Package ‘lsmeans’

August 28, 2017

Type Package
Title Least-Squares Means
Version 2.27-2
Date 2017-08-28
Depends estimability, methods, R (>= 3.2)
Suggests car, lattice, MCMCpack, mediation, multcompView, ordinal,
   pbkrtest (>= 0.4-1), CARBayes, coxme, gee, geepack, glmmADMB,
   lme4.0, lme4, lmerTest (>= 2.0.32), MASS, MCMCglmm, nnet, pscl,
   rsm, rstan, rstanarm, survival
Imports graphics, stats, utils, nlme, coda (>= 0.17), multcomp, plyr,
   mvtnorm, xtable (>= 1.8-2)
Additional_repositories http://glmmadmb.r-forge.r-project.org/repos,
   http://lme4.0.r-forge.r-project.org/repos
BugReports https://github.com/rvlenth/lsmeans/issues
LazyData yes
ByteCompile yes
Description Obtain least-squares means for many linear, generalized linear,
   and mixed models. Compute contrasts or linear functions of least-squares
   means, and comparisons of slopes. Plots and compact letter displays.
License GPL-2 | GPL-3
NeedsCompilation no
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Repository CRAN
Date/Publication 2017-08-28 14:41:31 UTC
**R topics documented:**

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**Description**

This package provides methods for obtaining so-called least-squares means for factor combinations in a variety of fitted linear models. It can also compute contrasts or linear combinations of these least-squares means, (several standard contrast families are provided), and in addition can estimate and contrast slopes of trend lines. Popular adjustments for multiple-comparisons are provided, as well as graphical ways of displaying the results.

---

**Details**

- **Package**: lsmeans
- **Type**: Package
- **License**: GPL-2
- **Other information**: See DESCRIPTION
Overview

Concept  Least-squares means (see Searle et al. 1980, who prefer the term “predicted marginal means” (PMM)) are popular for summarizing linear models that include factors. For balanced experimental designs, they are just the marginal means. For unbalanced data, they in essence estimate what you would have observed that the data arisen from a balanced experiment.

Reference grids  The implementation in lsmeans relies on our own concept of a reference grid, which is an array of factor and predictor levels. Predictions are made on this grid, and least-squares means are defined as averages of these predictions over zero or more dimensions of the grid. The function ref.grid explicitly creates a reference grid (ref.grid object) that can subsequently be used to obtain least-squares means. The update method is used to change its properties.

Our reference-grid framework expands slightly upon Searle et al.’s definitions of PMMs, in that it is possible to include multiple levels of covariates in the grid.

Models supported  Many linear models are supported by the package, including lm, glm, aovlist, and mlm in the stats package, as well as fitted-model objects from several contributed packages including nime, lme4, survival, and geepack. The help page for models provides more details, including, in some cases, additional ref.grid arguments that might affect the subsequent analysis. Also, some models require other packages be installed in order to obtain all the available features.

Least-squares means  The lsmeans function computes least-squares means given a ref.grid object or a fitted model, and a specification indicating what factors to include. The lstrends function creates the same sort of results for estimating and comparing slopes of fitted lines. Both return an lsmobj object very much like a reference grid, but with possibly fewer factors involved.

Summaries and analysis  The summary method may be used to display a ref.grid or an lsmobj. Special-purpose summaries are available via confint and test, the latter of which can also do a joint test of several estimates. The user may specify by variables, multiplicity-adjustment methods, confidence levels, etc., and if a transformation or link function is involved, may reverse-transform the results to the response scale.

Contrasts and comparisons  The contrast method is used to obtain contrasts among the estimates; several standard contrast families are available such as deviations from the mean, polynomial contrasts, and comparisons with one or more controls. Another lsmobj object is returned, which can be summarized or further analyzed. For convenience, a pairs method is provided for the case of pairwise comparisons. Related to this is the cld method, which provides a compact letter display for grouping pairs of means that are not significantly different. cld requires the multcompView package.

Graphs  The plot method will display side-by-side confidence intervals for the estimates, and/or ‘comparison arrows’ whereby the significance of pairwise differences can be judged by how much they overlap. The lsmip function displays estimates like an interaction plot, multi-paneled if there are by variables. These graphics capabilities require the lattice package be installed.

multcomp interface  The as.glht function and glht method for lsmobjs provide an interface to the glht function in the multcomp package, thus providing for more exacting simultaneous estimation or testing. The package also provides an ls method that works as an alternative to mcp in a call to glht.


**Additional information**

Examples and discussion are available via vignette("using-1smeans", package="lsmeans"). Some features of the lsmeans require (or are enhanced by) additional packages that are loaded when needed. Since they are not “required” packages, they are not automatically installed with lsmeans. We highly recommend that users also install the following packages: multcomp (if cld, glht, or as.glht are to be used), multcompView (for cld), lattice (for plot and lsmip), and lmerTest or pbkrtest (for models fitted by the lme4 package).

Starting with lsmeans version 2, a new object framework based on reference grids is used that increases flexibility and provides for extending its capabilities to additional model objects. Use vignette("1smeans-changes") for information on the user impact of these changes.

It is possible to write your own interfaces for models not yet supported by lsmeans. See the help page extending-1smeans and vignette("extending") for details on how to do this.

**Author(s)**

Russell V. Lenth (author), Maxime Hervé (contributor)

Maintainer: Russ Lenth <russell-lenth@uiowa.edu>

**References**


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<table>
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<th>auto.noise</th>
<th>Auto Pollution Filter Noise</th>
</tr>
</thead>
</table>

**Description**

Three-factor experiment comparing pollution-filter noise for two filters, three sizes of cars, and two sides of the car.

**Usage**

    auto.noise

**Format**

A data frame with 36 observations on the following 4 variables.

- **noise** Noise level in decibels - a numeric vector.
- **size** The size of the vehicle - an ordered factor with levels S, M, L.
- **type** Type of anti-pollution filter - a factor with levels Std and Octel
- **side** The side of the car where measurement was taken – a factor with levels L and R.
Details

The data are from a statement by Texaco, Inc., to the Air and Water Pollution Subcommittee of the Senate Public Works Committee on June 26, 1973. Mr. John McKinley, President of Texaco, cited an automobile filter developed by Associated Octel Company as effective in reducing pollution. However, questions had been raised about the effects of filters on vehicle performance, fuel consumption, exhaust gas back pressure, and silencing. On the last question, he referred to the data included here as evidence that the silencing properties of the Octel filter were at least equal to those of standard silencers.

Source

The dataset was imported from the Data and Story Library - http://lib.stat.cmu.edu/DASL/Datafiles/airpollutionfiltersdat.html (sic). However, the factor levels were assigned meaningful names, and the observations were sorted in random order as if this were the run order of the experiment.

References


Examples

```r
require(lsmeans)
noise.lm <- lm(noise ~ size * type * side, data = auto.noise)

# Interaction plot of predictions
lsmip(noise.lm, type ~ size | side)

# Confidence intervals
plot( lsmeans(noise.lm, ~ size | side*type) )
```

---

cld

Compact letter display of pairwise comparisons

Description

Extract and display information on all pairwise comparisons of least-squares means.

Usage

```r
## S3 method for class 'ref.grid'
cld(object, details = FALSE, sort = TRUE, by, alpha = 0.05,
     Letters = c("1234567890", LETTERS, letters), reversed = FALSE, ...)

## S3 method for class 'lsm.list'
cld(object, ..., which = 1)
```
Arguments

object     An object of class ref.grid
details    Logical value determining whether detailed information on tests of pairwise
            comparisons is displayed
sort       Logical value determining whether the LS means are sorted before the compar-
            isons are produced. When sort is TRUE, the results are displayed in increasing
            order if reversed is FALSE (the default), or in decreasing order if reversed is
            TRUE.
by         Character value giving the name or names of variables by which separate fami-
            lies of comparisons are tested. If NULL, all means are compared. If missing, and
            a by variable was used in creating object, it is used as the by variable in cld.
alpha      Numeric value giving the significance level for the comparisons
Letters    Character vector of letters to use in the display. Any strings of length greater
            than 1 are expanded into individual characters
reversed   Logical value (passed to multcompLetters in the multcompView package.) If
            TRUE, the order of use of the letters is reversed. In addition, if both sort and
            reversed are TRUE, the sort order of results is reversed.
...        Arguments passed to contrast (for example, an adjust method)
which      When object is a list, this determines which element is analyzed.

Details

This function uses the Piepho (2004) algorithm (as implemented in the multcompView package) to
generate a compact letter display of all pairwise comparisons of least-squares means. The function
obtains (possibly adjusted) \( P \) values for all pairwise comparisons of means, using the contrast
function with method = "pairwise". When a \( P \) value exceeds alpha, then the two means have at
least one letter in common.

Value

When details == FALSE, an object of class summary.ref.grid (which inherits from data.frame)
showing the summary of LS means with an added column named .groups with the cld information.
When details == TRUE, a list the object just described, as well as the summary of the contrast
results showing each comparison, its estimate, standard error, \( t \) ratio, and adjusted \( P \) value.

Note

This function requires the multcompView package to be installed. Otherwise an error message is
produced.

Author(s)

Russell V. Lenth

References

Hans-Peter Piepho (2004) An algorithm for a letter-based representation of all pairwise compar-
contrast

See Also
cld in the multcomp package

Examples

```r
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.lsm <- lsmeans(warp.lm, ~ tension | wool)
cld(warp.lsm)  # implicitly uses by = "wool"
cld(warp.lsm, by = "tension")  # overrides implicit 'by'

# Mimic grouping bars and compare all 6 means
cld(warp.lsm, by = NULL, Letters = "|||\|\|\|\|\|", alpha = .01)
```

Description

These methods provide for analyses of ref.grid objects, or follow-up analyses of lsmobj objects: Contrasts, pairwise comparisons, tests, and confidence intervals.

Usage

```r
## S3 method for class 'ref.grid'
contrast(object, method = "eff", interaction = FALSE,
         by, offset = NULL, name = "contrast",
         options = getOption("lsmeans")$contrast, adjust, ...)

## S3 method for class 'lsm.list'
contrast(object, ..., which = 1)

## S3 method for class 'ref.grid'
test(object, null = 0, joint = FALSE,
      verbose = FALSE, rows, by, ...)

## S3 method for class 'ref.grid'
confint(object, parm, level = 0.95, ...)

## S3 method for class 'ref.grid'
pairs(x, reverse = FALSE, ...)

## S3 method for class 'ref.grid'
coef(object, ...)
```

Arguments

```r
object, x An object of class "ref.grid" or its extension, "lsmobj".
```
<table>
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<th>Description</th>
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<tr>
<td>method</td>
<td>Character value giving the root name of a contrast method (e.g. &quot;pairwise&quot;). Alternatively, a named list of contrast coefficients that must each conform to the number of least-squares means in each by group. This is just like the contr argument in <code>lsmeans</code>. To identify the available methods, see <code>ls(&quot;package:lsmeans&quot;, pat=&quot;.1smc&quot;)</code>. You may define your own <code>.1smc</code> function and use its root name as method. If interaction is of character type, this argument is ignored.</td>
</tr>
<tr>
<td>interaction</td>
<td>Character vector or logical value. In multi-factor situations with interaction = FALSE, the factor combinations are treated as levels of a single “uber-factor”, and the contrast specified in method is applied to it. Otherwise, interaction contrasts are computed: Contrasts are generated for each factor separately, one at a time; and these contrasts are applied to the object (the first time around) or to the previous result (subsequently). (Any factors specified in by are skipped.) The final result comprises contrasts of contrasts, or, equivalently, products of contrasts for the factors involved. Processing is done in the order of appearance in <code>object@levels</code>. With interaction = TRUE, method (if specified as character) is used for each contrast. If interaction is a character vector, the elements specify the respective contrast method(s); they are recycled as needed.</td>
</tr>
<tr>
<td>by</td>
<td>Character names of variable(s) to be used for “by” groups. The contrasts or joint tests will be evaluated separately for each combination of these variables. If object was created with by groups, those are used unless overridden. Use by = NULL to use no by groups at all.</td>
</tr>
<tr>
<td>offset</td>
<td>Numeric vector of the same length as each by group. These values are added to their respective linear estimates. This argument is ignored when interaction is not FALSE.</td>
</tr>
<tr>
<td>name</td>
<td>Name to use to label the contrasts in table headings or subsequent contrasts of the returned object. This argument is ignored when interaction is not FALSE.</td>
</tr>
<tr>
<td>options</td>
<td>If non-NULL, a named list of arguments to pass to update, just after the object is constructed.</td>
</tr>
<tr>
<td>adjust</td>
<td>Method to use for adjusting P values. This is passed to summary. This argument is available in contrast for historical reasons; but it is better style to specify the adjustment method, along with other testing options such as side, as part of options.</td>
</tr>
<tr>
<td>joint</td>
<td>Logical value. If FALSE, the arguments are passed to summary with infer=c(FALSE,TRUE). If TRUE, a joint test of the hypothesis L beta = null is performed, where L is <code>object@linfct</code> and beta is the vector of fixed effects estimated by <code>object@betahat</code>. This will be either an F test or a chi-square (Wald) test depending on whether degrees of freedom are available.</td>
</tr>
<tr>
<td>rows</td>
<td>Integer values. The rows of L to be tested in the joint test. If missing, all rows of L are used. If not missing, by variables are ignored.</td>
</tr>
<tr>
<td>null</td>
<td>Numeric value specifying the null value(s) being tested against. It may be either a single value, in which case it is used as the null value for all linear functions under test; or a numeric vector of length equal to the number of linear functions.</td>
</tr>
<tr>
<td>parm</td>
<td>This is ignored, but it is a required argument of the generic confint method.)</td>
</tr>
</tbody>
</table>
verbose Logical value. If TRUE and joint==TRUE, a table of the effects being tested is printed.

level Numeric value of the desired confidence level.

which When object is a list of lsmobj objects, this specifies which member of the list is analyzed.

reverse Logical value determining whether "pairwise" or "revpairwise" pairwise comparisons are generated.

