length as estimate
upper	an optional numeric vector, of the same length as estimate
alternative	a single character string, one of "two.sided", "less", "greater"
lines	an optional numeric (vector) giving the position(s) of line(s) to draw into the plots orthogonal to the confidence intervals
lineslty	possible a vector of line type of the lines, see the options for lty in par
lineslwd	possible a vector of line width of the lines, see the options for lwd in par
linescol	possible a vector of line colors of the lines
CIvert	logical indicating, whether confidence intervals shall be drawn horizontal (default), or vertical (if set to TRUE)
CIlty	single value, to specify the line type used for the CI, see options for lty in ?par
CIlwd	single value, to specify the width type used for the CI, see options for lty in ?par
ICex	single value, to specify the extension of sympols in the plot, see options for cex in ?par
Cicol	single value, to specify the color used for the CI
CIlength	single numeric value, to be passed to the argument length in function arrows ; to specify the lengths of the CI bounds (in inches); defaults to 0.08 and 0.05 for less than 25 and more than 25 CIs, respectively

HL a logical, if TRUE (default), plot margins of the are adjusted depending on the length of the names by appropriate calls to `par`; this might be incompatible with combining the plot with others in the same device. If set to FALSE, its up to the user to choose appropriate plot margins by calling to `par`.

... further arguments to be passed to `plot`

See Also

[plotCI](#), for more convenient methods

Examples

```
est<-c(1,2,3)
names(est)<-c("A", "B", "C")
lw=c(0,1,2)
up=c(2,3,4)

plotCII(estimate=est, lower=lw, upper=up)

plotCII(estimate=est, lower=lw, upper=up,
lines=c(-1,0,1),
lineslty=c(3,1,3),
lineslwd=c(1,2,1))

#####

names(est)<-c("very long names",
"e v e n l o n g e r n a m e s", "C")

plotCII(estimate=est, lower=lw, upper=up,
Cicol=c("black","green","red"),
HL=TRUE
)

#####

names(est)<-c("very long names",
"e v e n l o n g e r n a m e s", "C")

plotCII(estimate=est, lower=lw, upper=up,
Cicol=c("black","green","red"),
HL=TRUE
)

op<-par(no.readonly = TRUE)

layout(matrix(1:2, ncol=1))
```

```

par(mar=c(5,14,3,1))

plotCII(estimate=est, lower=lw, upper=up,
  main="Lala 1",
  Cicol=c("black","green","red"),
  lines=-1,
  HL=FALSE
)

plotCII(estimate=est, lower=lw, upper=up,
  main="Lala 2",
  Cicol=c("black","green","red"),
  lines=c(0,1),
  HL=FALSE
)

par(op)

```

poly3ci

Simultaneous confidence intervals for contrasts of poly-3-adjusted tumour rates

Description

Function to calculate simultaneous confidence intervals for several contrasts of poly-3-adjusted tumour rates in a oneway layout. Assuming a data situation as in Peddada(2005) or Bailer and Portier (1988). Simultaneous asymptotic CI for contrasts of tumour rates, assuming that standard normal approximation holds.

Usage

```

poly3ci(time, status, f, type = "Dunnett",
  cmat = NULL, method = "BP", alternative = "two.sided",
  conf.level = 0.95, dist="MVN", k=3, ...)

```

Arguments

time	a numeric vector of times of death of the individuals
status	a logical (or numeric, consisting of 0,1 only) vector giving the tumour status at time of death of each individual, where TRUE (1) = tumour present, FALSE (0) = no tumour present
f	a factor, giving the classification variable
type	a character string, giving the name of a contrast method, as defined in <code>contrMat(multcomp)</code>
cmat	a optional contrast matrix

method	a single character string, specifying the method for adjustment, with options: "BP" (Bailer Portier: assuming poly-3-adjusted rates are binomial variables), "BW" (Bieler, Williams: delta method as in Bieler and Williams (1993)) "ADD1" (as Bailer Portier, including an add1-adjustment on the raw tumour rates) "ADD2" (as Bailer Portier, including an add2-adjustment on the raw tumour rates following Agresti and Caffo (2000) for binomials)
alternative	a single character string
conf.level	a single numeric value, simultaneous confidence level
dist	a character string, "MVN" invokes multiplicity adjustment via the multivariate normal distribution, "N" invokes use of quantiles of the univariate normal distribution
k	the exponent to calculate survival adjusted proportions, default is k=3
...	further arguments to be passed; currently only base, to be passed to <code>contrMat</code> to choose the control group with <code>type="Dunnett"</code>

Value

A object of class "poly3ci", a list containing:

conf.int	a matrix with 2 columns: lower and upper confidence bounds, and M rows
alternative	character string, as input
conf.level	single numeric value, as input
quantile	the quantile used to construct the CIs
estimate	a numeric vector with the point estimates of the contrasts
time	as input
status	as input
f	as input
method	as input
cmat	as input, with <code>colnames=</code> factor levels of f
sample.est	a list containing sample estimates

Note

Please note that all methods here described are only approximative, and might violate the nominal level in certain situations. Please note further that appropriateness of the point estimates, and consequently of tests and confidence intervals is based on the assumptions in Bailer and Portier (1988), which might be a matter of controversies.

Author(s)

Frank Schaarschmidt

References

The implemented methodology is described in:

Schaarschmidt, F., Sill, M., and Hothorn, L.A. (2008): Approximate Simultaneous confidence intervals for multiple contrasts of binomial proportions. *Biometrical Journal* 50, 782-792.

Background references are:

Assumption for poly-3-adjustment: Bailer, J.A. and Portier, C.J. (1988): Effects of treatment-induced mortality and tumor-induced mortality on tests for carcinogenicity in small samples. *Biometrics* 44, 417-431.

Peddada, S.D., Dinse, G.E., and Haseman, J.K. (2005): A survival-adjusted quantal response test for comparing tumor incidence rates. *Applied Statistics* 54, 51-61.

Bieler, G.S. and Williams, R.L. (1993): Ratio estimates, the Delta Method, and quantal response tests for increased carcinogenicity. *Biometrics* 49, 793-801.

Statistical procedures and characterization of the coverage probabilities are described in: Sill, M. (2007): Approximate simultaneous confidence intervals for multiple comparisons of binomial proportions. Master thesis, Institute of Biostatistics, Leibniz University Hannover.

Examples

```
#####
### Methyleugenol example in Schaarschmidt et al. (2008) ###
#####
# load the data:

data(methyl)

# The results in Table 5 (Schaarschmidt et al. 2008) can be
# reproduced by calling:

methylW<-poly3ci(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Williams", method = "ADD1", alternative="greater" )

methylW

methylWT<-poly3test(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Williams", method = "ADD1", alternative="greater" )

methylWT

plot(methylW, main="Simultaneous CI for \n Poly-3-adjusted tumour rates")

# The results in Table 6 can be reproduced by calling:
```

```

methylD<-poly3ci(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Dunnett", method = "ADD1", alternative="greater" )

methylD

methylDT<-poly3test(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Dunnett", method = "ADD1", alternative="greater" )

methylDT

plot(methylD, main="Simultaneous CI for Poly-3-adjusted tumour rates", cex.main=0.7)

#####

# unadjusted CI

methylD1<-poly3ci(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Dunnett", method = "ADD1", dist="N" )

methylD1

plot(methylD1, main="Local CI for Poly-3-adjusted tumour rates")

```

poly3est

Only for internal use.

