

Package ‘tukeytrend’

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Type Package

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Title Tukeys Trend Test via Multiple Marginal Models

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Author Frank Schaarschmidt [aut, cre],
Christian Ritz [aut],
Ludwig Hothorn [ctb]

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

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SimComp, HSAUR3

VignetteBuilder knitr

Description Provides wrapper functions to the multiple marginal model function `mmm()` of package ‘multcomp’ to implement the trend test of Tukey, Ciminera and Heyse (1985) <DOI:10.2307/2530666> for general parametric models.

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asglht

Coerce object of class 'tukeytrend' to 'glht'

Description

Transfers output of functions `tukeytrendfit` and `tukeytrendformula` to `glht`.

Usage

```
asglht(object, df = "mean", ...)
```

Arguments

<code>object</code>	an object of class 'tukeytrend' as results from functions <code>tukeytrendfit</code> , or <code>tukeytrendformula</code>
<code>df</code>	defines whether and how/which degrees of freedom are passed to <code>glht</code> for the computation of multivariate quantile. Can be a single character string, that names a function ("mean", "min", ...) to summarize the individual models' degrees of freedom to a single degree of freedom, defaults to computing the mean df of all the input models. Setting <code>df=NULL</code> invokes that no degree of freedom is passed to <code>glht</code> , i.e. asymptotic inference by use of multivariate normal quantiles. You may also specify a single integer <code>df</code> in this argument, which is then passed to <code>glht</code> .
<code>...</code>	further arguments to be passed to <code>glht</code> , e.g. <code>alternative</code>

Details

Passes the elements `$mmm` from `tukeytrend` output to `glht`, argument `model`, and `$mlf` to argument `linfct`, computes summary statistics for the vector of degrees of freedom in `tukeytrend` and passes it to `glht`, argument `df`. All other arguments are passed via `...` to `glht`; be aware that mis-spelled arguments may be lost in `glht`.

Value

An object of class `glht`.

Examples

```
data(litter, package="multcomp")
dl <- litter
dl$dosen <- as.numeric(as.character(dl$dose))
fit <- lm(weight ~ dosen + number, data=dl)
ttlitter <- tukeytrendfit(fit, dose="dosen",
  scaling=c("ari", "ord", "log"))

# instead of transferring individual elements from the
# tukeytrend output into the arguments of glht , ...
# comp1 <- glht(model=ttlitter$mmm, linfct=ttlitter$mlf,
```

```
# df=round(mean(ttlitter$df), alternative="less")
# summary(comp1)

# ... you can just use:
comp2 <- asglht(ttlitter, alternative="less")
summary(comp2)
```

 combtt

Combine several pobjects of class 'tukeytrend'

Description

Concatenate several objects of class 'tukeytrend', for example to perform inference for multiple marginal models with different endpoints or including different covariates.

Usage

```
combtt(...)
```

Arguments

... names of tukeytrend objects (for multiple endpoint, or with different covariates), separated by comma

Value

Same structure as output of tuketrendfit, see [tukeytrendfit](#), [tukeytrendformula](#)

Examples

```
# Simulated data for normal and binomial response

dat <- data.frame(y = rnorm(20,0,1),
  succ=rbinom(n=20, size=10, prob=0.3),
  dose=rep(c(0,1,2,5,10), each=4))

# linear models with 3 different
# scalings of the dose variable

fitn <- lm(y~dose, data=dat)
ttn <- tukeytrendfit(fitn, dose="dose",
  scaling=c("ari", "ord", "arilog"))

# generalized linear models with 3
# different scalings of the dose variable
fitb <- glm(cbind(succ, 10-succ)~dose, data=dat, family=binomial)
ttb <- tukeytrendfit(fitb, dose="dose",
```

```

scaling=c("ari", "ord", "arilog"))

# concatenate the normal and binomial models
ttnb <- combtt(ttn, ttb)

# joint inference for the 6 regression slopes
summary(asglht(ttnb))

```

dosescalett

Rescaling dose variables acc. to Tukey et al. (1985)

Description

Add rescaled dose variables to a data.frame.