... Additional arguments passed to summary or to a contrast function.

Details

Though contrast is ordinarily used to create true contrasts (whose coefficients sum to zero), it may be used to estimate any linear function of the LS means; and offset expands this capability further by allowing additive constants. pairs is equivalent to contrast with method = "pairwise".

confint and test (when joint==FALSE) are equivalent to calling summary with infer=c(TRUE, FALSE) and infer=c(FALSE, TRUE), respectively.

When using test to do a joint test of L beta = null, an error is thrown if any row of L is non-estimable. It is permissible for the rows of L to be linearly dependent as long as null == 0; a reduced set of contrasts is tested. Linear dependence and nonzero null cause an error.

Value

contrast and pairs return an object of class "lsmobj", which is an extension of "ref.grid". Consequently, they may be used as arguments to other "lsmobj" or "ref.grid" methods. The user may, for example, compute contrasts of contrasts, or re-summarize a set of confidence intervals with a different by grouping or confidence level. The "grid" for the returned value is simply the set of variables that identify the results. For example, contrast's return value is a reference grid for one factor named contrast.

cconfint and test (when Joint==FALSE) return an object of class summary.ref.grid. When JOINT==TRUE, test returns a "summary.ref.grid" object (extends "data.frame") with the test statistic, degrees of freedom, and P value for each by group.

When object is the result of a call to contrast or pairs, the coef method returns a data.frame. The initial columns are the factor combinations that were contrasted (i.e. the grid for the object originally specified in the call to contrast), and the remaining columns (named c.1, c.2, ...) contain the contrast coefficients that were applied to the corresponding predictions. If object was not produced via contrast, NULL is returned, along with a message.

Author(s)

Russell V. Lenth

See Also

Additional "lsmobj" methods having their own help pages are cld and glht. Also, the summary and other methods for "ref.grid" objects also work for "lsmobj" objects.
Examples

require(lsmeans)
warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)
warp.lsm <- lsmeans(warp.lm, ~ tension | wool)

# Polynomial contrasts of tension, by wool
(warp.pl <- contrast(warp.lsm, "poly", name = "order"))
# Same results with a different adjustment
summary(warp.pl, adjust = "fdr")

# Jointly test the tension effects for each wool
test(warp.pl, joint = TRUE, by = "wool")

# Compare the two contrasts for each order
contrast(warp.pl, "revpairwise", by = "order")

# User-provided contrasts, ignoring the previous by grouping
contrast(warp.lsm,
  list(c1=c(1,0,0,-1,0,0), c2=c(1,1,1,-1,-1,-1)/3),
  by = NULL)

# Compare consecutive tension*wool comb's as treatment with 6 levels
contrast(warp.lsm, "consec", by = NULL)

# Interaction contrasts (comparisons of linear and quadratic contrasts)
(int.con <- contrast(warp.lsm, interaction = c("poly", "consec"), by = NULL))

# See the contrast coefficients used by the previous call
coeff(int.con)

---

feedlot                Feedlot data

Description

This is an unbalanced analysis-of-covariance example, where one covariate is affected by a factor.
Feeder calves from various herds enter a feedlot, where they are fed one of three diets. The weight
of the animal at entry is the covariate, and the weight at slaughter is the response.

Usage

data(feedlot)

Format

A data frame with 67 observations on the following 4 variables.

herd  a factor with levels 9 16 3 32 24 31 19 36 34 35 33, designating the herd that a feeder calf
came from.
diet a factor with levels Low Medium High: the energy level of the diet given the animal.
swt a numeric vector: the weight of the animal at slaughter.
ewt a numeric vector: the weight of the animal at entry to the feedlot.

Details

The data arise from a Western Regional Research Project conducted at New Mexico State University. Calves born in 1975 in commercial herds entered a feedlot as yearlings. Both diets and herds are of interest as factors. The covariate, ewt, is thought to be dependent on herd due to different genetic backgrounds, breeding history, etc. The levels of herd ordered to similarity of genetic background.

Note: There are some empty cells in the cross-classification of herd and diet.

Source


Examples

require(lsmeans)
feedlot.lm <- lm(swtn ~ ewt + herd * diet, data = feedlot)

# Obtain LS-means with a separate reference value of ewt for each
# herd. This reproduces the last part of Table 2 in the reference
lsmeans(feedlot.lm, ~ diet | herd, cov.reduce = ewt ~ herd)

fiber

Fiber data

Description

Fiber data from Montgomery Design (8th ed.), p.656 (Table 15.10). Useful as a simple analysis-of-covariance example.

Usage

fiber

Format

A data frame with 15 observations on the following 3 variables.
machine a factor with levels A B C. The primary factor of interest.
strength a numeric vector. The response variable.
diameter a numeric vector. A covariate.
Details

The goal of the experiment is to compare the mean breaking strength of fibers produced by the three machines. When testing this, the technician also measured the diameter of each fiber, and this measurement may be used as a concomitant variable to improve precision of the estimates.

Source


Examples

```r
require(lsmeans)
fiber.lm <- lm(strength ~ diameter + machine, data=fiber)
ref.grid(fiber.lm)

# Covariate-adjusted means and comparisons
lsmeans(fiber.lm, pairwise ~ machine)
```

---

**glht**  
**lsmeans support for glht**

Description

These functions and methods provide an interface between lsmeans and the glht function for simultaneous inference in the multcomp package.

Usage

```r
## S3 method for class 'ref.grid'
as.glht(object, ...)
## S3 method for class 'lsm.list'
as.glht(object, ..., which = 1)

## S3 method for class 'glht.list'
coef(object, ...)
## S3 method for class 'glht.list'
confint(object, ...)
## S3 method for class 'glht.list'
plot(x, ...)
## S3 method for class 'glht.list'
summary(object, ...)
## S3 method for class 'glht.list'
vcov(object, ...)

lsm(...)
pmm(...)
```
Arguments

- `object, x` An object of the required class.
- `which` Numeric index of which element of the lsm.list to use.
- `...` Additional arguments to other methods.

Details

`lsmeans` (and `pmmeans`, which is identical) are meant to be called only from "glht" as its second (`linfct`) argument. It works similarly to `mcp` except with `specs` (and optionally by and contr arguments) provided as in a call to `lsmeans` or `pmmeans`.

When there is a non-NULL by variable (either explicitly or implicitly), each “by” group is passed separately to `glht` and returned as a list of "glht" objects. For convenience, this is classed as "glht.list", and appropriate methods `coef`, `confint`, `plot`, `summary`, and `vcov` are provided.

Value

`as.glht` returns an object of class `glht`, or of class `glht.list` if by is non-NULL. The latter is simply a list of `glht` objects, and the provided methods `coef`, `confint`, `plot`, `summary`, and `vcov` simply `lapply` the corresponding methods for class "glht".

Note

There is also a `glht` method for class `ref.grid`, but it is far preferable to use `as.glht` instead, as its model argument is redundant.

Author(s)

Russell V. Lenth

See Also

`lsmeans, glht`

Examples

```r
require(lsmeans)
require(multcomp)

warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)

# Using 'lsm'
summary(glht(warp.lm, lsm(pairwise ~ tension | wool)))

# Same, but using an existing 'lsmeans' result
warp.lsmobj <- lsmeans(warp.lm, ~ tension | wool)
summary(as.glht(pairs(warp.lsmobj)))

# Same contrasts, but treat as one family
summary(as.glht(pairs(warp.lsmobj), by = NULL))
```
Least-squares means (or predicted marginal means)

Description

Compute least-squares means (predicted marginal means) for specified factors or factor combinations in a linear model, and optionally comparisons or contrasts among them.

Usage

```r
## S3 method for class 'character'
lsmeans(object, specs, ...)
## (used when 'specs' is 'character')

## S3 method for class 'character.ref.grid'
lsmeans(object, specs, by = NULL,
        fac.reduce = function(coefs) apply(coefs, 2, mean), contr,
        options = getOption("lsmeans")$lsmeans, weights, trend, ...)
## (used when 'object' is a 'ref.grid' and 'specs' is 'character')

## S3 method for class 'list'
lsmeans(object, specs, ...)
## (used when 'specs' is a 'list')

## S3 method for class 'formula'
lsmeans(object, specs, contr.list, trend, ...)
## (used when 'specs' is a 'formula')

lstrends(model, specs, var, delta.var = 0.01 * rng, data,
         transform = c("none", "response"), ...)

lsmobj(bhat, V, levels, linfct, df = NA, post.beta = matrix(NA), ...)

pmmmeans(...)
pmtrends(...)
pmmobj(...)
```

Arguments

- **object**: An object of class ref.grid; or a fitted model object that is supported, such as the result of a call to lm or lmer. Many fitted-model objects are supported; see link(models) for details.
- **specs**: A character vector specifying the names of the predictors over which LS-means are desired. specs may also be a formula or a list (optionally named) of valid specs. Use of formulas is described in the Details section below.
- **by**: A character vector specifying the names of predictors to condition on.
lsmeans

fac.reduce  A function that combines the rows of a matrix into a single vector. This implements the “marginal averaging” aspect of least-squares means. The default is the mean of the rows. Typically if it is overridden, it would be some kind of weighted mean of the rows. If fac.reduce is nonlinear, bizarre results are likely, and LS means will not be interpretable. If the weights argument is non-missing, fac.reduce is ignored.

contr  A list of contrast coefficients to apply to the least-squares means – or the root name of an .lsmc function that returns such coefficients. In addition, contr = "cld" is an alternative way to invoke the cld function. See contrast for more details on contrasts. NOTE: contr is ignored when specs is a formula.

contr.list  A named list of lists of contrast coefficients, as for contr. This is used only in the formula method; see Details below.

options  If non-NULL, a named list of arguments to pass to update, just after the object is constructed.

weights  Numeric vector, numeric matrix, or character string specifying weights to use in averaging predictions. If a vector, its length must equal the number of predictions to be averaged to obtain each least-squares mean. If a matrix, each row of the matrix is used in turn, wrapping back to the first row as needed. When in doubt about what is being averaged (or how many), first call with weights = "show.levels".

If a string, it should partially match one of the following:

"equal"  Use an equally weighted average.

"proportional"  Weight in proportion to the frequencies (in the original data) of the factor combinations that are averaged over.

"outer"  Weight in proportion to each individual factor’s marginal frequencies. Thus, the weights for a combination of factors are the outer product of the one-factor margins.

"cells"  Weight according to the frequencies of the cells being averaged.

"flat"  Give equal weight to all cells with data, and ignore empty cells.

"show.levels"  This is a convenience feature for understanding what is being averaged over. Instead of a table of LS means, this causes the function to return a table showing the levels that are averaged over, in the order they appear.

Outer weights are like the 'expected' counts in a chi-square test of independence, and will yield the same results as those obtained by proportional averaging with one factor at a time. All except "cells" uses the same set of weights for each mean. In a model where the predicted values are the cell means, cell weights will yield the raw averages of the data for the factors involved. Using "flat" is similar to "cells", except nonempty cells are weighted equally and empty cells are ignored.

Note: If a nested structure exists (see the nests argument in ref.grid), then averaging is done separately over each nesting group; thus, these groups are potentially of different sizes. Accordingly, it is unsafe to specify numerical weights.