Description

Poly-3- adjusted point and variance estimates for long term carcinogenicity data

Usage

```
poly3est(time, status, tmax, method = "BP", k=NULL)
```

Arguments

time	a numeric vector of times of death of the individuals
status	a logical (or numeric, consisting of 0,1 only) vector giving the tumour status at time of death of each individual, where TRUE (1) = tumour present, FALSE (0) = no tumour present
tmax	a single numeric vector, the final time of the trial
method	a single character string, specifying the method for adjustment, with options: "BP" (Bailer Portier: assuming poly-3-adjusted rates are binomial variables), "BW" (Bieler, Williams: delta method as in Bieler-Williams (1993)) "ADD1"

(as Bailer Portier, including an add1-adjustment on the raw tumour rates) "ADD2"
(as Bailer Portier, including an add2-adjustment on the raw tumour rates following Agresti Caffo (2000) for binomials)

k a single numeric value, the exponent to calculate survival adjusted proportions according to Bailer and Portier (1988), defaults to 3

Details

Only for internal use of [poly3test](#) and [poly3ci](#).

Value

A list containing:

Y	number of tumours
n	number of individuals
estimate	poly-3-adjusted rates according to Bailer, Portier (1988)
weight	a vector of poly-3-adjusted weights, of length n
estp	poly-3-adjusted rate (according to method)
nadj	adjusted n (sum of weights)
varp	variance estimate (according to method)
varcor	variance estimate, if necessary corrected such that estimates of 0 can not occur

Note

Please note, that appropriateness of the point estimates seriously depends on whether the assumptions in Bailer and Portier are met or not.

References

Bailer, J.A. and Portier, C.J. (1988): Effects of treatment-induced mortality and tumor-induced mortality on tests for carcinogenicity in small samples. *Biometrics* 44, 417-431.

poly3estf

Only for internal use.

Description

Poly-3- adjusted point and variance estimates for long term carcinogenicity data if data are given as a numeric time vector, a logical status vector and a factor containing a grouping variable

Usage

```
poly3estf(time, status, tmax=NULL, f, method = "BP", k=NULL)
```

Arguments

time	a numeric vector of times of death of the individuals
status	a logical (or numeric, consisting of 0,1 only) vector giving the tumour status at time of death of each individual, where TRUE (1) = tumour present, FALSE (0) = no tumour present
tmax	a single numeric value, the time of sacrifice in the trial, or the last last time of death, defaults to the maximal value observed in time
f	a factor of the same length as time, status, giving the levels of a grouping variable in a one-way layout
method	a single character string, specifying the method for adjustment, with options: "BP" (Bailer Portier: assuming poly-3-adjusted rates are binomial variables), "BW" (Bieler, Williams: delta method as in Bieler-Williams (1993)) "ADD1" (as Bailer Portier, including an add1-adjustment on the raw tumour rates) "ADD2" (as Bailer Portier, including an add2-adjustment on the raw tumour rates following Agresti Caffo (2000) for binomials)
k	a single numeric value, the exponent to calculate survival adjusted proportions according to Bailer and Portier (1988), defaults to 3

Details

For internal use.

Value

A list containing:

Y	a numeric vector, groupwise number of tumours
n	a numeric vector, groupwise number of individuals
estimate	a numeric vector, groupwise poly-3-adjusted rates according to Bailer, Portier (1988)
weight	a numeric vector of poly-3-adjusted weights
estp	a numeric vector, groupwise poly-3-adjusted rate (according to method)
nadj	adjusted n (sum of weights)
varp	a numeric vector, groupwise variance estimate (according to method)
varcor	a numeric vector, groupwise variance estimate, if necessary corrected such that estimates of 0 can not occur
names	a character vector, the levels of the grouping variable f
k	a single numeric value, as input

Note

See [poly3est](#)

References

See [poly3est](#)

Examples

```
data(bronch)

poly3estf(status=bronch$Y, time=bronch$time, f=bronch$group, k=3)

poly3estf(status=bronch$Y, time=bronch$time, f=bronch$group, k=5)
```

poly3table	<i>Summarize long term carcinogenicity data</i>
------------	---

Description

Function to summarize data of long term carcinogenicity trials in a text format. Data are assumed to consist of (1) a dichotomous variable, defining whether the tumour of interest was present in an individual animal at time of death, and (2) a numeric variable containing the time of death of an individual animal, and (3) a grouping factor.

Usage

```
poly3table(time, status, f, tumour = NULL, symbol = "*")
```

Arguments

time	a numeric vector, containing the time of death of an individual
status	a logical (or dichotomous categorical) vector
f	a factor, specifying treatment groups
tumour	the value which status obtains if a tumour is present in an individual at time of death
symbol	symbol to indicate presence of tumour in the text representation

Value

A named list, containing a character string for each group

Author(s)

Frank Schaarschmidt

Examples

```
data(methyl)
methyl
poly3table(time=methyl$death, status=methyl$tumour,
  f=methyl$group, tumour = 1, symbol = "*")
```

poly3test

*Approximate simultaneous test for poly-3-adjusted tumour rates***Description**

P-value of maximum test and adjusted p-values for M contrasts of I groups in a one-way layout. Based on approximation of the true distribution of the M test statistics by an M-variate normal distribution.

Usage

```
poly3test(time, status, f, type = "Dunnett",
  cmat = NULL, method = "BP", alternative = "two.sided",
  dist="MVN", k=3, ...)
```

Arguments

time	a numeric vector of times of death of the individuals
status	a logical (or numeric, consisting of 0,1 only) vector giving the tumour status at time of death of each individual, where TRUE (1) = tumour present, FALSE (0) = no tumour present
f	a factor of the same length as time, status, giving the levels of a grouping variable in a one-way layout
type	a character string specifying the contrast type
cmat	an optional user defined contrast matrix of dimension MxI
method	a single character string, specifying the method for adjustment, with options: "BP" (Bailer Portier: assuming poly-3-adjusted rates are binomial variables), "BW" (Bieler, Williams: delta method as in Bieler-Williams (1993)) "ADD1" (as Bailer Portier, including an add1-adjustment on the raw tumour rates) "ADD2" (as Bailer Portier, including an add2-adjustment on the raw tumour rates following Agresti Caffo (2000) for binomials)
alternative	a character string specifying the direction of the alternative hypothesis
dist	a character string, where "MVN" invokes the computation of p-values using the multivariate normal distribution, and "N" invokes use p-value computation using the univariate normal distribution
k	a single numeric value, the exponent to calculate survival adjusted proportions according to Bailer and Portier (1988), defaults to 3
...	further arguments to be passed; currently only base, to choose the control group with type="Dunnett"

Details

Testversion.