Usage

```

dosescalett(data, dose,
scaling = c("ari", "ord", "log", "arilog"),
sep = "", d0shift = 1)

```

Arguments

data	a data.frame
dose	a single character string, naming the (numeric) variable in data that is to be rescaled
scaling	A vector of character strings, naming the options for rescaling the variable specified in dose: "ari": no rescaling, "ord": ranks of dose levels, "log": log-transformed dose levels, "arilog": log-transformed dose levels, with interpolated dose score for 0, "high.vs.low": dose coerced to a factor, and only highest and lowest dose level retained, all others set NA, "treat": dose coerced to a factor, all levels retained with the option to apply multiple contrast tests to the treatment levels
sep	an optional separator for the names of the rescaled variables
d0shift	an optional factor, that is multiplied with the interpolated dose score for dose = 0 in option scaling="arilog"; ignored in all other options for rescaling

Value

A list containing

data	the input data.frame, with rescaled variables added as columns
transnam	a vector of names of the re-scaled variables
scaling	the scaling options, as input

lmer2lm	<i>Coerce objects of class lmerMod or lme to lm</i>
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Description

Coerce fitted mixed effect models of class lmerMod or lme to an lm-like object using a pseudo data approach (Ritz et al., 2017)

Usage

```
lmer2lm(lmerFit)
```

Arguments

lmerFit	a fitted object of class "lmerMod" or "lme", as can be obtained from packages "lme4" and "nlme"
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Details

The returned object resembles a fitted model of class "lm" such that its asymptotic representation corresponds to that of the originally fitted mixed model, see Ritz et al. (2017) for details.

Author(s)

Christian Ritz

References

Christian Ritz, Rikke Pilmann Laursen and Camilla Trab Damsgaard (2017): Simultaneous inference for multilevel linear mixed models - with an application to a large-scale school meal study: Appl. Statist. 66, Part 2, pp. 295-311.

tukeytrendfit	<i>Fit multiple marginal models by updating a given fitted model object with re-scaled dose variable.</i>
---------------	---

Description

Wrapper function to re-fit a given model after different rescalings of a single dose variable. The refitted models are combined into a list that is suitable as input to the multiple marginal model function of package multcomp, mmm.

Usage

```
tukeytrendfit(fit, dose,
  scaling = c("ari", "ord", "log", "arilog", "highvslow", "treat"),
  ctype = NULL, ddf = c("residual", "KR", "PB"), d0shift = 1)
```

Arguments

<code>fit</code>	A fitted model object, currently, the classes <code>lm</code> , <code>glm</code> , <code>lmer</code> , and <code>nlme</code> are supported. The left hand side of the models formula must contain a single (numeric) variable that is treated as a dose variable in Tukeys trend test. It may contain further variables that are not changed when refitting the model.
<code>dose</code>	A single character string, naming a numeric variable in the models formula. This variable is rescaled acc. to the options in <code>scaling</code> and the model in <code>fit</code> is then refitted with the rescaled dose variable.
<code>scaling</code>	A vector of character strings, naming the options for rescaling the variable specified in <code>dose</code> : <code>"ari"</code> : no rescaling, <code>"ord"</code> : ranks of dose levels, <code>"log"</code> : log-transformed dose levels, <code>"arilog"</code> : log-transformnd dose levels, with interpolated dose score for 0, <code>"high.vs.low"</code> : dose coerced to a factor, and only highest and lowest dose level retained, all others set NA, <code>"treat"</code> : dose coerced to a factor, all levels retained with the option to apply multiple contrast tests to the treatment levels
<code>ctype</code>	optional character string naming a contrast type for multiple comparisons between dose levels, when <code>scaling="treat"</code> . Options are <code>"Dunnett"</code> , <code>"William"</code> etc., see <code>?contrMat</code> in package <code>multcomp</code> . Argument <code>ctype</code> is ignored if <code>scaling</code> does not involve option <code>"treat"</code> .
<code>ddf</code>	single character string, defining the option for the degree of freedom in inference after model fitting. By default, <code>"residual"</code> degrees of freedom will be used for all models. <code>"KR"</code> : For models of class <code>"lmerMod"</code> (fitted by <code>"lmer"</code> from package <code>"lme4"</code>), Kenward-Roger degrees of freedom can be computed (based on methods from package <code>"pbkrtest"</code>); <code>"PB"</code> : For models of class <code>"lme"</code> (fitted by <code>"lme"</code> from package <code>"nlme"</code>), containment degrees of freedom as defined by Pinheiro and Bates can be extracted.
<code>d0shift</code>	an optional factor, that is multiplied with the interpolated dose score for <code>dose = 0</code> in option <code>scaling="arilog"</code> ; ignored in all other options for rescaling

Value

A list with elements

<code>mmm</code>	a list of fitted models, after rescaling the dose variable
<code>mlf</code>	a list of matrices defining a linear functions of model parameters for each model in <code>mmm</code> , defining the parameter of interest for inference in function <code>mlf</code> and <code>glht</code>
<code>df</code>	a vector of degrees of freedom, one for each model in <code>mmm</code>

and information of the model `typ` and call of the initial model

Author(s)

Frank Schaarschmidt and Christian Ritz (providing internal functions to interface objects of class `"lmerMod"` and `"lme"`)

References

Tukey JW, Ciminera JL, Heyse JF (1985). Testing the statistical certainty of a response to increasing doses of a drug. *Biometrics* 41(1), 295-301.