Note: If weights were used in fitting the model, then weight totals are used in place of frequencies in these schemes.
lsmeans

If weights is used, fac.reduce is ignored.

trend  Including this argument is an alternative way of calling lstrends with trend as its var argument and object as its model.

model   A supported model object (not a ref.grid).

var     Character giving the name of a variable with respect to which a difference quotient of the linear predictors is computed. In order for this to be useful, var should be a numeric predictor that interacts with at least one factor in specs. Then instead of computing least-squares means, we compute and compare the slopes of the var trend over levels of the specified other predictor(s). As in least-squares means, marginal averages are computed when some variables in the reference grid are excluded for the specification.

The user may specify some monotone function of one variable, e.g., var = "log(dose)". If so, the chain rule is applied. Note that, in this example, if model contains log(dose) as a predictor, we will be comparing the slopes estimated by that model, whereas specifying var = "dose" would perform a transformation of those slopes.

delta.var The value of h to use in forming the difference quotient \((f(x+h) - f(x))/h\). Changing it (especially changing its sign) may be necessary to avoid numerical problems such as logs of negative numbers. The default value is 1/100 of the range of var over the dataset.

data    As in ref.grid, you may use this argument to supply the dataset used in fitting the model, for situations where it is not possible to reconstruct the data. Otherwise, leave it missing.

transform In lstrends, if object has a response transformation, then specifying transform = "response" will cause lstrends to calculate the trends after back-transforming to the response scale. This is done using the chain rule, and standard errors are estimated via the delta method. With transform = "none" (the default), the trends are calculated on the scale of the linear predictor, without back-transforming it. This argument works similarly to the transform argument in ref.grid (but without a "log" option), in that the returned object is re-gridded to the new scale (see also regrid).

bhat     Numeric. Vector of regression coefficients.

V        Square matrix. Covariance matrix of bhat

levels Named list or vector. Levels of factor(s) that define the estimates defined by linfct. If not a list, we assume one factor named "level"

linfct   Matrix. Linear functions of bhat for each combination of levels

df       Numeric or function with arguments \((x, dfargs)\). If a number, that is used for the degrees of freedom. If a function, it should return the degrees of freedom for \(\sum(x*bhat)\); if additional parameters are needed, include them in \(...\) as dfargs (not abbreviated).

post.beta Matrix whose columns comprise a sample from the posterior distribution of the regression coefficients (so that typically, the column averages will be bhat). A 1 x 1 matrix of NA indicates that such a sample is unavailable.
Additional arguments passed to other methods or to \texttt{ref.grid}. For example, \texttt{vcov} may be used to override the default covariance estimate, and some models allow additional options. Some models require data to be given explicitly. See the help pages for \texttt{ref.grid} and \texttt{models}. In addition, if the model formula contains references to variables that are not predictors, you must provide a \texttt{params} argument with a list of their names; see the example below for \texttt{Oatssq.lm}.

**Details**

Least-squares means (also called predicted marginal means) are predictions from a linear model over a \textit{reference grid}, or marginal averages thereof. They have been popularized by \texttt{SAS} (SAS Institute, 2012). The \texttt{ref.grid} function identifies/creates the reference grid upon which \texttt{lsmeans} is based.

For those who prefer the term “predicted marginal means”, courtesy wrappers \texttt{pmmeans}, \texttt{pmtrends}, and \texttt{pmmobj} are provided that behave identically to those that start with \texttt{ls}, except that estimates are relabeled accordingly (e.g., \texttt{lsmean} becomes \texttt{pmmean}).

If \texttt{specs} is a formula, it should be of the form \texttt{~ specs, ~ specs | by, contr ~ specs, or contr ~ specs | by}. The formula is parsed and the variables therein are used as the arguments \texttt{specs}, \texttt{by}, and \texttt{contr} as indicated. The left-hand side is optional, but if specified it should be the name of a contrast family (e.g., \texttt{pairwise}) or of a sub-list of \texttt{contr.list}. Operators like \texttt{*} or \texttt{:} are necessary to delineate names in the formulas, but otherwise are ignored.

In the special case where the mean (or weighted mean) of all the predictions is desired, specify \texttt{specs} as \texttt{~ 1} or \texttt{"1"}.

A number of standard contrast families are provided. They can be identified as functions having names ending in \texttt{.lsmc} – use

\begin{verbatim}
ls("package:lsmeans", pat=".lsmc")
\end{verbatim}

to list them. See the documentation for \texttt{pairwise.lsmc} and its siblings for details. You may write your own \texttt{.lsmc} function for custom contrasts.

The function \texttt{lsmobj} may be used to construct an object just like one returned by \texttt{lsmeans} from user-specified coefficients, covariance matrix, levels (or row labels), linear functions for each row, and degrees of freedom. After the object is constructed, it is \texttt{updated} with any additional arguments in \ldots.

**Value**

When \texttt{specs} is a character vector or one-sided formula, an object of class \texttt{lsmobj}. A number of methods are provided for further analysis, including \texttt{summary}, \texttt{confint}, \texttt{test}, \texttt{contrast}, \texttt{pairs}, and \texttt{cld}.

When \texttt{specs} is a list or a formula having a left-hand side, the return value is an \texttt{lsm.list} object, which is simply a list of \texttt{lsmobj} objects. Methods for \texttt{lsm.list} objects are the same as those for \texttt{lsmobj}, but they apply to only one member of the list, determined by its which argument.

**Side effect:** When \texttt{object} is a model, a reference grid is constructed and it is saved as \texttt{.Last.ref.grid} in the user’s environment (unless this is disabled via \texttt{\'lsm.option(save.ref.grid = FALSE)\'}}). This makes it possible to check what reference grid was used, or to use it as the object in future \texttt{lsmeans} calls (and bypass reconstructing it). Similarly, \texttt{lstrends} also saves its reference grid (but for predicting difference quotients) as \texttt{.Last.ref.grid}. 


Note

If the model formula contains variables that are not predictors (e.g., degree of a polynomial, knots for a spline, etc.), you must add a `params` argument to the call.

Note

While using `specs` as a two-sided formula or a list is a convenient way to get a lot of results with minimal effort, it can also create confusion when additional arguments are provided, because not all arguments may be applied to all the results produced (see examples). Thus, the safer route is to do things incrementally.

Note

`lsmeans` and its relatives can produce fatal errors or incorrect results with models containing splines (e.g., `ns`) and other smoothers because the required information to reconstruct their basis is not always available. A model with `poly` involving two or more predictors will almost always produce misleading results without any warning; but `poly(..., raw = TRUE)` will work correctly.

Note

For a `ref.grid` or `lsmobj` object created in `lsmeans` version 2.10 or earlier, the information needed by the `weights` argument is not present; so a message is displayed and averaging is done using `fac.reduce`.

Author(s)

Russell V. Lenth

References


See Also

`ref.grid`, `.Last.ref.grid`, `models`, `pairwise.lsmc`, `glht`, `lsm.options`

Examples

```r
require(lsmeans)

### Covariance example (from Montgomery Design (8th ed.), p.656)
# Uses supplied dataset 'fiber'
fiber.lm <- lm(strength ~ diameter + machine, data = fiber)

# adjusted means and comparisons, treating machine C as control
(fiber.lsm <- lsmeans(fiber.lm, "machine"))
contrast(fiber.lsm, "trt.vs.ctrlk")
# Or get both at once using
```
### lsmeans

```r
# lsmeans (fiber.lm, "machine", contr = "trt.vs.ctrlk")

### Factorial experiment
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
(warp.lsm <- lsmeans(warp.lm, ~ wool | tension,
    options = list(estName = "pred.breaks")))
pairs(warp.lsm) # remembers 'by' structure
contrast(warp.lsm, method = "poly", by = "wool")

### Unbalanced split-plot example ###
#-- The imbalance is imposed deliberately to illustrate that
#-- the variance estimates become biased
require(nlme)
Oats.lme <- lme(yield ~ factor(nitro) + Variety,
    random = ~1 | Block/ Variety,
    subset = -c(1,2,3,5,8,13,21,34,55), data = Oats)
(O oats.anal <- lsmeans(Oats.lme, list(poly ~ nitro, pairwise ~ Variety)))

### Issues with lists of specs

# Using 'pmmeans' wrapper ...

pmmeans(warp.lm, ~ wool,
    options = list(infer = c(TRUE, TRUE), null = 22, side = ">"))

### Weights
# See what's being averaged over in the above
lsmeans(Oats.lme, ~ nitro, cov.reduce = FALSE, weights = "show.levels")

# Give three times the weight to Marvellous
lsmeans(Oats.lme, ~ nitro, cov.reduce = FALSE, weights = c(1,3,1))

# Overall mean
lsmeans(Oats.lme, ~ 1, weights = "equal")
lsmeans(Oats.lme, "1", weights = "cells")

### Model with a quadratic trend for 'nitro'
# Also illustrates use of 'params' argument to list non-predictors

# Predictions at each unique 'nitro' value in the dataset
lsmeans(Oatsq.lm, ~ nitro, cov.reduce = FALSE, params = "deg")

### Trends
fiber.lm <- lm(strength ~ diameter*machine, data=fiber)
# Obtain slopes for each machine ...
(fiber.lst <- lstrends(fiber.lm, "machine", var="diameter") )
```
# ... and pairwise comparisons thereof
pairs(fiber.lst)

# Suppose we want trends relative to sqrt(diameter)...
listrends(fiber.lm, - machine | diameter, var = "sqrt(diameter)",
at = list(diameter = c(20, 30)))

# Given summary statistics for 4 cities computed elsewhere,
# obtain multiple comparisons of their means using the
# Satterthwaite method
ybar <- c(47.6, 53.2, 88.9, 69.8)
s <- c(12.1, 19.5, 22.8, 13.2)
n <- c(44, 11, 37, 24)
se2 = s^2 / n
Satt.df <- function(x, dfargs)
  sum(x * dfargs$v)^2 / sum((x * dfargs$v)^2 / (dfargs$n - 1))
city.pmm <- pmmobj(bhat = ybar, V = diag(se2),
  levels = list(city = LETTERS[1:4]), linfct = diag(c(1,1,1,1)),
  df = Satt.df, dfargs = list(v = se2, n = n), estName = "mean")
city.pmm
contrast(city.pmm, "revpairwise")

# See also many other examples in documentation for
# 'contrast', 'cld', 'glht', 'lsmip', 'ref.grid', 'MOats',
# 'nutrition', etc., and in the vignettes

---

**lsmip**

*Least-squares (predicted marginal) means interaction plot*

**Description**

This function creates an interaction plot of the least-squares means based on a fitted model and a simple formula specification.

**Usage**

```r
## Default S3 method:
lsmip(object, formula, type,
  pch = c(1,2,6,7,9,10,15:20),
  lty = 1, col = NULL, plotit = TRUE, ...)
lsmip(...)
```

**Arguments**

- **object** An object of class `lsmobj`, or a fitted model of a class supported by `lsmeans`. 
Formula of the form `trace.factors ~ x.factors | by.factors`. The least-squares means are plotted against `x.factors` for each level of `trace.factors`. `by.factors` is optional, but if present, it determines separate panels. Each element of this formula may be a single factor in the model, or a combination of factors using the `*` operator.

As in `predict`, this determines whether we want to inverse-transform the predictions (`type="response"`) or not (any other choice). The default is "link", unless the "predict.type" option is in force; see `lsm.options`.

The plotting characters to use for each group (i.e., levels of `trace.factors`). They are recycled as needed.

The line types to use for each group. Recycled as needed.

The colors to use for each group, recycled as needed. If not specified, the default trellis colors are used.

If `TRUE`, the plot is displayed. Otherwise, one may use the "lattice" attribute of the returned object and print it, perhaps after additional manipulation.

Additional arguments passed to `lsmeans` or to `xyplot`.

If `object` is a fitted model, `lsmeans` is called with an appropriate specification to obtain least-squares means for each combination of the factors present in `formula` (in addition, any arguments in `...` that match `at`, `trend`, `cov.reduce`, or `fac.reduce` are passed to `lsmeans`). Otherwise, if `object` is an `lsmobj` object, its first element is used, and it must contain one `lsmean` value for each combination of the factors present in `formula`.

The wrapper `pmmip` is provided for those who prefer the term ‘predicted marginal means’.

(Invisibly), a `data.frame` with the table of least-squares means that were plotted, with an additional "lattice" attribute containing the trellis object for the plot.

This function uses the `xyplot` function in the lattice package (an error is returned if lattice is not installed). Conceptually, it is equivalent to `interaction.plot` where the summarization function is the least-squares means.