Value

An object of class "poly3test", a list containing:

teststat	a numeric vector of teststatistics of length M
pval	a single numeric p-value, the p-value of the maximum test (minimum p-value)
p.val.adj	a vector of length M, the adjusted p-values of the single contrasts
alternative	a single character vector, as the input
dist	a character string specifying which distribution
time	as input
status	as input
f	as input
method	as input
cmat	used contrast matrix
sample.est	a list containing sample estimates

Note

Please note that all methods here described are only approximative, and might violate the nominal level in certain situations. Please note further that appropriateness of the point estimates, and consequently of tests and confidence intervals is based on the assumptions in Bailer and Portier (1988), which might be a matter of controversies.

References

Assumptions corresponding to the poly-k-adjustment:

Bailer, J.A. and Portier, C.J. (1988): Effects of treatment-induced mortality and tumor-induced mortality on tests for carcinogenicity in small samples. *Biometrics* 44, 417-431.

Peddada, S.D., Dinse, G.E., and Haseman, J.K. (2005): A survival-adjusted quantal response test for comparing tumor incidence rates. *Applied Statistics* 54, 51-61.

Statistical procedures and characterization of coverage probabilities are described in: Sill, M. (2007): Approximate simultaneous confidence intervals for multiple comparisons of binomial proportions. Master thesis, Institute of Biostatistics, Leibniz University Hannover.

Examples

```
# poly-3-adjusted tumour rates with a potential
# down-turn effect for the highest dose group "4":

data(methyl)

# many-to-one:
methylD<-poly3test(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Dunnett", method = "ADD1")

methylD
```

```

# Williams-Contrast:
methylW<-poly3test(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Williams", method = "ADD1", alternative="greater" )

methylW

# Changepoint-Contrast:
methylCh<-poly3test(time=methyl$death, status=methyl$tumour,
  f=methyl$group, type = "Change", method = "ADD1", alternative="greater" )

methylCh

```

powerbinom

Approximate power for multiple contrast tests of binomial proportions.

Description

Approximative power calculation for multiple contrast tests of binomial proportions (based on risk differences), based on probabilities of the multivariate standard normal distribution.

Usage

```

powerbinom(p, n, alpha = 0.05, type = "Dunnett", cmat = NULL,
  rhs = 0, alternative = c("two.sided", "less", "greater"),
  ptype = c("global", "anypair", "allpair"), method = "Wald", crit = NULL, ...)

```

Arguments

p	a numeric vector of assumed true proportions under the alternative hypothesis where <code>length(p)</code> is the number of (treatment) groups in the multiple comparison problem
n	a vector (of integer values), the number of observations per (treatment) group, must have the same length as p
alpha	a single numeric value, the alpha-level of the test, default is 0.05
type	a single character string naming the type of multiple contrast test to be applied on the proportions p; will be ignored if argument <code>cmat</code> is specified; for possible choices, see contrMat
cmat	optional, a matrix of contrast coefficients (with number of columns = number of treatment groups and number of rows = number of comparisons); if <code>cmat</code> is not specified, the type of multiple contrast test to be applied on the proportions p is chosen according to argument <code>type</code>
rhs	a (vector of) numeric value(s) between 0 and 1, specifying the right hand sides of the alternative hypotheses, default is 0, if values greater or less than 0 are specified, power for shifted tests (non-inferiority, superiority) is computed; Note that this makes primarily sense for one-sided hypotheses, in case of <code>rhs!=0</code> and <code>alternative="two.sided"</code> , a warning is given.

<code>alternative</code>	a character string, specifying the direction of the alternative hypotheses, options are "two.sided", "less", "greater"; <code>alternative</code> primarily controls for which directional decisions power is calculated, in case that argument <code>crit</code> is not specified, <code>alternative</code> also controls the choice of the critical value.
<code>ptype</code>	a single character string, naming the type of rejection probability to be computed; options are "global" for the overall rejection probability (irrespective which contrasts are under the null or the alternative), "anypair" for the rejection probability considering only those contrasts under the alternative, "global" for the probability that all elementary alternatives are rejected.
<code>method</code>	a character string, the method for variance estimation in test / confidence interval construction: one of "Wald", "ADD1", "ADD2".
<code>crit</code>	a single numeric value to serve as equicoordinate critical point in the multiple test; if it is not specified, it is computed as a quantile of the multivariate normal distribution based on the specifications in arguments <code>n</code> , <code>cmat</code> (or <code>type</code>); note that for alternatives 'two.sided' and 'greater', <code>crit</code> should be a single positive value, while for alternative 'less', <code>crit</code> should be a single negative value.
<code>...</code>	further arguments, which are passed to the functions <code>qmvnorm</code> and <code>pmvnorm</code> , mainly to control the computation errors, see GenzBretz for details

Details

For (standard type and user-defined) multiple contrast tests of proportions in 2xk contingency tables assuming the binomial distribution and k treatment groups, different types of rejection probabilities in the multiple testing problem can be computed. Based on a central multivariate normal distribution, equicoordinate critical points for the test are computed, different critical points for the test can be specified in `crit`.

Via computing probabilities of non-central multivariate normal distributions `pmvt(mvtnorm)` one can calculate:

The global rejection probability (`power="global"`), i.e. the probability that at least one of the elementary null hypotheses is rejected, irrespective, whether this (or any contrast!) is under the corresponding elementary alternative). As a consequence this probability involves elementary type-II-errors for those contrasts which are under their elementary null hypothesis.

The probability to reject at least one of those elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="anypair"`). Technically, this is achieved by omitting those contrasts under the elementary null and compute the rejection probability for a given critical value from a multivariate normal distribution with reduced dimension. Note that for 'two-sided' alternatives, type III-errors (rejection of the two-sided null in favor of the wrong direction) are included in the power.

The probability to reject all elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="allpair"`). Also here, for 'two-sided' alternatives type III-error contribute to the computed 'allpair power'. Note further that two-sided allpair power is simulated based on multivariate normal random numbers.

Value

A list consisting of the following items:

power	a numeric value the computed power, with estimated computational error as an attribute
p	the input vector of expected groupwise proportions
n	the input vector of group sample sizes
conexp	a data frame containing the contrast matrix, the expected values of the contrasts given p (expContrast), the right hand sides of the hypotheses (rhs, as input), the expected values of the test statistics corresponding to the contrasts and rhs, and a column of logical values indicating whether the corresponding contrasts was under the alternative (under HA)
crit	a single numeric value, the critical value used for power computation
alternative	a single character string, as input
ptype	a single character string, as input
alpha	a single numeric value, as input

Warning

Note that the the tests for which power is computed as well as the power computation itself relies on simple approximations of binomial distribution by normal distributions. It is known that the test procedures do not control FWER for small n and extreme proportions p. Hence, computed power may substantially deviate from the true power of the methods a) due to the fact that the tests have sizes deviating from the nominal level, b) due to insufficient approximation in the power calculation. Simulations show that absolute deviations of approximate power from simulated power are large if low values of $n \cdot p$ or $n \cdot (1-p)$ are involved and if power is low (i.e. $\text{power} < 0.3$).