Pipper CB, Ritz C, Bisgaard H (2012). A versatile method for confirmatory evaluation of the effects of a covariate in multiple models. *JRSSC - Applied Statistics* 61, 315-326.

See Also

Functions `glht` and `mmm` in package `multcomp`; [tukeytrendformula](#) for starting with a model formula and `data.frame`; [combt](#) for concatenating several objects resulting from this function, e.g. for multiple endpoints or with different covariates; [asglht](#) for direct conversion to a `glht`-object

Examples

```
data(litter, package="multcomp")

# variable of interest is 'dose', but
# we may want to include 'number' as a covariate

dl <- litter
dl$dosen <- as.numeric(as.character(dl$dose))

fit <- lm(weight ~ dosen + number, data=dl)

# as it is unclear on which scale of dosage,
# dose-response-shape is met best, we may try different
# re-scalings of the dose variable, including Dunnett- type multiple contrast test

ttlitter <- tukeytrendfit(fit, dose="dosen",
  scaling=c("ari", "ord", "log", "treat"), ctype="Dunnett")

summary(asglht(ttlitter))
```

tukeytrendformula *Fit multiple marginal models with differently re-scaled dose variable.*

Description

Wrapper function to fit a given model after different rescalings of a single dose variable. The fitted models are combined into a list that is suitable as input to the multiple marginal model function of package `multcomp`, `mmm`.

Usage

```
tukeytrendformula(formula, data, model = "lm", dose,
  scaling = c("ari", "ord", "log", "arilog", "treat", "treatHL"),
  ctype = NULL, ddf = c("residual", "KR", "PB"), d0shift = 1, ...)
```

Arguments

formula	formula object suitable for the model function specified in model, the left hand side of the formula should contain at least one (numeric) variable, that is to be re-scaled in the model fits
data	data.frame containing the variables of interest
model	character string, naming the function for model fitting, currently "lm", "glm", "lmer", "lme" are supported
dose	A single character string, naming a numeric variable in the models formula. This variable is rescaled acc. to the options in scaling and the model in fit is then refitted with the rescaled dose variable.
scaling	A vector of character strings, naming the options for rescaling the variable specified in dose: "ari": no rescaling, "ord": ranks of dose levels, "log": log-transformed dose levels, "arilog": log-transformed dose levels, with interpolated dose score for 0, "high.vs.low": dose coerced to a factor, and only highest and lowest dose level retained, all others set NA, "treat": dose coerced to a factor, all levels retained with the option to apply multiple contrast tests to the treatment levels
ctype	optional character string naming a contrast type for multiple comparisons between dose levels, when scaling="treat". Options are "Dunnett", "William" etc., see ?contrMat in package multcomp. Argument ctype is ignored if scaling does not involve option "treat".
ddf	single character string, defining the option for the degree of freedom in inference after model fitting. By default, "residual" degrees of freedom will be used for all models. "KR": For models of class "lmerMod" (fitted by "lmer" from package "lme4"), Kenward-Roger degrees of freedom can be computed (based on methods from package "pbkrtest"); "PB": For models of class "lme" (fitted by "lme" from package "nlme"), containment degrees of freedom as defined by Pinheiro and Bates can be extracted.
d0shift	an optional factor, that is multiplied with the interpolated dose score for dose = 0 in option scaling="arilog"; ignored in all other options for rescaling
...	arguments passed to the model fitting function named in model

Value

A list with elements

mmm	a list of fitted models, after rescaling the dose variable
m1f	a list of matrices defining a linear functions of model parameters for each model in mmm, defining the parameter of interest for inference in function m1f and glht
df	a vector of degrees of freedom, one for each model in mmm

and information of the model type and call of the initial model

Author(s)

Frank Schaarschmidt and Christian Ritz (providing internal functions to interface objects of class "lmerMod" and "lme")

References

Tukey JW, Ciminera JL, Heyse JF (1985). Testing the statistical certainty of a response to increasing doses of a drug. *Biometrics* 41(1), 295-301.

Pipper CB, Ritz C, Bisgaard H (2012). A versatile method for confirmatory evaluation of the effects of a covariate in multiple models. *JRSSC - Applied Statistics* 61, 315-326.

Examples

```
data(litter, package="multcomp")

# compare

dl <- litter
dl$dosen <- as.numeric(as.character(dl$dose))

ttlitter <- tukeytrendformula(weight ~ dosen + number, data=dl, model="lm", dose="dosen",
  scaling=c("ari", "ord", "log", "treat"), ctype="Dunnett")

summary(asglht(ttlitter))
```

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