Russell V. Lenth

`interaction.plot`
Examples

```r
require(lsmeans)
require(lattice)

#--- Two-factor example
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)

# Following plot is the same as the usual interaction plot of the data
lsmip(warp.lm, wool ~ tension)

#--- Three-factor example
noise.lm = lm(noise ~ size * type * side, data = auto.noise)

# Separate interaction plots of size by type, for each side
lsmip(noise.lm, type ~ size | side)

# One interaction plot, using combinations of size and side as the x factor
lsmip(noise.lm, type ~ side * size)

# One interaction plot using combinations of type and side as the trace factor
# customize the colors, line types, and symbols to suggest these combinations
lsmip(noise.lm, type * side ~ size, lty=1:2, col=1:2, pch=c(1,1,2,2))

# 3-way interaction is significant, but doesn't make a lot of visual difference...
noise.lm2 = update(noise.lm, ~ . ~ size:type:side)
lsmip(noise.lm2, type * side ~ size, lty=1:2, col=1:2, pch=c(1,1,2,2))
```

---

make.tran  Response transformations

Description

Create the needed information to perform transformations of the response variable, including inverting the transformation and estimating variances of back-transformed predictions via the delta method. `make.tran` is similar to `make.link`, but it covers additional transformations. The result can be used as an environment in which the model is fitted, or as the tran argument in `update.ref.grid` (when the given transformation was already applied in an existing model).

Usage

```r
make.tran(type = c("genlog", "power", "boxcox", "sympower", "asin.sqrt"), param = 1)
```

# See Details for additional auto-detected transformations

Arguments

- **type**
  - The name of the transformation. See Details.

- **param**
  - Numeric parameter for the transformation. Optionally, it may be a vector of two numeric values; the second element specifies an alternative base or origin for certain transformations. See Details.
Details

The functions `lsmeans`, `ref.grid`, and related ones automatically detect response transformations that are recognized by examining the model formula. These are `log`, `log2`, `log10`, `sqrt`, `logit`, `probit`, `cauchit`, `cloglog`; as well as (for a response variable \( y \)) `asin(sqrt(y))`, `asinh(sqrt(y))`, and `sqrt(y) + sqrt(y+1)`. In addition, any constant multiple of these (e.g., `2*sqrt(y)`) is auto-detected and appropriately scaled (see also the `tran.mut` argument in `update.ref.grid`).

A few additional character strings may be supplied as the `tran` argument in `update.ref.grid`: "identity", "1/mu^2", "inverse", "reciprocal", "asin sqrt", and "asinh sqrt".

More general transformations may be provided as a list of functions and supplied as the `tran` argument as documented in `update.ref.grid`. The `make.tran` function returns a suitable list of functions for several popular transformations. Besides being usable with `update`, the user may use this list as an enclosing environment in fitting the model itself, in which case the transformation is auto-detected when the special name `linkfun` (the transformation itself) is used as the response transformation in the call. See the examples below.

Most of the transformations available in "make.tran" require a parameter, specified in `param`; we use \( p \) to denote this parameter, and \( y \) to denote the response variable, in subsequent expressions. The type argument specifies the following transformations:

- "genlog" Generalized logarithmic transformation: \( \log(y + p) \), where \( y > -p \)
- "power" Power transformation: \( y^p \), where \( y > 0 \). When \( p = 0 \), "log" is used instead
- "boxcox" The Box-Cox transformation (unscaled by the geometric mean): \( (y^p - 1)/p \), where \( y > 0 \). When \( p = 0 \), \( \log(y) \) is used.
- "sympower" A symmetrized power transformation on the whole real line: \( \text{abs}(y)^p \ast \text{sign}(y) \). There are no restrictions on \( y \), but we require \( p > 0 \) in order for the transformation to be monotone and continuous.
- "sqrt" Arcsin-square-root transformation: \( \sin(-1)(y/p)^{1/2} \). Typically, the parameter \( p \) equal to 1 for a fraction, or 100 for a percentage.

The user may include a second element in `param` to specify an alternative origin (other than zero) for the "power", "boxcox", or "sympower" transformations. For example, `type = "power", param = c(1.5, 4)` specifies the transformation \( (y - 4)^{1.5} \). In the "genpower" transformation, a second `param` element may be used to specify a base other than the default natural logarithm. For example, `type = "genlog", param = c(.5, 10)` specifies the \( \log_{10}(y + .5) \) transformation.

For purposes of back-transformation, the ‘\( \sqrt{y} + \sqrt{y+1} \)’ transformation is treated exactly the same way as ‘\( 2*\sqrt{y} \)’, because both are regarded as estimates of \( 2\sqrt{\mu} \).

Value

A list having at least the same elements as that returned by `make.link`. The `linkfun` component is the transformation itself.

Note

We modify certain `make.link` results in transformations where there is a restriction on valid prediction values, so that reasonable inverse predictions are obtained for no matter what. For example, if a `sqrt` transformation was used but a predicted value is negative, the inverse transformation is zero rather than the square of the prediction. A side effect of this is that it is possible for one or both confidence limits, or even a standard error, to be zero.
MOats

Author(s)
Russell V. Lenth

See Also
make.link, update

Examples

require("lsmeans")

# Fit a model using an oddball transformation:
bctrans <- make.tran("boxcox", 0.368)
warp.bc <- with(bctrans,
  lm(linkfun(breaks) ~ wool * tension, data = warpbreaks))
# Obtain back-transformed LS means:
lsmeans(warp.bc, ~ tension | wool, type = "response")

## Not run:
## An existing model 'mod' was fitted with a log(y + 1) transformation...
mod.rg <- update(ref.grid(mod), tran = make.tran("genlog", 1))
lsmeans(mod.rg, "treatment")

## End(Not run)

MOats

Oats data in multivariate form

Description
This is the Oats dataset provided in the nlme package, but it is rearranged as one multivariate observation per plot.

Usage
data(MOats)

Format
A data frame with 18 observations on the following 3 variables.

Variety a factor with levels Golden Rain, Marvellous, Victory
Block an ordered factor with levels VI < V < III < IV < II < I
yield a matrix with 4 columns, giving the yields with nitrogen concentrations of 0, .2, .4, and .6.
Details

These data arise from a split-plot experiment reported by Yates (1935) and used as an example in Pinheiro and Bates (2000) and other texts. Six blocks were divided into three whole plots, randomly assigned to the three varieties of oats. The whole plots were each divided into 4 split plots and randomized to the four concentrations of nitrogen.

Source

The dataset Oats in the nlme package.

References


Examples

require(lsmeans)
MOats.lm <- lm (yield ~ Block + Variety, data = MOats)
MOats.rg <- ref.grid (MOats.lm, mult.name = "nitro")
lsmeans(MOats.rg, ~ nitro | Variety)

Models supported in lsmeans

Description

Here we document what model objects may be used with lsmeans, and some special features of some of them. We start with those in the stats package; the other packages follow in alphabetical order.

Certain objects are affected by optional arguments to functions that construct ref.grid or lsmobj objects, including ref.grid, lsmeans, lstrends, and lsmip. When “arguments” are mentioned in the subsequent object-by-object documentation, we are talking about arguments in these constructors.

Additional models can be supported by writing appropriate recover.data and lsm.basis methods. See extending.lsmeans and vignette("extending") for details.

stats package

lm, aov, glm No extended features. Note that the lm support often extends to a number of model objects that inherit from it, such as rlm in the MASS package and rsm in the rsm package.
mlm, maov, manova When there is a multivariate response, the different responses are treated as if they were levels of a factor – named rep.meas by default. The mult.name argument may be used to change this name. The mult.levs argument may specify a named list of one or more sets of levels. If this has more than one element, then the multivariate levels are expressed as combinations of the named factor levels via the function expand.grid.
Support for these objects is limited. To avoid strong biases in the predictions, the contrasts attribute of all factors should be of a type that sums to zero – for example, "contr.sum", "contr.poly", or "contr.helmert" but not "contr.treatment". Only intra-block estimates of covariances are used. That is, if a factor appears in more than one error stratum, only the covariance structure from its lowest stratum is used in estimating standard errors. Degrees of freedom are obtained using the Satterthwaite method. In general, aovlist support is best with balanced designs, and due caution in the use of contrasts. If a vcov. argument is supplied, it must yield a single covariance matrix for the unique fixed effects, and the degrees of freedom are set to NA.

Support for mixed objects has been removed. Version 0.14 and later of afex provides new object classes with their own lsmeans support.

The additional mode argument has possible values of "response", "link", "precision", "phi.link", "variance", and "quantile", which have the same meaning as the type argument in predict.betareg – with the addition that "phi.link" is like "link", but for the precision portion of the model. When mode = "quantile" is specified, the additional argument quantile (a numeric scalar or vector) specifies which quantile(s) to compute; the default is 0.5 (the median). Also in "quantile" mode, an additional variable quantile is added to the reference grid, and its levels are the values supplied.

The user must supply (via the data argument) the dataset used in fitting the model. As with other MCMC-based objects, the summaries and such are frequentist, but the as.mcmc method provides a posterior sample of the desired quantities.

No additional options. Support for these models is experimental; may throw errors or incorrect results.

Currently, gam objects are not supported. Past versions of lsmeans appeared to support gam models owing to inheritance from lm, but the results were incorrect because spline features were ignored. We now explicitly trap gam objects to avoid these misleading analyses.

These models all have more than one covariance estimate available, and it may be selected by supplying a string as the vcov.method argument. It is partially matched with the available choices; thus, for example, ‘vcov = "n"’ translates to ‘vcov.method = "naive"’

Available covariance estimates are specified in vcov.method as "robust" (the default) and "naive".
**geeglm, geese** Available covariance estimates are specified in `vcov.method` as "vbeta" (the default), "vbeta.naiv", "vbeta.jls", or "vbeta.fij". The aliases "robust" (for "vbeta") and "naive" (for "vbeta.naiv") are also accepted.

If a matrix or function is supplied as `vcov.method`, it is interpreted as a `vcov` specification as described for ... in `ref.grid`.

**glmmADMB package**

**glmmadmb** No extended features.

**lme4 package**

**lmerMod** There is an optional `mode` argument that defaults to `get.lsm.option("lmer_df")` (which in turn defaults to "satterthwaite"). The possible values are "satterthwaite", "kenward-roger", and "asymptotic" (these are partially matched and case-insensitive). With "satterthwaite", d.f. are obtained using code from the `lmerTest` package, if installed. With "kenward-roger", d.f. are obtained using code from the `pbkrtest` package, if installed. With "asymptotic", or if the needed package is not installed, d.f. are set to NA.

A by-product of the Kenward-Roger method is that the covariance matrix is adjusted using `vcovAdj`. This can require considerable computation; so to avoid that overhead, the user should opt for the Satterthwaite or asymptotic method; or, for backward compatibility, may disable the use of `pbkrtest` via `lsm.options(disable.pbkrtest=TRUE)` (this does not disable the `pbkrtest` package entirely, just its use in `lsmeans`). The computation time required depends roughly on the number of observations, N, in the design matrix (because a major part of the computation involves inverting an N x N matrix). Thus, `pbkrtest` is automatically disabled if N exceeds the value of `get.lsm.option("pbkrtest.limit")`. If desired, the user may use `lsm.options` to adjust this limit from the default of 3000.

The `df` argument may be used to specify some other degrees of freedom. Note that if `df` and `method = "satterthwaite"` are both specified, the covariance matrix is adjusted but the K-R degrees of freedom are not used.

**glmerMod** No degrees of freedom are available for these objects, so tests and confidence intervals are asymptotic.

**lme4.0 package**

**mer** Only asymptotic results are available (no d.f.).

**MASS package**

**glmmPQL** Supported by virtue of inheritance from `lme` in the `nlme` package.
**glm.nb** Supported by virtue of inheritance from `glm`.

**polr** There are two optional arguments: `mode` and `rescale` (which defaults to \( \{c(0, 1)\} \)). For details, see the documentation below regarding the support for the `ordinal` package, which produces comparable objects (but since `polr` does not support scale models, `mode="scale"` is not supported). Tests and confidence intervals are asymptotic.

**rlm** Supported by virtue of inheritance from `lm`.
MCMCglmm package

MCMCglmm Currently, I have found no way to reconstruct the data based on information in the object; thus, you must provide the dataset via the data argument. In addition, the contrasts specifications are not recoverable from the object, so the system default must match what was actually used in fitting the model. The usual summary, test, etc. methods provide frequentist analyses of the results based on the posterior means and covariances. However, an as.mcmc method is provided that creates an mcmc object that can be summarized or plotted using the coda package. It provides a posterior sample of lsmeans for the given reference grid, based on the posterior sample of the fixed effects from the MCMCglmm object.