Author(s)

Frank Schaarschmidt

References

Genz A, Bretz F (1999): Numerical computation of multivariate t-probabilities with application to power calculation of multiple contrasts. *Journal of Statistical Computation and Simulation*, 63, 4, 361-378. *Bretz F, Hothorn LA (2002):* Detecting dose-response using contrasts: asymptotic power and sample size determination for binomial data. *Statistics in Medicine*, 21, 22, 3325-3335.

Examples

```
# Assume, one wants to perform a test for increasing trend
# using Williams type of contrasts among I=5 groups
# (e.g. 4 doses and one control).
# Proportions are assumed to have values
# pi=(0.1,0.12,0.14,0.14,0.2) under the alternative.

powerbinom(p=c(0.1, 0.12, 0.14, 0.14, 0.2),
n=c(100,100,100,100,100), type = "Williams",
alternative = "greater")
```

```

powerbinom(p=c(0.1, 0.12, 0.14, 0.14, 0.2),
n=c(150,150,150,150,150), type = "Williams",
alternative = "greater")

powerbinom(p=c(0.1, 0.12, 0.14, 0.14, 0.2),
n=c(190,140,140,140,140), type = "Williams",
alternative = "greater")

# probability to show for at least one group (2,3,4)
# a significant reduction versus control (1)

powerbinom(p=c(0.3, 0.15, 0.15, 0.15),
n=c(140,140,140,140), type = "Dunnett",
alternative = "less")

# probability to show for at least one group (2,3,4)
# a significant reduction versus control (1) of more
# than 0.05 percent

powerbinom(p=c(0.3, 0.15, 0.15, 0.15),
n=c(140,140,140,140), type = "Dunnett",
alternative = "less", rhs=-0.05)

# probability to show for all groups (2,3,4)
# a significant reduction versus control (1) of more
# than 0.05 percent

powerbinom(p=c(0.3, 0.15, 0.15, 0.15),
n=c(140,140,140,140), type = "Dunnett",
alternative = "less", rhs=-0.05, ptype="allpair")

```

powermcpn

Approximative power calculation for multiple contrast tests

Description

Approximative power calculation for multiple contrast tests which are based on normal approximation.

Usage

```
powermcpn(ExpTeststat, corrH1, crit, alternative = c("two.sided", "less", "greater"),
alpha = 0.05, ptype = c("global", "anypair", "allpair"), ...)
```

Arguments

ExpTeststat numeric vector: the expectation of the test statistics under the alternative

<code>corrH1</code>	a numeric matrix, the correlation matrix of the teststatistics under the alternative, must have same number of columns and rows as length of <code>ExpTeststat</code>
<code>crit</code>	a single numeric value, the critical value of the test; if not specified a critical value is computed as an equicoordinate $(1-\alpha)$ quantile of a central multivariate normal distribution with correlation <code>corrH1</code>
<code>alternative</code>	a character string, specifying the direction of the alternative hypotheses, options are "two.sided", "less", "greater"; <code>alternative</code> primarily controls for which directional decisions power is calculated, in case that argument <code>crit</code> is not specified, <code>alternative</code> also controls the choice of the critical value.
<code>alpha</code>	a single numeric value: the FWER for the multiple test, defaults to 0.05
<code>ptype</code>	a single character string, naming the type of rejection probability to be computed; options are "global" for the overall rejection probability (irrespective which contrasts are under the null or the alternative), "anypair" for the rejection probability considering only those contrasts under the alternative, "global" for the probability that all elementary alternatives are rejected.
<code>...</code>	further arguments, which are passed to the functions <code>qmvnorm</code> and <code>pmvnorm</code> , mainly to control the computation errors, see <code>help GenzBretz(mvtnorm)</code> for details

Details

For (standard type and user-defined) multiple contrast tests based on approximation with the multivariate normal distribution, three types of rejection probabilities in the multiple testing problem can be computed.

The global rejection probability (`power="global"`), i.e. the probability that at least one of the elementary null hypotheses is rejected, irrespective, whether this (or any contrast!) is under the corresponding elementary alternative). As a consequence this probability involves elementary type-II-errors for those contrasts which are under their elementary null hypothesis.

The probability to reject at least one of those elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="anypair"`). Technically, this is achieved by omitting those contrasts under the elementary null and compute the rejection probability for a given critical value from a multivariate normal distribution with reduced dimension. Note that for 'two-sided' alternatives, type III-errors (rejection of the two-sided null in favor of the wrong direction) are included in the power.

The probability to reject all elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="allpair"`). Also here, for 'two-sided' alternatives type III-error contribute to the computed 'allpair power'. Note further that two-sided allpair power is simulated based on multivariate normal random numbers.

Value

A list consisting of the following items:

<code>power</code>	a numeric value the computed power, with estimated computational error as an attribute
--------------------	--

conexp	the expected values of the test statistics corresponding to the contrasts and rhs, and a column with [0,1] values, indicating whether the corresponding contrasts was under the alternative (1) or under the null (0)
crit	a single numeric value, the critical value used for power computation
alternative	a single character string, as input
ptype	a single character string, as input
alpha	a single numeric value, as input

Author(s)

Frank Schaarschmidt

powermcpt	<i>Testversion. Power calculation for multiple contrast tests (1-way ANOVA model)</i>
-----------	---

Description

Testversion. Calculate the power of a multiple contrast tests of k means in a model with homogeneous Gaussian errors, using the function pmvt(mvtnorm) to calculate multivariate t probabilities. Different options of power definition are "global": the overall rejection probability (the probability that at the elementary null is rejected for at least one contrast, irrespective of being under the elementary null or alternative), "anypair": the probability to reject any of the elementary null hypotheses for those contrasts that are under the elementary alternatives, "allpair": and the probability that all elementary nulls are rejected which are indeed under the elementary nulls. See Sections 'Details' and 'Warnings'!

Usage

```
powermcpt(mu, n, sd, cmat = NULL, rhs=0, type = "Dunnett",
  alternative = c("two.sided", "less", "greater"), alpha = 0.05,
  ptype = c("global", "anypair", "allpair"), crit = NULL, ...)
```

Arguments

mu	a numeric vector of expected values in the k treatment groups
n	a numeric vector of sample sizes in the k treatment groups
sd	a single numeric value, specifying the expected standard deviation of the residual error
cmat	optional specification of a contrast matrix; if specified, it should have as many columns as there are groups in arguments mu and n and it should have at least 2 rows, if specified, argument type is ignored, if not specified, the contrast is determined by argument type
rhs	numeric vector, specifying the right hand side of the hypotheses to test, defaults to 0, other specifications lead to tests of non-inferiority and superiority.