MCMCpack package (and perhaps others)

mcmc Certain linear-model or mixed-model objects are of class mcmc, and contain a sample from the posterior distribution of fixed-effect coefficients. In some cases (e.g., results of MCMCregress and MCMCpoisson), the object may include a "call" attribute that lsmeans can use to reconstruct the data and obtain a basis for the least-squares means. If not, a formula and data argument are provided that may help produce the right results. In addition, the contrasts specifications are not recoverable from the object, so the system default must match what was actually used in fitting the model. As for other MCMC-based objects, the summaries and such are frequentist, but the as.mcmc method provides a posterior sample of the desired quantities.

nlme package

gls No additional features. Degrees of freedom are computed using N - p in object$dim. This is consistent with nlme::summary.gls but seems questionable.

lme Degrees of freedom are obtained using a containment method, i.e., the minimum of those elements of object$fixDF$X receiving nonzero weight (but with a correction to the lme object's intercept df). (This is similar to SAS's containment method, but I believe SAS does it incorrectly when the estimands are not contrasts.) The optional argument sigmaAdjust (defaults to TRUE) will adjust standard errors like in summary.lme when the model is fitted using the "ML" method. Note: sigmaAdjust is comparable to adjustSigma in summary.lme but it is renamed to avoid conflicting with adjust.

nlme Support is provided for inferences on parameters named in the fixed part of the model. The user must specify param in the call and give the name of a parameter that appears in the right-hand side of a fixed formula. Degrees of freedom are obtained using the containment-like method described above for lme.

nnet package

multinom The reference grid includes a pseudo-factor with the same name and levels as the multinomial response. There is an optional mode argument which should match "prob" or "latent". With mode = "prob", the reference-grid predictions consist of the estimated multinomial probabilities. The "latent" mode returns the linear predictor, recentered so that it averages to zero over the levels of the response variable (similar to sum-to-zero contrasts). Thus each latent variable can be regarded as the log probability at that level minus the average log probability over all levels.

Please note that, because the probabilities sum to 1 (and the latent values sum to 0) over the multivariate-response levels, all sensible results from lsmeans must involve that response as...
one of the factors. For example, if \( \text{resp} \) is a response with \( k \) levels, \( \text{lsmeans}(\text{model}, \sim \text{resp} | \text{trt}) \) will yield the estimated multinomial distribution for each \( \text{trt} \); but \( \text{lsmeans}(\text{model}, \sim \text{trt}) \) will just yield the average probability of \( 1/k \) for each \( \text{trt} \).

**ordinal package**

clm, clmm  The reference grid will include all variables that appear in the main model as well as those in the scale or nominal models. There are two optional arguments: `mode` (a character string) and `rescale` (which defaults to \( c(0,1) \)). `mode` should match one of "latent" (the default), "linear.predictor", "cum.prob", "exc.prob", "prob", "mean.class", or "scale".

With `mode = "latent"`, the reference-grid predictions are made on the scale of the latent variable implied by the model. The scale and location of this latent variable are arbitrary, and may be altered via `rescale`. The predictions are multiplied by `rescale[2]`, then `rescale[1]` is added. Keep in mind that the scaling is related to the link function used in the model; for example, changing from a probit link to a logistic link will inflate the latent values by around \( \frac{\pi}{\sqrt{3}} \), all other things being equal. `rescale` has no effect for other values of `mode`.

With `mode = "linear.predictor"` mode = "cum.prob", and `mode = "exc.prob"`, the boundaries between categories (i.e., thresholds) in the ordinal response are included in the reference grid as a pseudo-factor named `cut`. The reference-grid predictions are then of the cumulative probabilities at each threshold (for `mode = "cum.prob"`), exceedance probabilities (one minus cumulative probabilities, for `mode = "exc.prob"`), or the link function thereof (for `mode = "linear.predictor"`).

With `mode = "prob"`, a pseudo-factor with the same name as the model’s response variable is created, and the predictions are of the probabilities of each class of the ordinal response. With "mean.class", the returned results are means of the ordinal response, interpreted as a numeric value from 1 to the number of classes, using the "prob" results as the estimated probability distribution for each case.

With `mode = "scale"`, and the fitted object incorporates a scale model, least-squares means are obtained for the factors in the scale model instead of the response model. The grid is constructed using only the factors in the scale model.

Any grid point that is non-estimable by either the location or the scale model (if present) is set to \( \text{NA} \), and any LS-means involving such a grid point will also be non-estimable. A consequence of this is that if there is a rank-deficient scale model, and then all latent responses become non-estimable because the predictions are made using the average log-scale estimate. Tests and confidence intervals are asymptotic.

**pscl package**

hurdle, zeroinfl  Two optional arguments – `mode` and `lin.pred` – are provided. The `mode` argument has possible values "response" (the default), "count", "zero", or "prob". `lin.pred` is logical and defaults to `FALSE`.

With `lin.pred = FALSE`, the results are comparable to those returned by \( \text{predict(..., type = "response")}, \text{predict(..., type = "count")}, \text{predict(..., type = "zero")}, \text{or predict(..., type = "prob")}[1] \).

See the documentation for `predict.hurdle` and `predict.zeroinfl`.

The option `lin.pred = TRUE` only applies to `mode = "count"` and `mode = "zero"`. The results returned are on the linear-predictor scale, with the same transformation as the link function in that part of the model. The predictions for a reference grid with `mode = "count"`,
lin.pred = TRUE, and type = "response" will be the same as those obtained with lin.pred = FALSE and mode = "count"; however, any LS means derived from these grids will be different, because the averaging is done on the log-count scale and the actual count scale, respectively – thereby producing geometric means versus arithmetic means of the predictions.

If the vcov. argument is used (see details in ref.grid), it must yield a matrix of the same size as would be obtained using vcov.hurdle or vcov.zeroinfl with its model argument set to ("full", "count", "zero") in respective correspondence with mode of ("mean", "count", "zero"). If vcov. is a function, it must support the model argument.

**rms package**

**Potential masking issue** Both rms and lsmeans offer contrast methods, and whichever package is loaded later masks the other. Thus, you may need to call lsmeans::contrast or rms::contrast explicitly to access the one you want.

**Objects inheriting from rms** Standard support is provided. However, with models having more than one intercept (e.g. from orm), a mode argument is provided that works similarly to that for the ordinal package. The available modes are "middle" (the default), "latent", "linear.predictor", "cum.prob", "exc.prob", "prob", and "mean.class". All are as described for the ordinal package, except as noted below.

With mode = "middle" (this is the default), the middle intercept is used, comparable to the default for rms's Predict function. This is quite similar in concept to mode = "latent", where all intercepts are averaged together.

Results for mode = "linear.predictor" are reversed from those in the ordinal package, because orm models predict the link function of the upper-tail (exceedance) probabilities.

With mode = "prob", a pseudo-factor is created having the same name as the model response variable, but its levels are always integers ‘1, 2, …’ regardless of the levels of the original response.

**rstanarm package**

**stanreg** Support for models fitted using stan_xxx is similar to that for models fitted by xxx, where supported, except that posterior samples are also available. For example, stan_glms models are treated like stats::glm, and stan_polr results are similar to those for MASS::polr (including its available mode and rescale options). Models fitted using stan_biglm, stan_betareg, and stan_gamm4 are currently not supported. The user may use as.mcmc, as.mcmc.list, or as.stanfit on the ref.grid or lsmobj produced to obtain posterior samples of LS means, contrasts, etc.

**survival package**

**survreg, coxph** No extended features.

**Author(s)**

Russell V. Lenth
See Also

ref.grid, lsm.basis

nutrition Nutrition data

Description

This observational dataset involves three factors, but where several factor combinations are missing. It is used as a case study in Milliken and Johnson, Chapter 17, p.202. (You may also find it in the second edition, p.278.)

Usage

nutrition

Format

A data frame with 107 observations on the following 4 variables.

age  a factor with levels 1, 2, 3, 4. Mother’s age group.
group  a factor with levels FoodStamps, NoAid. Whether or not the family receives food stamp assistance.
race  a factor with levels Black, Hispanic, White. Mother’s race.
gain  a numeric vector (the response variable). Gain score (posttest minus pretest) on knowledge of nutrition.

Details

A survey was conducted by home economists “to study how much lower-socioeconomic-level mothers knew about nutrition and to judge the effect of a training program designed to increase their knowledge of nutrition.” This is a messy dataset with several empty cells.

Source


Examples

```r
require(lsmeans)
nutr.aov <- aov(gain ~ (group + age + race)^2, data = nutrition)

# Summarize predictions for age group 3
nutr.lsm <- lsmmeans(nutr.aov, ~ race * group,
                     at = list(age="3"))
```
Description

This example dataset on sales of oranges has two factors, two covariates, and two responses. There is one observation per factor combination.

Usage

data(orange)

Format

A data frame with 36 observations on the following 6 variables.

store  a factor with levels 1 2 3 4 5 6. The store that was observed.
day   a factor with levels 1 2 3 4 5 6. The day the observation was taken (same for each store).
price1 a numeric vector. Price of variety 1.
price2 a numeric vector. Price of variety 2.
sales1 a numeric vector. Sales (per customer) of variety 1.
sales2 a numeric vector. Sales (per customer) of variety 2.

Source


References


Examples

require(lsmeans)

# Example on p.244 of Littell et al.
orange.lm <- lm(sales ~ price1*day, data = orange)
lsmeans(orange.lm, "day")

# Example on p.246
lsmeans(orange.lm, "day", at = list(price1 = 0))
pairwise.lsmc

Contrast families

Description

These functions return standard sets of contrast coefficients. The name of any of these functions (with the .lsmc omitted) may be used as the method argument in `contrast`, or as the contr argument or left-hand side of a spec formula in `lsmeans`.

Usage

```r
pairwise.lsmc(levs, ...)  # All pairwise
revpairwise.lsmc(levs, ...)  # Reverse-pairwise
tukey.lsmc(levs, reverse = FALSE)

poly.lsmc(levs, max.degree = min(6, k - 1))

trt.vs.ctrl.lsmc(levs, ref = 1)  # Treatments vs control
trt.vs.ctrl1.lsmc(levs, ...)  # Treatments vs control 1
trt.vs.ctrlk.lsmc(levs, ...)  # Treatments vs control k
dunnett.lsmc(levs, ref = 1)

consec.lsmc(levs, reverse = FALSE, ...)  # Consecutive
mean_chg.lsmc(levs, reverse = FALSE, ...)

eff.lsmc(levs, ...)  # Treatment effects
del.eff.lsmc(levs, ...)
```

Arguments

- `levs`: Vector of factor levels
- `...`: Additional arguments, ignored but needed to make these functions interchangeable
- `max.degree`: The maximum degree of the polynomial contrasts in `poly.lsmc`
- `reverse`: Logical value to determine the direction of comparisons, e.g., pairwise (if TRUE) or reverse-pairwise (if FALSE) comparisons.
- `ref`: Reference level (or control group) in `trt.vs.ctrl.lsmc`

Details

Each contrast family has a default multiple-testing adjustment as noted below. These adjustments are often only approximate; for a more exacting adjustment, use the interfaces provided to `glht` in the `multcomp` package.

`pairwise.lsmc`, `revpairwise.lsmc`, and `tukey.lsmc` generate contrasts for all pairwise comparisons among least-squares means at the levels in `levs`. The distinction is in which direction they are subtracted. For factor levels A, B, C, D, `pairwise.lsmc` generates the comparisons A-B, A-C,
A-D, B-C, B-D, and C-D, whereas `revpairwise.lsmc` generates B-A, C-A, C-B, D-A, D-B, and D-C. `tukey.lsmc` invokes `pairwise.lsmc` or `revpairwise.lsmc` depending on `reverse`. The default multiplicity adjustment method is "tukey", which is approximate when the standard errors differ.

`poly.lsmc` generates orthogonal polynomial contrasts, assuming equally-spaced factor levels. These are derived from the `poly` function, but an ad hoc algorithm is used to scale them to integer coefficients that are (usually) the same as in published tables of orthogonal polynomial contrasts. The default multiplicity adjustment method is "none".

`tutryNlsmc` invokes `pairwiseNlsmc` or `revpairwiseNlsmc` depending on `reverse`. The default multiplicity adjustment method is `BtukeyB`, which is approximate when the standard errors differ.

`polyNlsmc` generates orthogonal polynomial contrasts, assuming equally-spaced factor levels. These are derived from the `poly` function, but an ad hoc algorithm is used to scale them to integer coefficients that are (usually) the same as in published tables of orthogonal polynomial contrasts. The default multiplicity adjustment method is "none".