<code>type</code>	a single character string, naming one of the contrast types available in <code>contrMat(multcomp)</code> ; argument is ignored if <code>cmat</code> is specified
<code>alternative</code>	a single character string, specifying the direction of the alternative hypothesis, one of <code>"two.sided"</code> , <code>"less"</code> , <code>"greater"</code> . Note that this argument governs how the multivariate t probabilities are evaluated as well as the computation of the critical value if none is specified (i.e. default <code>crit=NULL</code>)
<code>alpha</code>	a single numeric value, familywise type I error to be controlled, is ignored if argument <code>crit</code> is specified
<code>pptype</code>	a single character string, naming the type of rejection probability to be computed; options are <code>"global"</code> for the global rejection probability, <code>"anypair"</code> for the rejection probability considering only those contrasts under the alternative, <code>"global"</code> for the probability that all elementary alternatives are rejected.
<code>crit</code>	a single numeric value to serve as equicoordinate critical point in the multiple test; if it is not specified, it is computed as a quantile of the multivariate t distribution based on the specifications in arguments <code>n</code> , <code>cmat</code> (or <code>type</code>); note that for alternatives <code>'two.sided'</code> and <code>'greater'</code> , <code>crit</code> should be a single positive value, while for alternative <code>'less'</code> , <code>crit</code> should be a single negative value.
<code>...</code>	further arguments, which are passed to the functions <code>qmv</code> and <code>pmvt</code> , mainly to control the computation errors, see <code>help GenzBretz(mvtnorm)</code> for details

Details

In a homoscedastic Gaussian model with k possibly different means compared by (user-defined) multiple contrast tests, different types of rejection probabilities in the multiple testing problem can be computed. Based on a central multivariate t distribution with $df = \text{sum}(n) - k$ appropriate equicoordinate critical points for the test are computed, different critical points can be specified in `crit`. Computing probabilities of non-central multivariate t distributions `pmvt(mvtnorm)` one can calculate:

The global rejection probability (`power="global"`), i.e. the probability that at least one of the elementary null hypotheses is rejected, irrespective, whether this (or any contrast!) is under the corresponding elementary alternative). As a consequence this probability involves elementary type-II-errors for those contrasts which are under their elementary null hypothesis.

The probability to reject at least one of those elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="anypair"`). Technically, this is achieved by omitting those contrasts under the elementary null and compute the rejection probability for a given critical value from a multivariate t distribution with reduced dimension. Note that for `'two-sided'` alternatives, type III-errors (rejection of the two-sided null in favor of the wrong direction) are included in the power.

The probability to reject all elementary null hypotheses which are indeed under their corresponding elementary alternatives (`power="allpair"`). Also here, for `'two-sided'` alternatives type III-error contribute to the computed `'allpair power'`. Note further that two-sided `allpair power` is simulated based on multivariate t random numbers.

Value

A list consisting of the following items:

power	a numeric value the computed power, with estimated computational error as an attribute
mu	the input vector of expected values of group means
n	the input vector of group sample sizes
conexp	a data frame containing the contrast matrix, the expected values of the contrasts given mu (expContrast), the right hand sides of the hypotheses (rhs, as input), the expected values of the test statistics corresponding to the contrasts and rhs, and a column of logical values indicating whether the corresponding contrasts was under the alternative (under HA)
crit	a single numeric value, the critical value used for power computation
alternative	a single character string, as input
ptype	a single character string, as input
alpha	a single numeric value, as input

Warning

This is a test version, which has roughly (but not for an extensive number of settings) been checked by simulation. Any reports of errors/odd behaviour/amendments are welcome.

Author(s)

Frank Schaarschmidt

References

Genz A, Bretz F (1999): Numerical computation of multivariate t-probabilities with application to power calculation of multiple contrasts. *Journal of Statistical Computation and Simulation*, 63, 4, 361-378. *Bretz F, Hothorn LA (2002):* Detecting dose-response using contrasts: asymptotic power and sample size determination for binomial data. *Statistics in Medicine*, 21, 22, 3325-3335. *Bretz F, Hayter AJ and Genz A (2001):* Critical point and power calculations for the studentized range test for generally correlated means. *Journal of Statistical Computation and Simulation*, 71, 2, 85-97. *Dilba G, Bretz F, Hothorn LA, Guizard V (2006):* Power and sample size computations in simultaneous tests for non-inferiority based on relative margins. *Statistics in Medicien* 25, 1131-1147.

Examples

```
powermcpt(mu=c(3,3,5,7), n=c(10,10,10,10), sd=2, type = "Dunnett",
  alternative ="greater", ptype = "global")
powermcpt(mu=c(3,3,5,7), n=c(10,10,10,10), sd=2, type = "Williams",
  alternative ="greater", ptype = "global")

powermcpt(mu=c(3,3,5,7), n=c(10,10,10,10), sd=2, type = "Dunnett",
  alternative ="greater", ptype = "anypair")
powermcpt(mu=c(3,3,5,7), n=c(10,10,10,10), sd=2, type = "Williams",
  alternative ="greater", ptype = "anypair")

powermcpt(mu=c(3,4,5,7), n=c(10,10,10,10), sd=2, type = "Dunnett",
```

```

alternative = "greater", ptype = "allpair")
powermcpt(mu=c(3,2,1,-1), n=c(10,10,10,10), sd=2, type = "Dunnett",
alternative = "greater", ptype = "allpair")

```

PQln	<i>Generalized pivotal quantity for log expected values of several lognormal samples.</i>
------	---

Description

Computes samples of the generalized pivotal quantities for the log of expected values of lognormal samples according to Krishnamoorthy and Mathew (2003). For internal use in `Indci`, `lnrci`.

Usage

```
PQln(nx, mlx, varlx, B = 10000)
```

Arguments

<code>nx</code>	a numeric (integer) vector, the sample sizes for each of K groups
<code>mlx</code>	a numeric vector, the sample estimates of the log mean for each of K groups
<code>varlx</code>	a numeric vector, the (unbiased) sample estimates of the log variance for each of K groups
<code>B</code>	a single (integer) value, the number of samples to draw, defaults to 10000

Details

Generates B samples from independent standard normal and χ^2 distributions for each of K independent groups in a one-way layout and computes the generalized pivotal quantities for the log expected value for each of the groups, according to Krishnamoorthy and Mathew (2003). The input vectors `nx`, `mlx`, `varlx` must be vectors of length K.

Value

A BxK matrix

Author(s)

Frank Schaarschmidt

References

Krishnamoorthy K, Mathew T (2003): Inferences on the means of lognormal distributions using generalized p-values and generalized confidence intervals. *Journal of Statistical Planning and Inference* 115, 103-121.

print.binomORci	<i>Print method for binomORci</i>
-----------------	-----------------------------------

Description

Print method for objects of class "binomORci" as can be created by calling binomORci.

Usage

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

Arguments

x	an object of class "binomRDci"
...	...

Value

A print out.

print.binomRDci	<i>Print method for binomRDci</i>
-----------------	-----------------------------------

Description

Print method for objects of class "binomRDci" as can be created by calling binomRDci.

Usage

```
## S3 method for class 'binomRDci'
print(x, digits = 4, ...)
```

Arguments

x	an object of class "binomRDci"
digits	a single integer value specifying the number of digits for rounding
...	further arguments to be passed to "print.default"

Value

A print out.