`trtNvsNctrlNlsmc` and its relatives generate contrasts for comparing one level (or the average over specified levels) with each of the other levels. The argument `ref` should be the `index(es)` of the reference level(s). `trtNvsNctrl1Nlsmc` is the same as `trtNvsNctrl` with a reference value of 1, and `trtNvsNctrlkNlsmc` is the same as `trtNvsNctrl` with a reference value of length(`levs`). `dunnettNlsmc` is the same as `trtNvsNctrl`. The default multiplicity adjustment method is "dunnetttx", a close approximation to the Dunnett adjustment.

`consecNlsmc` and `mean_chgNlsmc` are useful for contrasting treatments that occur in sequence. For a factor with levels A, B, C, D, E, `consecNlsmc` generates the comparisons B-A, C-B, and D-C, while `mean_chgNlsmc` generates the contrasts (B+C+D)/3 - A, (C+D)/2 - (A+B)/2, and D - (A+B+C)/3. With `reverse = TRUE`, these differences go in the opposite direction.

`effNlsmc` and `del1.effNlsmc` generate contrasts that compare each level with the average over all levels (in `effNlsmc`) or over all other levels (in `del1.effNlsmc`). These differ only in how they are scaled. For a set of `k` lsmeans, `del1.effNlsmc` gives weight 1 to one lsmean and weight `-1/(k-1)` to the others, while `effNlsmc` gives weights `(k-1)/k` and `-1/k` respectively, as in subtracting the overall lsmean from each lsmean. The default multiplicity adjustment method is "fdr". This is a Bonferroni-based method and is slightly conservative; see `p.adjust`

**Value**

A data frame, each column containing contrast coefficients for `levs`. The "desc" attribute is used to label the results in `lsmeans`, and the "adjust" attribute gives the default adjustment method for multiplicity.

**Note**

You may create your own contrast functions, using these as guides. A function named `mycontr.lsmc` may be invoked in `lsmeans` via, e.g.,

```r
lsmeans(~ var{factor}), mycontr ~ var{factor})
```

The "desc", "adjust", and "offset" attributes are optional; if present, these are passed to `contrast`. If absent, the root name of the function is used as "desc", and no adjustment is requested for `p` values. See the examples.

**Author(s)**

Russell V. Lenth

**See Also**

`lsmeans`, `glht`
Examples

### View orthogonal polynomials for 4 levels
poly.lsmc(1:4)

### Setting up a custom contrast function
helmert.lsmc <- function(levs, ...)
  
  M <- as.data.frame(contr.helmert(levs))
  names(M) <- paste(levs[-1], "vs earlier")
  attr(M, "desc") <- "Helmert contrasts"
  
  M

warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)
lsmmeans(warp.lm, helmert ~ tension | wool)

---

recover.data  Support functions for creating a reference grid

Description

This documents the methods used to create a `ref.grid` object from a fitted model.

Usage

```r
recover.data(object, ...)
## S3 method for class 'call'
recover.data(object, trms, na.action,
            data = NULL, params = NULL, ...)
```

lsm.basis(object, trms, xlev, grid, ...)

Arguments

- **object**: An object returned from a model-fitting function.
- **trms**: The `terms` component of object
- **xlev**: Named list of levels of factors in the model frame. This should *not* include levels of factors created in the model itself, e.g., by including a `factor` call in the model formula.
- **grid**: A `data.frame` containing predictor values at which predictions are needed.
- **na.action**: Integer vector of indices of observations to ignore; or `NULL` if none
- **data**: Data frame. Usually, this is `NULL`. However, if non-null, this is used in place of the reconstructed dataset. It must have all of the predictors used in the model, and any factor levels must match those used in fitting the model.
- **params**: Character vector giving the names of any variables in the model formula that are *not* predictors. An example would be a variable `knots` specifying the knots to use in a spline model.
- **...**: Additional arguments passed to other methods.
Details

To create a reference grid, the ref.grid function needs to reconstruct the data used in fitting the model, and then obtain a matrix of linear functions of the regression coefficients for a given grid of predictor values. These tasks are performed by calls to recover.data and lsm.basis respectively.

To extend lsmeans's support to additional model types, one need only write S3 methods for these two functions. The existing methods serve as helpful guidance for writing new ones. Most of the work for recover.data can be done by its method for class "call", providing the terms component and na.action data as additional arguments. Writing an lsm.basis method is more involved, but the existing methods (e.g., lsmeans:::lsm.basis.lm) can serve as models. See the “Value” section below for details on what it needs to return. Also, certain recover.data and lsm.basis methods are exported from lsmeans, so if your object is based on another model-fitting object, it may be that all that is needed is to call one of these exported methods and perhaps make modifications to the results. Contact the developer if you need others of these exported.

If the model has a multivariate response, bhat needs to be “flattened” into a single vector, and X and V must be constructed consistently.

In models where a non-full-rank result is possible (often you can tell by seeing if there is a singular.ok argument in the model-fitting function), summary and predict check the estimability of each prediction, using the nonest.basis function in the estimability package.

The models already supported are detailed in models. Some packages may provide additional lsmeans support for its object classes.

Value

recover.data should return a data.frame containing all the variables in the original data that appear as predictors in the model. Several attributes need to be included as well; see the code for lsmeans:::recover.data.lm.

lsm.basis should return a list with the following elements:

X  The matrix of linear functions over grid, having the same number of rows as grid and the number of columns equal to the length of bhat.

bhat  The vector of regression coefficients for fixed effects. This should include any NAs that result from rank deficiencies.

nbasis  A matrix whose columns form a basis for non-estimable functions of beta, or a 1x1 matrix of NA if there is no rank deficiency.

V  The estimated covariance matrix of bhat.

dffun  A function of (k, dfargs) that returns the degrees of freedom associated with sum(k * bhat).

dfargs  A list containing additional arguments needed for dffun.

Optional hooks

Some models may need something other than standard linear estimates and standard errors. If so, custom functions may be pointed to via the items misc$estHook, misc$vcovHook and misc$postGridHook. If just the name of the hook function is provided as a character string, then it is retrieved using get.

The estHook function should have arguments ‘(object, do.se, tol, ...)’ where object is the ref.grid or lsmobj object, do.se is a logical flag for whether to return the standard error,
and tol is the tolerance for assessing estimability. It should return a matrix with 3 columns: the estimates, standard errors (NA when do.se==FALSE), and degrees of freedom (NA for asymptotic). The number of rows should be the same as 'object@linfct'. The vcovHook function should have arguments ‘(object, tol, ...)’ as described. It should return the covariance matrix for the estimates. Finally, postGridHook, if present, is called at the very end of ref.grid; it takes one argument, the constructed object, and should return a suitably modified ref.grid object.

**Additional functions**

A few additional functions used in the lsmeans codebase are exported as they may be useful to package developers. See details near the end of the vignette "extending".

**Author(s)**

Russell V. Lenth

**See Also**

models, ref.grid, ref.grid-class

**Examples**

```r
## Not run:
require(lsmeans)

# Fit a 2-factor model with two empty cells
warpings.lm <- lm(breaks ~ wool*tension,
       data = warpbreaks, subset = -(16:40))

lsmeans:::recover.data.lm(warpings.lm, data = NULL)
grid = with(warpbreaks,
       expand.grid(wool = levels(wool), tension = levels(tension)))
lsmeans:::lsm.basis.lm(warpings.lm, delete.response(terms(warpings.lm)),
       warpings.lm$xlevels, grid)

## End(Not run)
```

---

**ref.grid**

Create a reference grid from a fitted model

**Description**

Using a fitted model object, determine a reference grid for which least-squares means are defined. The resulting ref.grid object encapsulates all the information needed to calculate LS means and make inferences on them.
Usage

ref.grid(object, at, cov.reduce = mean, mult.name, mult.levs, options = get.lsm.option("ref.grid"), data, df, type, transform = c("none", "response", "mu", "unlink", "log"), nesting, ...)

Arguments

object An object produced by a supported model-fitting function, such as lm. Many models are supported. See models.

at Optional named list of levels for the corresponding variables

cov.reduce A function, logical value, or formula; or a named list of these. Each covariate not specified in at is reduced according to these specifications. If a single function, it is applied to each covariate. If logical and TRUE, mean is used. If logical and FALSE, it is equivalent to specifying ‘function(x) sort(unique(x))’, and these values are considered part of the reference grid; thus, it is a handy alternative to specifying these same values in at.

If a formula (which must be two-sided), then a model is fitted to that formula using lm; then in the reference grid, its response variable is set to the results of predict for that model, with the reference grid as newdata. (This is done after the reference grid is determined.) A formula is appropriate here when you think experimental conditions affect the covariate as well as the response.

If cov.reduce is a named list, then the above criteria are used to determine what to do with covariates named in the list. (However, formula elements do not need to be named, as those names are determined from the formulas’ left-hand sides.) Any unresolved covariates are reduced using "mean".

Any cov.reduce specification for a covariate also named in at is ignored.

mult.name Character, the name to give to the “factor” whose levels delineate the elements of a multivariate response. If this is provided, it overrides the default name, e.g., “rep.meas” for an mlm object or “cut” for a polr object.

mult.levs A named list of levels for the dimensions of a multivariate response. If there is more than one element, the combinations of levels are used, in expand.grid order. The (total) number of levels must match the number of dimensions. If mult.name is specified, this argument is ignored.

options If non-NULL, a named list of arguments to pass to update, just after the object is constructed.

data A data.frame to use to obtain information about the predictors (e.g., factor levels). If missing, then recover.data is used to attempt to reconstruct the data.

df This is a courtesy shortcut, equivalent to specifying options(df = df). See update.

type If provided, this is saved as the “predict.type” setting. See update
transform If other than "none", the reference grid is reconstructed via `regrid` with the given transform argument. See Details.

nesting If the model has nested fixed effects, this may be specified here via a named list specifying the nesting structure. Specifying nesting overrides the nesting structure that may be automatically detected. See Details.

... Optional arguments passed to `lsm.basis`, such as `vcov`. (see Details below) or options for certain models (see `models`).

Details

The reference grid consists of combinations of independent variables over which predictions are made. Least-squares means are defined as these predictions, or marginal averages thereof. The grid is determined by first reconstructing the data used in fitting the model (see `recover.data`), or by using the `data.frame` provided in context. The default reference grid is determined by the observed levels of any factors, the ordered unique values of character-valued predictors, and the results of `cov.reduce` for numeric predictors. These may be overridden using `at`.

Ability to support a particular class of object depends on the existence of `recover.data` and `lsm.basis` methods – see `extending-lsmeans` for details. The call methods("recover.data") will help identify these.

In certain models, (e.g., results of `glmer.nb`), it is not possible to identify the original dataset. In such cases, we can work around this by setting data equal to the dataset used in fitting the model, or a suitable subset. Only the complete cases in data are used, so it may be necessary to exclude some unused variables. Using data can also help save computing, especially when the dataset is large. In any case, data must represent all factor levels used in fitting the model. It cannot be used as an alternative to at. (Note: If there is a pattern of NAs that caused one or more factor levels to be excluded when fitting the model, then data should also exclude those levels.)

By default, the variance-covariance matrix for the fixed effects is obtained from object, usually via its `vcov` method. However, the user may override this via a `vcov` argument, specifying a matrix or a function. If a matrix, it must be square and of the same dimension and parameter order of the fixed effects. If a function, must return a suitable matrix when it is called with `object` as its only argument.

Nested factors: `ref.grid` tries to discern which factors are nested in other factors, but it is not always obvious, and if it misses some, the user must specify this structure via nesting; or later using `update`. Each member of nesting should be a character vector of the name(s) of grouping factors; and the name for that member should be that of the factor that is nested therein; for example, `list(city = c("state", "country"))`. Having a nesting structure affects marginal averaging in `lsmeans` in that it is done separately for each level (or combination thereof) of the grouping factors.

There is a subtle difference between specifying `type = "response"` and `transform = "response"`. While the summary statistics for the grid itself are the same, subsequent use in `lsmeans` will yield different results if there is a response transformation. With `type = "response"`, LS means are computed by averaging together predictions on the linear-predictor scale and then back-transforming to the response scale; while with `transform = "response"`, the predictions are already on the response scale so that the LS means will be the arithmetic means of those response-scale predictions. To add further to the possibilities, geometric means of the response-scale predictions are obtainable via `transform = "log", type = "response"`.

The most recent result of `ref.grid`, whether called directly or indirectly via `lsmeans`, `lstrends`, or some other function that calls one of these, is saved in the user's environment as `.Last.ref.grid`.
This facilitates checking what reference grid was used, or reusing the same reference grid for further calculations. This automatic saving is enabled by default, but may be disabled via `lsm.options(save.ref.grid = FALSE)` and re-enabled by specifying `TRUE`.