`print.binomRDtest` *Print method for binomRDtest*

Description

Print method for objects of class "binomRDtest" as can be created by calling `binomRDtest`.

Usage

```
## S3 method for class 'binomRDtest'  
print(x, digits = 4, ...)
```

Arguments

<code>x</code>	an object of class "binomRDtest"
<code>digits</code>	a single integer value specifying the number of digits for rounding
<code>...</code>	further arguments to be passed to "print.default"

Value

A print out.

`print.binomRRci` *Print method for binomRRci*

Description

Print method for objects of class "binomRRci" as can be created by calling `binomRDci`.

Usage

```
## S3 method for class 'binomRRci'  
print(x, digits = 4, ...)
```

Arguments

<code>x</code>	an object of class "binomRRci"
<code>digits</code>	a single integer value specifying the number of digits for rounding
<code>...</code>	further arguments to be passed to "print.default"

Value

A print out.

print.poly3ci	<i>Print method for poly3ci</i>
---------------	---------------------------------

Description

Print method for objects of class "poly3ci" as can be created by calling poly3ci

Usage

```
## S3 method for class 'poly3ci'  
print(x, digits = 4, ...)
```

Arguments

x	an object of class "poly3ci"
digits	a single integer value specifying the number of digits for rounding
...	further arguments to be passed to "print.default"

Value

A print out.

print.poly3est	<i>Print function for poly3est</i>
----------------	------------------------------------

Description

Print function for objects of class "poly3est", as can be created by calling "poly3estf"

Usage

```
## S3 method for class 'poly3est'  
print(x, digits = 4, ...)
```

Arguments

x	An object of class "poly3est", as can be created by calling poly3estf
digits	a single integer, the number of digits for rounding of the results
...	currently not used...

print.poly3test *Print method for poly3test*

Description

Print method for objects of class "poly3test" as can be created by calling poly3test.

Usage

```
## S3 method for class 'poly3test'  
print(x, digits = 4, ...)
```

Arguments

x	an object of class "poly3test"
digits	a single integer value specifying the number of digits for rounding
...	further arguments to be passed to "print.default"

Value

A print out.

print.Shannonci *Print function for Shannonci*

Description

Print out the confidence intervals calculated by Shannonci.

Usage

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

Arguments

x	An object of class "Shannonci", as produced by calling Shannonci
...	currently only digits is passed to print

Value

A print out.

See Also

For a more detailed print out, use [summary.Shannonci](#)

print.Simpsonci	<i>Print function for Simpsonci</i>
-----------------	-------------------------------------

Description

Print out the confidence intervals calculated by Simpsonci.

Usage

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

Arguments

x	An object of class "Simpsonci", as produced by calling Simpsonci
...	currently only digits is passed to print

Value

A print out.

scalechoice	<i>Only for internal use.</i>
-------------	-------------------------------

Description

How to define scale parameters of the Weibull distribution in the poly-k-setting

Usage

```
scalechoice(shape = 3, pm = 0.2, t = 1)
```

Arguments

shape	shape parameter (usually shape=3)
pm	a numeric value ($0 < p < 1$), probability that an individual dies before time t
t	a numeric value, time t

Value

a single numeric value, giving the scale parameter needed to fulfill $pm(t)$ for the current definition of the Weibull distribution.

Note

Scale parameter is defined differently than that in Peddada (2005)

See Also

?pweibull

SCSrank	<i>Compute a rectangular simultaneous confidence set from a sample of a joint empirical distribution.</i>
---------	---

Description

Given a large sample of N values from an M dimensional joint empirical distribution, the rank based method of Besag et al. (1995) is used to compute a rectangular M -dimensional 'confidence' set that includes $N \cdot \text{conf.level}$ values of the sample.

Usage

```
SCSrank(x, conf.level = 0.95, alternative = "two.sided", ...)
```

Arguments

<code>x</code>	an $N \times M$ matrix containing N sampled values of the M dimensional distribution of interest
<code>conf.level</code>	the simultaneous confidence level, a single numeric value between 0 and 1, defaults to 0.95 for simultaneous 95 percent sets
<code>alternative</code>	a single character string related to hypotheses testing, "two.sided" invokes two-sided confidence sets, "less" invokes sets with upper limits only and "greater" invokes sets with lower limits only,
<code>...</code>	currently ignored

Value

an $M \times 2$ (`alternative="two.sided"`) matrix containing the lower and upper confidence limit for the M dimensions, in case of `alternative="less"`, `alternative="greater"` the lower and upper bounds are replaced by `-Inf` and `Inf`, respectively.

Author(s)

Frank Schaarschmidt

References

Besag J, Green P, Higdon D, Mengersen K (1995). Bayesian Computation and Stochastic Systems. *Statistical Science* 10, 3-66. Mandel M, Betensky RA. Simultaneous confidence intervals based on the percentile bootstrap approach. *Computational Statistics and Data Analysis* 2008; 52(4): 2158-2165.

Examples

```
x <- cbind(rnorm(1000,1,2), rnorm(1000,0,2), rnorm(1000,0,0.5), rnorm(1000,2,1))
dim(x)
cm <- rbind(c(-1,1,0,0), c(-1,0,1,0), c(-1, 0,0,1))
xd <- t(apply(x, 1, function(x){crossprod(t(cm), matrix(x))}))
pairs(xd)

SCSrank(xd, conf.level=0.9)
```

Shannon	<i>Shannon index</i>
---------	----------------------

Description

Calculates the Shannon-Wiener index.

Usage

```
Shannon(p)
```

Arguments

`p` A numeric vector of proportions.

Shannonci	<i>Confidence intervals for multiple contrasts of Shannon indices</i>
-----------	---

Description

Calculates simultaneous and local confidence intervals for differences of Shannon indices.

Usage

```
Shannonci(X, f, cmat = NULL, type = "Dunnett", alternative = "two.sided",
  conf.level = 0.95, dist = "MVN", ...)
```

Arguments

<code>X</code>	a data.frame of dimensions n times p with integer entries, where n is the number of samples and p is the number of species
<code>f</code>	a factor variable of length n, grouping the observations in X
<code>cmat</code>	an contrast matrix; the number of columns should match the number of levels in f
<code>type</code>	a single character string, currently one of "Dunnett", "Tukey", "Sequen"
<code>alternative</code>	a single character string, one of "two.sided", "less" (upper bounds), "greater" (lower bounds)
<code>conf.level</code>	the confidence level of the simultaneous (or local) confidence intervals
<code>dist</code>	a single character string, defining the type of quantiles to be used for interval calculation; "MVN" invokes simultaneous intervals, "N" invokes unadjusted confidence intervals with coverage probability conf.level for each of them
<code>...</code>	further arguments to be passed; currently only base is used, a single integer value, specifying which group to be taken as the control in case that type="Dunnett", ignored otherwise

Details

This function implements confidence intervals described by Fritsch and Hsu (1999) for the difference of Shannon indices between several groups. Deviating from Fritsch and Hsu, quantiles of the multivariate normal distribution are used which take the sample variance into account.