Value

An S4 object of class "ref.grid" (see ref.grid-class). These objects encapsulate everything needed to do calculations and inferences for least-squares means, and contain nothing that depends on the model-fitting procedure. As a side effect, the result is also saved as `.Last.ref.grid` (in the global environment, unless this variable is found in another position).

Author(s)

Russell V. Lenth

See Also

See also `summary` and other methods for the returned objects. Reference grids are fundamental to `lsmeans`. Click here for more on the `ref.grid` class. Supported models are detailed in `models`.

Examples

```r
require(lsmeans)

fiber.lm <- lm(strength ~ machine*diameter, data = fiber)
ref.grid(fiber.lm)
summary(ref.grid(fiber.lm))

ref.grid(fiber.lm, at = list(diameter = c(15, 25)))

# Not run:
# We could substitute the sandwich estimator vcovHAC(fiber.lm)
# as follows:
require(sandwich)
summary(ref.grid(fiber.lm, vcov. = vcovHAC))

# End(Not run)

# If we thought that the machines affect the diameters
# (admittedly not plausible in this example), then we should use:
ref.grid(fiber.lm, cov.reduce = diameter~machine)

# Multivariate example
MOats.lm = lm(yield ~ Block + Variety, data = MOats)
ref.grid(MOats.lm, mult.name = "nitro")
# silly illustration of how to use 'mult.levs'
ref.grid(MOats.lm, mult.levs = list(T=LETTERS[1:2], U=letters[1:2]))
```
A reference grid encapsulates everything needed to compute least-squares means, independently of the underlying model object. The "lsmobj" class is a minor extension of "ref.grid" where the linear predictors for the reference grid are transformed in some linear way such as marginal averages or contrasts.

Objects from the Classes

Objects of class "ref.grid" are most commonly created by calling the ref.grid function.

Objects of class "lsmobj" are created by calling lsmeans or a related function such as contrast.

Slots

model.info: Object of class "list" containing the elements call (the call that produced the model), terms (its terms object), and xlev (factor-level information)

roles: Object of class "list" containing at least the elements predictors, responses, and multresp. These are character vectors of names of these variables.

grid: Object of class "data.frame" containing the combinations of the variables that define the reference grid. In addition, there is an auxiliary column named ".wgt." holding the observed frequencies or weights for each factor combination (excluding covariates). If the model has one or more offset() calls, there is another auxiliary column named ".offset.". Auxiliary columns are not considered part of the reference grid. (However, any variables included in offset calls are in the reference grid.)

levels: Object of class "list" with each entry containing the distinct levels of variables in the reference grid. Note that grid is obtained by applying the function expand.grid to this list

matlevs: Object of class "list" Like levels but has the levels of any matrices in the original dataset. Matrix columns must always be reduced to a single value for purposes of the reference grid

linfct: Object of class "matrix" giving the linear functions of the regression coefficients for predicting each element of the reference grid. The rows of this matrix go in one-to-one correspondence with the rows of grid, and the columns with elements of bhat

bhat: Object of class "numeric" with the regression coefficients. If there is a multivariate response, this must be flattened to a single vector, and linfct and V redefined appropriately. Important: bhat must include any NA values produced by collinearity in the predictors. These are taken care of later in the estimability check.

nbasis: Object of class "matrix" with the basis for the non-estimable functions of the regression coefficients. Every LS mean will correspond to a linear combination of rows of linfct, and that result must be orthogonal to all the columns of nbasis in order to be estimable. This will be NULL if everything is estimable

V: Object of class "matrix", the symmetric variance-covariance matrix of bhat
dffun, dfargs: Objects of class "function" and "list" respectively. dffun(k, dfargs) should return the degrees of freedom for the linear function sum(k*bhat), or NA if unavailable.

misc: A list containing additional information used by methods. These include at least the following: estName (the label for the estimates of linear functions), and the default values of infer, level, and adjust to be used in the summary method. Elements in this slot may be modified if desired using the update method.

post.beta: A matrix containing a sample from the posterior distribution of the regression coefficients; or a 1 x 1 matrix of NA if this is not available. When it is non-trivial, the as.mcmc method returns post.beta times t(linfct), which is a sample from the posterior distribution of the LS means.

**Extends**

Class "lsmobj" extends Class "ref.grid", directly. There is hardly a difference between these classes except for how the slots linfct and grid are obtained, and their show methods.

**Methods**

All methods for these objects are S3 methods except for show.

show: Prints the results of str for ref.grid objects, and summary for lsmobj objects.

str: Displays a brief listing of the variables and levels defining the grid.

summary: Displays a summary of estimates, standard errors, degrees of freedom, and optionally, tests and/or confidence intervals.

lsmeans: Computes least-squares means and creates an "lsmobj" object.

confint: Confidence intervals for lsmeans.

test: Hypothesis tests.

cld: Compact-letter display for tests of pairwise comparisons

contrast: Contrasts among lsmeans.

pairs: A special case of contrasts for pairwise comparisons.

update: Change defaults used primarily by summary, such as transformation, p-value adjustment, and confidence level.

**Author(s)**

Russell V. Lenth

**See Also**

ref.grid, lsmeans

**Examples**

showClass("ref.grid")
showClass("lsmobj")
Methods for ref.grid objects

Description

Use these methods to summarize, print, plot, or examine objects of class "ref.grid". They also apply to the class "lsmobj", which is an extension of "ref.grid".