Note, that this approach ignores the variability of the individual samples.

Value

A list containing the elements:

<code>conf.int</code>	a matrix, containing the lower and upper confidence limits in the columns
<code>quantile</code>	a single numeric value, the quantile used for interval calculation
<code>estimate</code>	a matrix, containing the point estimates of the contrasts in its column
<code>cmat</code>	the contrast matrix used
<code>methodname</code>	a character string, for printing
<code>sample.estimate</code>	A list of sample estimates as returned by estShannonf

and some of the input arguments

Author(s)

Frank Schaarschmidt

References

Fritsch, KS, and Hsu, JC (1999): Multiple Comparison of Entropies with Application to Dinosaur Biodiversity. *Biometrics* 55, 1300-1305.

See Also

[Simpsonci](#) for simultaneous and local intervals of differences of the Simpson index

Examples

```

data(HCD)

HCDcounts<-HCD[,-1]
HCDf<-HCD[,1]

# Comparison to the confidence bounds shown in
# Fritsch and Hsu (1999), Table 5, "Standard normal".

cmat<-rbind(
  "HM-HU"=c(0,1,-1),
  "HL-HM"=c(1,-1,0),
  "HL-HU"=c(1,0,-1)
)

Shannonci(X=HCDcounts, f=HCDf, cmat=cmat,
  alternative = "two.sided", conf.level = 0.9, dist = "N")

# Note, that the calculated confidence intervals
# differ from those published by Fritsch and Hsu (1999),
# whenever Lower is involved.

# Comparison to the lower cretaceous,
# unadjusted confidence intervals:

Shannonci(X=HCDcounts, f=HCDf, type = "Dunnett",
  alternative = "greater", conf.level = 0.9, dist = "N")

# Stepwise comparison between the strata,
# unadjusted confidence intervals:

ShannonS<-Shannonci(X=HCDcounts, f=HCDf, type = "Sequen",
  alternative = "greater", conf.level = 0.9, dist = "N")

ShannonS

summary(ShannonS)

plot(ShannonS)

# A trend test based on multiple contrasts:

```

```

cmatTREND<-rbind(
  "U-LM"=c(-0.5,-0.5,1),
  "MU-L"=c(-1,0.5,0.5),
  "U-L"=c(-1,0,1)
)

TrendCI<-Shannonci(X=HCDcounts, f=HCDf, cmat=cmatTREND,
  alternative = "greater", conf.level = 0.95, dist = "MVN")
TrendCI

plot(TrendCI)

```

Simpson

Calculate the Simpson index

Description

Calculate the Simpson index from a vector of proportions.

Usage

```
Simpson(p)
```

Arguments

p a numeric vector of proportions

Simpsonci

Confidence intervals for differences of Simpson indices

Description

Calculates simultaneous and local confidence intervals for differences of Simpson indices.

Usage

```
Simpsonci(X, f, cmat = NULL, type = "Dunnett", alternative = "two.sided", conf.level = 0.95, dist = "MVN")
```

Arguments

<code>X</code>	a data.frame of dimensions n times p with integer entries, where n is the number of samples and p is the number of species
<code>f</code>	a factor variable of length n, grouping the observations in X
<code>cmat</code>	an contrast matrix; the number of columns should match the number of levels in f
<code>type</code>	a single character string, currently one of "Dunnett", "Tukey", "Sequen"
<code>alternative</code>	a single character string, one of "two.sided", "less" (upper bounds), "greater" (lower bounds)
<code>conf.level</code>	the confidence level of the simultaneous (or local) confidence intervals
<code>dist</code>	a single character string, defining the type of quantiles to be used for interval calculation; "MVN" invokes simultaneous intervals, "N" invokes unadjusted confidence intervals with coverage probability <code>conf.level</code> for each of them
<code>...</code>	further arguments to be passed; currently only <code>base</code> is used, a single integer value, specifying which group to be taken as the control in case that <code>type="Dunnett"</code> , ignored otherwise

Details

This function implements confidence intervals described by Rogers and Hsu (1999) for the difference of Shannon indices between several groups. Deviating from Fritsch and Hsu, quantiles of the multivariate normal distribution are used which take the sample variance into account.

Note, that this approach ignores the variability of the individual samples.

Value

A list containing the elements:

<code>conf.int</code>	a matrix, containing the lower and upper confidence limits in the columns
<code>quantile</code>	a single numeric value, the quantile used for interval calculation
<code>estimate</code>	a matrix, containing the point estimates of the contrasts in its column
<code>cmat</code>	the contrast matrix used
<code>methodname</code>	a character string, for printing
<code>sample.estimate</code>	

A list of sample estimates as returned by `estShannonf`

and some of the input arguments.

Author(s)

Frank Schaarschmidt

References

Rogers, JA and Hsu, JC (2001): Multiple Comparisons of Biodiversity. *Biometrical Journal* 43, 617-625.

See Also[Shannonci](#)**Examples**

```
data(HCD)

HCDcounts<-HCD[,-1]
HCDf<-HCD[,1]

# Rogers and Hsu (2001), Table 2:
# All pair wise comparisons:

Simpsonci(X=HCDcounts, f=HCDf, type = "Tukey",
  conf.level = 0.95, dist = "MVN")

# Rogers and Hsu (2001), Table 3:
# Comparison to the lower cretaceous:

Simpsonci(X=HCDcounts, f=HCDf, type = "Dunnett",
  alternative = "less", conf.level = 0.95, dist = "MVN")

# Note, that the confidence bounds here differ
# from the bounds in Rogers and Hsu (2001)
# in the second digit, whenever the group Upper
# is involved in the comparison.

# Stepwise comparison between the strata:

SimpsonS<-Simpsonci(X=HCDcounts, f=HCDf, type = "Sequen",
  alternative = "greater", conf.level = 0.95, dist = "MVN")

SimpsonS
summary(SimpsonS)

plot(SimpsonS)

# # # Hell Creek Dinosaur data:
# # Is there a downward trend in biodiversity during the
# # Cretaceous period?

# A trend test based on multiple contrasts:

cmatTREND<-rbind(
  "U-LM"=c(-0.5,-0.5,1),
  "MU-L"=c(-1,0.5,0.5),
  "U-L"=c(-1,0,1)
)
```

```
TrendCI<-Simpsonci(X=HCDcounts, f=HCDf, cmat=cmatTREND, alternative = "greater", conf.level = 0.95, dist = "MVN")
TrendCI

plot(TrendCI)
```

summary.binomORci *Detailed print out for binomORci*

Description

Produces a more detailed print out of objects of class "binomORci", including summary statistics, the used contrast matrix and the confidence intervals.

Usage

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

Arguments

object	an object of class "binomORci" as created by function binomORci
...	...

Value

A print out.

Examples

```
x<-c(1,3,6,7,5)
n<-c(30,30,30,30,30)
names<-LETTERS[1:5]

ORD<-binomORci(x=x, n=n, names=names,
  type="Dunnett", alternative="greater")
summary(ORD)

ORW<-binomORci(x=x, n=n, names=names,
  type="Williams", alternative="greater")
summary(ORW)
```

summary.binomRDci *Detailed print out for binomRDci*

Description

Produces a more detailed print out of objects of class "binomRDci", including summary statistics, the used contrast matrix and the confidence intervals.

Usage

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

Arguments

object	an object of class "binomRDci" as created by function binomRDci
...	further arguments to be passed to summary, currently only digits for rounding is supported

Value

A print out.

Examples

```
data(liarozole)  
head(liarozole)  
LiWi<-binomRDci(Improved ~ Treatment, data=liarozole,  
type="Williams")  
LiWi  
summary(LiWi)
```

summary.binomRDtest *Detailed print out for binomRDtest*

Description

Produces a more detailed print out of objects of class "binomRDtest", including summary statistics, the used contrast matrix and the p-values.

Usage

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

Arguments

object an object of class "binomRDtest" as created by function [binomRDtest](#)
... further arguments to be passed to summary, currently only digits for rounding is supported

Value

A print out.

Examples

```
ntrials <- c(40,20,20,20)  
xsucceses <- c(1,2,2,4)  
names(xsucceses) <- LETTERS[1:4]  
test<-binomRDtest(x=xsucceses, n=ntrials, method="ADD1",  
  type="Changepoint", alternative="greater")  
  
test  
  
summary(test)
```

summary.binomRRci *Detailed print out for binomRRci*

Description

Produces a more detailed print out of objects of class "binomRRci", including summary statistics, the used contrast matrix and the confidence intervals.

Usage

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

Arguments

```
object      an object of class "binomRRci" as created by function binomRRci
...         further arguments to be passed to summary, currently only digits for rounding
           is supported
```

Value

A print out.

Examples

```
data(liarozole)

head(liarozole)

LiDu<-binomRRci(Improved ~ Treatment, data=liarozole,
  type="Dunnett", alternative="greater")

LiDu

summary(LiDu)
```

summary.poly3est *Detailed print out for poly3est*

Description

Summary statistics for long-term carcinogenicity data, including poly-3-estimates. For internal use.

Usage

```
## S3 method for class 'poly3est'
summary(object, ...)
```

Arguments

```
object      An object of class "poly3est", as can be obtained by poly3est
...         further argument for the print out, as e.g. digits for rounding
```

Details

For internal use.

Value

A print out.

Author(s)

Frank Schaarschmidt

Examples

```
data(methyl)
head(methyl)
```

```
estk3<-poly3estf(time=methyl$death, status=methyl$tumour, f=methyl$group)
summary(estk3)
```

```
estk5<-poly3estf(time=methyl$death, status=methyl$tumour, f=methyl$group, k=5)
summary(estk5)
```

summary.Shannonci *Summary for Shannonci*

Description

Produces a detailed print out of the results of the function Shannonci.

Usage

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

Arguments

object An object of class "Shannonci", see [Shannonci](#)
... further arguments to be passed to print, currently only digits

Value

A print out, comprising a table of the (possibly aggregated) data used for estimation, the sample estimates for the Shannon index with bias corrected and raw values, its variance estimates, the used contrast matrix, and the confidence intervals.

Examples

```

data(HCD)

HCDcounts<-HCD[,-1]
HCDf<-HCD[,1]

# Comparison to the confidence bounds shown in
# Fritsch and Hsu (1999), Table 5, "Standard normal".

cmat<-rbind(
  "HM-HU"=c(0,1,-1),
  "HL-HM"=c(1,-1,0),
  "HL-HU"=c(1,0,-1)
)

ShannonS<-Shannonci(X=HCDcounts, f=HCDf, type = "Sequen",
  alternative = "greater", conf.level = 0.9, dist = "N")

summary(ShannonS)

```

summary.Simpsonci	<i>Summary function for Simpsonci</i>
-------------------	---------------------------------------

Description

Produces a detailed print out of the results of function Simpsonci.

Usage

```

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

```

Arguments

object	an object of class "Simpsonci" as obtained by calling Simpsonci
...	further arguments to be passed to print and round: currently only digits

Value

A print out, comprising a table of the (possibly aggregated) data used for estimation, the sample estimates for the Simpsons index, and its variance estimates, the used contrast matrix, and the confidence intervals.

Examples

```

data(HCD)

HCDcounts<-HCD[,-1]
HCDf<-HCD[,1]

SimpsonS<-Simpsonci(X=HCDcounts, f=HCDf, type = "Sequen",
  alternative = "greater", conf.level = 0.95, dist = "MVN")

summary(SimpsonS)

```

Waldci

Simultaneous Wald confidence intervals

Description

General function for simultaneous CIs in a one-way layout using multivariate normal distribution.

Usage

```
Waldci(cmat, estp, varp, varcor, alternative = "two.sided", conf.level = 0.95, dist="MVN")
```

Arguments

cmat	Contrast matrix of dimension $M \times I$, with M = the number of contrasts, I = the number of samples
estp	numeric vector of point estimates of length I , with I = the number of samples
varp	numeric vector of variance estimates of length I , to be used for interval construction
varcor	numeric vector of variance estimates of length I
alternative	character string
conf.level	single numeric vector
dist	a character string, "MVN" invokes multiplicity adjustment via the multivariate normal distribution, "N" invokes use of quantiles of the univariate normal distribution

Details

Mainly for internal use.

Value

A list containing:

<code>conf.int</code>	a matrix with 2 columns: lower and upper confidence bounds, and M rows
<code>alternative</code>	character string, as input
<code>conf.level</code>	single numeric value, as input
<code>quantile</code>	the quantile used to construct the CIs

Author(s)

Frank Schaarschmidt

See Also

For user level implementations see: [binomRDci](#), [binomORci](#), [poly3ci](#)

Waldtest

Simultaneous Wald tests

Description

General function for adjusted p-values for an UIT in a one-way layout using multivariate normal distribution.

Usage

```
Waldtest(estp, varp, cmat, alternative = "greater", dist="MVN")
```

Arguments

<code>estp</code>	numeric vector of point estimates of length I, with I = the number of samples
<code>varp</code>	numeric vector of variance estimates of length I, to be used for interval construction
<code>cmat</code>	Contrast matrix of dimension MxI, with M = the number of contrasts, I= the number of samples
<code>alternative</code>	character string
<code>dist</code>	a character string, where "MVN" invokes the computation of p-values using the multivariate normal distribution, and "N" invokes use p-value computation using the univariate normal distribution

Value

A list containing:

<code>teststat</code>	a numeric vector of teststatistics of length M
<code>pval</code>	a single numeric p-value, the p-value of the maximum test (minimum p-value)
<code>p.val.adj</code>	a vector of length M, the adjusted p-values of the single contrasts
<code>alternative</code>	a single character vector, as the input
<code>dist</code>	a character string specifying which distribution

Author(s)

Frank Schaarschmidt

See Also

For user level implementations see:

[binomRDtest](#), [poly3test](#)

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