Usage

```r
## S3 method for class 'ref.grid'
summary(object, infer, level, adjust, by, type, df,
         null, delta, side, ...)

## S3 method for class 'ref.grid'
predict(object, type, ...)

## S3 method for class 'ref.grid'
str(object, ...)

## S3 method for class 'ref.grid'
rbind(..., deparse.level = 1, adjust = "mvt")

## S3 method for class 'ref.grid'
x[i, adjust = "mvt", drop.levels = TRUE, ...]

## S3 method for class 'ref.grid'
print(x, ...)

## S3 method for class 'summary.ref.grid'
print(x, ..., digits = NULL, quote = FALSE, right = TRUE)

## S3 method for class 'ref.grid'
xtable(x, caption = NULL, label = NULL, align = NULL,
         digits = 4, display = NULL, auto = FALSE, ...)

## S3 method for class 'summary.ref.grid'
xtable(x, caption = NULL, label = NULL, align = NULL,
         digits = 4, display = NULL, auto = FALSE, ...)

## S3 method for class 'xtable.lsm'
print(x, type = getOption("xtable.type", "latex"),
       include.rownames = FALSE, sanitize.message.function = footnotesize, ...

## S3 method for class 'lsmobj'
plot(x, y, type, intervals = TRUE, comparisons = FALSE,
     alpha = 0.05, adjust = "tukey", int.adjust, ...)

## S3 method for class 'summary.ref.grid'
plot(x, y, horizontal = TRUE,
     ...)```
Arguments

object An object of class "ref.grid".

infer A vector of two logical values. The first determines whether confidence intervals are displayed, and the second determines whether t-tests and P-values are displayed. If only one value is provided, it is used for both.

level Confidence level for confidence intervals, if infer[1] is TRUE.

adjust Character value naming the method used to adjust p-values or confidence limits; or to adjust comparison arrows in plot. See Details.

by Character name(s) of variables to use for grouping. This affects the family of tests considered in adjusted P-values. The printed display of the summary is grouped by the by variables.

type Type of prediction desired (except in print.xtable). This only has an effect if there is a known transformation or link function. "response" specifies that the inverse transformation be applied. "mu" (or equivalently, "unlink") is usually the same as "response", but in the case where the model has both a link function and a response transformation, only the link part is back-transformed. Other valid values are "link", "lp", and "linear"; these are equivalent, and request that results be shown for the linear predictor, with no back-transformation. The default is "link", unless the "predict.type" option is in force; see lsm.options. Note that type is also an argument for the print.xtable method; it is passed to print.xtableList in the xtable package.

df If non-missing a constant number of degrees of freedom to use in constructing confidence intervals and P-values (NA specifies asymptotic results).

null Null hypothesis value(s) against which estimates are tested. May be a single value used for all, or a numeric vector of length equal to the number of tests in each family (i.e., by group in the displayed table).

delta Numeric value. If zero, ordinary tests of significance are performed. If positive, this specifies a threshold for testing equivalence (using the TOST or two-sided-test method), non-inferiority, or non-superiority, depending on side. See Details for how the test statistics are defined.
summary

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side Numeric or character value specifying whether the test is left-tailed (-1, "-", code"<", "left", or "nonsuperiority"); right-tailed (1, "+", ">", "right", or "noninferiority"); or two-sided (0, 2, "+", "two-sided", "both", "equivalence", or "+").

deparse.level This argument is needed by the generic rbind method, but ignored by its ref.grid method.

drop.levels Logical value to specify whether or not the levels slot should be recomputed based on a possibly reduced number of levels of factors in the grid.

x The object to be subsetted, printed, plotted, or converted.

y This argument is ignored.

i Integer index(es) of which linear functions to extract.

horizontal Determines orientation of plotted confidence intervals.

intervals If TRUE, confidence intervals are plotted for each estimate

comparisons If TRUE, “comparison arrows” are added to the plot, in such a way that the degree to which arrows overlap reflects as much as possible the significance of the comparison of the two estimates.

alpha The alpha argument to use in constructing comparison arrows.

int.adjust the multiplicity adjustment method for the plotted confidence intervals; if missing, it defaults to the object’s internal adjust setting (see update). (Note: the adjust argument in plot sets the adjust method for the comparison arrows, not the confidence intervals.)

transform Character value. If "response" or "mu", the inverse transformation is applied to the estimates in the grid (but if there is both a link function and a response transformation, "mu" back-transforms only the link part); if "log", the results are formulated as if the response had been log-transformed; if "none", predictions thereof are on the same scale as in object, and any internal transformation information is preserved. For compatibility with past versions, transform may also be logical; TRUE is taken as "response", and FALSE as "none".

inv.log.lbl Character value. This applies only when transform = "log", and is used to label the predictions if subsequently summarized with type = "response".

predict.type Character value. If provided, the returned object is first updated with the given type, e.g., "response".

names Logical scalar or vector specifying whether variable names are appended to levels in the column labels for the as.mcmc or as.mcmc.list result – e.g., column names of treat A and treat B versus just A and B. When there is more than one variable involved, the elements of names are used cyclically.

sep.chains Logical value. If TRUE, and there is more than one MCMC chain available, an mcmc.list object is returned by as.mcmc, with separate lsmeans posteriors in each chain.

..., digits, quote, right, caption, label, align, display, auto, include, rownames, sanitize.message For summaries, these are additional arguments passed to other methods including print.data.frame, xtablelist, print.xtablelist, update, or dotplot as appropriate. If not specified, appropriate defaults are used. For example, the default layout is one column of horizontal panels or one row of vertical panels.
Details

Defaults for summarization, etc.: The misc slot in object contains default values for by, infer, level, adjust, type, null, side, and delta. These defaults vary depending on the code that created the object. The update method may be used to change these defaults. In addition, any options set using `lsm.options(summary=...)` will trump those stored in the object's misc slot.

Transformations and links: With type="response", the transformation assumed can be found in 'object@misc$tran', and its label, for the summary is in 'object@misc$inv.lbl'. Any t or z tests are still performed on the scale of the linear predictor, not the inverse-transformed one. Similarly, confidence intervals are computed on the linear-predictor scale, then inverse-transformed.

Confidence-limit and P-value adjustments: The adjust argument specifies a multiplicity adjustment for tests or confidence intervals. This adjustment always is applied separately to each table or sub-table that you see in the printed output (see the details on rbind below for how to combine tables). The valid values of adjust are as follows:

"tukey" Uses the Studentized range distribution with the number of means in the family. (Available for two-sided cases only.)

"scheffe" Computes p values from the F distribution, according to the Scheffe critical value of $\sqrt{kF(k,d)}$, where d is the error degrees of freedom and k is (family size minus 1) for contrasts, and (number of estimates) otherwise. (Available for two-sided cases only.)

"sidak" Makes adjustments as if the estimates were independent (a conservative adjustment in many cases).

"bonferroni" Multiplies p values, or divides significance levels by the number of estimates. This is a conservative adjustment.

"dunnetttx" Uses an approximation to the Dunnett distribution for a family of estimates having pairwise correlations of 0.5 (as is true when comparing treatments with a control with equal sample sizes). The accuracy of the approximation improves with the number of simultaneous estimates, and is much faster than "mvt". (Available for two-sided cases only.)

"mvt" Uses the multivariate t distribution to assess the probability or critical value for the maximum of k estimates. This method produces the same p values and intervals as the default summary or confint methods to the results of as.glht. In the context of pairwise comparisons or comparisons with a control, this produces "exact" Tukey or Dunnett adjustments, respectively. However, the algorithm (from the mvtnorm package) uses a Monte Carlo method, so results are not exactly repeatable unless the random-number seed is used (see set.seed). As the family size increases, the required computation time will become noticeable or even intolerable, making the "tukey", "dunnetttx", or others more attractive.

"none" Makes no adjustments to the p values.

For P-value adjustments only, the Bonferroni-inequality-based adjustment methods in p.adjust are also available (currently, these include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", and "none"). If a p.adjust.methods method other than "bonferroni" or "none" is specified for confidence limits, the straight Bonferroni adjustment is used instead. Also, if an adjustment method is not appropriate (e.g., using "tukey" with one-sided tests, or with results that are not pairwise comparisons), a more appropriate method (usually "sidak") is substituted.

In some cases, confidence and p-value adjustments are only approximate – especially when the degrees of freedom or standard errors vary greatly within the family of tests. The "mvt" method is
always the correct one-step adjustment, but it can be very slow. One may use as.glht with methods in the multcomp package to obtain non-conservative multi-step adjustments to tests.

**Information:** The `str` method outputs a very brief summary of the object, whereas `levels` produces a `data.frame` containing the combinations of levels of predictor values that define the reference grid.

**xtable-related methods:** The `xtable` methods actually use `xtableList`, because of the ability to display messages such as those for P-value adjustments. These methods return an object of class "xtable.lsm" – an extension of "xtableList". Unlike other `xtable` methods, the number of digits defaults to 4; and degrees of freedom and t ratios are always formatted independently of digits. The print method uses `print.xtableList`, and any ... arguments are passed there.

**rbind** and **methods:** `rbind` can be used to combine two or more reference grids into one. The "[" method for `refGrids` may be used to obtain a subset. The primary reason for doing this would be to redefine the family of tests to which a P-value adjustment method applies. In `rbind`, the variables defined in the objects’ grids are merged into one grid, and the returned object has no “by” variables and the multiplicity adjustment method set to "mvt" (as this is likely the only appropriate one). `rbind` throws an error if there are any mismatches among the dimensions, fixed-effect coefficients, or covariance matrices.

**Non-estimable cases:** When the model is rank-deficient, each row x of object’s `linfct` slot is each checked for estimability. If `sum(x*bhat)` is found to be non-estimable, then an NA is displayed for the estimate (as well as any associated statistics). This check is performed using the orthonormal basis N in the `nbasis` slot for the null space of the rows of the model matrix. Estimability fails when `||N.x||^2/||x||^2` exceeds `tol`, which by default is 1e-8. You may change it via `lsm.options` by setting `estable.tol` to the desired value.

**More on tests:** When `delta = 0`, test statistics are of the usual form `(estimate - null)/SE`, or notationally, `t = (Q - θ_0)/SE` where Q is our estimate of θ; then left, right, or two-sided p values are produced.

When `delta` is positive, the test statistic depends on side as follows.

- Left-sided (nonsuperiority): `H_0 : θ ≥ θ_0 + δ` versus `H_1 : θ < θ_0 + δ`; `t = (Q - θ_0 - δ)/SE`. The p value is the lower-tail probability.
- Right-sided (noninferiority): `H_0 : θ ≤ θ_0 - δ` versus `H_1 : θ > θ_0 - δ`; `t = (Q - θ_0 + δ)/SE`. The p value is the upper-tail probability.
- Two-sided (equivalence): `H_0 : |θ - θ_0| ≥ δ` versus `H_1 : |θ - θ_0| < δ`; `t = (|Q - θ_0| - δ)/SE`. The p value is the lower-tail probability.

**Plots:** The `plot` method for "lsmobj" or "summary.ref.grid" objects (but not "ref.grid" objects themselves) produces a plot displaying confidence intervals for the estimates. If any by variables are in force, the plot is divided into separate panels. These functions use the `dotplot` function, and thus require that the `lattice` package be installed. For "summary.ref.grid" objects, the ... arguments in plot are passed only to `dotplot`, whereas for "lsmobj" objects, the object is updated using ... before summarizing and plotting.

In plots with `comparisons = TRUE`, the resulting arrows are only approximate, and in some cases may fail to accurately reflect the pairwise comparisons of the estimates – especially when estimates having large and small standard errors are intermingled in just the wrong way.

**Re-gridding:** The `regrid` function reparameterizes an existing `ref.grid` so that its `linfct` slot is the identity matrix and its `bhat` slot consists of the estimates at the grid points. If `transform` is `TRUE`, the inverse transform is applied to the estimates. Outwardly, the summary after applying
regrid is identical to what it was before (using ‘type=”response”’ if transform is TRUE). But
subsequent contrasts will be conducted on the transformed scale – which is the reason this function
exists. See the example below. In cases where the degrees of freedom depended on the linear
function being estimated, the d.f. from the reference grid are saved, and a kind of “containment”
method is substituted in the returned object whereby the calculated d.f. for a new linear function
will be the minimum d.f. among those having nonzero coefficients. This is kind of an ad hoc
method, and it can over-estimate the degrees of freedom in some cases.

MCMC samplers: When the object’s post.beta slot is non-trivial, as.mcmc will return an mcmc or
mcmc.list object that can be summarized or plotted using methods in the coda package. Alternatively, as.stanfit will return a stanfit object that can be summarized or plotted using methods
in the rstan package. You may use any of these functions regardless of what packages were originally used to implement the MCMC method. In these functions, post.beta is transformed by
post-multiplying it by t(linfct), creating a sample from the posterior distribution of LS means.
In as.mcmc, if sep.chains is TRUE and there is in fact more than one chain, an mcmc.list is re-
turned with each chain’s results. The as.mcmc.list method is guaranteed to return an mcmc.list,
even if it comprises just one chain. Note that stanfit objects are designed already for multiple
chains.

Value
The summary method for "ref.grid" objects returns an object of class "summary.ref.grid",
which extends "data.frame". xtable returns an object of class "xtable.lsm", as explained in
details. plot returns an object of class "trellis". vcov returns the covariance matrix of the
product of the object’s linfct and bhat slots. as.mcmc returns a coda mcmc object.

Author(s)
Russell V. Lenth

See Also
Methods for the closely related "lsmobj" class can be found in contrast, cld, and glht. For more
on Bonferroni-based P-value adjustments, see p.adjust. Also, test and confint are essentially
front-ends for summary, so additional examples may be found there.

Examples

require(lsmeans)
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.rg <- ref.grid(warp.lm)
str(warp.rg)
levels(warp.rg)
summary(warp.rg)

summary(warp.rg, by = "wool",
       infer = c(TRUE, FALSE), level = .90, adjust = "sidak")

# Do all pairwise comparisons within rows or within columns,
# all considered as one family of tests:
w.t <- pairs(lsmmeans(warp, ~ wool | tension))
t.w <- pairs(lsmmeans(warp, ~ tension | wool))
rtbind(w.t, t.w)

# Transformed response
sqwarp.rg <- ref.grid(update(warp.lm, sqrt(breaks) ~ .))
summary(sqwarp.rg)

# Back-transformed results - compare with summary of 'warp.rg'
summary(sqwarp.rg, type = "response")

# But differences of sqrts can't be back-transformed
summary(pairs(sqwarp.rg, by = "wool"), type = "response")

# We can do it via regrid
sqwarp.rg2 <- regrid(sqwarp.rg)
summary(sqwarp.rg2) # same as for sqwarp.rg with type = "response"
pairs(sqwarp.rg2, by = "wool")

# Logistic regression
# Reshape the Titanic data
Titan <- do.call("expand.grid", dimnames(Titanic)[-4])
Titan$Died <- matrix(Titanic, ncol=2)
Titan.glm <- glm(Died ~ (Class + Sex + Age)^2,
  family = binomial, data = Titan)
Titan.lsm <- lsmmeans(Titan.glm, ~ Class|Sex, at = list(Age="Adult"))
summary(Titan.lsm, type="response")
summary(pairs(Titan.lsm), type="response")

# Nonsuperiority test: Is any class no more likely to die than
# the 1st class passengers?
summary(contrast(Titan.lsm, "trt.vs.ctrl1"), delta = 1,
  adjust = "none", side = "<")

# Plot 90% CIs on the response scale
plot(Titan.lsm, type = "response", level = .90,
  xlab = "Predicted probability of drowning")

---

Set or retrieve options for objects and summaries in lsmeans

Description

Objects of class ref.grid or lsmobj contain several settings in their "misc" slot that affect primarily the defaults used by summary. This update method allows them to be changed more safely than by modifying this slot directly.

In addition, the user may set defaults for all objects using 'options(lsmeans = ...)', or more conveniently using the lsm.options and get.lsm.option functions documented here (or its courtesy wrappers, pmm.options and get.pmm.option for those who dislike the 'least-squares means' terminology).
Usage

## S3 method for class 'ref.grid'
update(object, ..., silent = FALSE)

lsm.options(...)
get.lsm.option(x, default = defaults.lsm[[x]])

pmm.options(...)
get.pmm.option(...)

Arguments

- **object**: An object of class ref.grid (or its extension, lsmobj).
- **...**: Arguments specifying elements' names and their new values.
- **silent**: If FALSE, a message is displayed for any unmatched names.
- **x**: Character string holding an option name for lsm.options.
- **default**: Return value if x is not found.

Details

**Using update**: In update, the names in ... are partially matched against those that are valid, and if a match is found, it adds or replaces the current setting. The valid names are

- **nesting** (named list) specifies the nesting structure. The names are those of nested factors, and the elements are character vectors of the factors they are nested in. See the nesting argument and Details section of ref.grid. The current nesting structure is displayed by link(str).

- **tran, tran2** (list or character) specifies the transformation which, when inverted, determines the results displayed by summary, predict, or lsmip when type="response". The value may be the name of a standard transformation from make.link or additional ones supported by name, such as log2; or, for a custom transformation, a list containing at least the functions linkinv (the inverse of the transformation) and mu.eta (the derivative thereof). The make.tran function returns such lists for a number of popular transformations. See the help page of make.tran for details as well as information on the additional named transformations that are supported. tran2 is just like tran except it is a second transformation (i.e., a response transformation in a generalized linear model).

- **tran.mult**: Multiple for tran. For example, for the response transformation ‘2*sqrt(y)’ (or ‘sqrt(y) + sqrt(y + 1)’, for that matter), we should have tran = "sqrt" and tran.mult = 2. If absent, a multiple of 1 is assumed.

- **estName** (character) is the column label used for displaying predictions or LS means.

- **inv.lbl** (character) is the column label to use for predictions or LS means when type="response".

- **by.vars** (character vector or NULL) the variables used for grouping in the summary, and also for defining subfamilies in a call to contrast.

- **pri.vars** (character vector) are the names of the grid variables that are not in by.vars. Thus, the combinations of their levels are used as columns in each table produced by summary.

- **alpha** (numeric) is the default significance level for tests, in summary as well as cld and plot when ‘intervals = TRUE’

- **adjust** (character) is the default for the adjust argument in summary.
estType (character) is the type of the estimate. It should match one of ‘c("prediction","contrast","pairs")’. This is used along with "adjust" to determine appropriate adjustments to P values and confidence intervals.

famSize (integer) is the nmeans parameter for ptukey when adjust="tukey".

infer (logical vector of length 2) is the default value of infer in summary.

level (numeric) is the default confidence level. level, in summary

df (numeric) overrides the default degrees of freedom with a specified single value.

null (numeric) null hypothesis for summary or test (taken to be zero if missing).

side (numeric or character) side specification for for summary or test (taken to be zero if missing).

delta (numeric) delta specification for summary or test (taken to be zero if missing).

predict.type (character) sets the default method of displaying predictions in summary, predict, and lsmip. Valid values are "link" (with synonyms "lp" and "linear"), or "response".

avgd.over (character vector) are the names of the variables whose levels are averaged over in obtaining marginal averages of predictions, i.e., LS means. Changing this might produce a misleading printout, but setting it to character(0) will suppress the “averaged over” message in the summary.

initMesg (character) is a string that is added to the beginning of any annotations that appear below the summary display.

methDesc (character) is a string that may be used for creating names for a list of lsmobj objects.

(any slot name) If the name matches an element of slotNames(object), that slot is replaced by the supplied value, if it is of the required class (otherwise an error occurs). Note that all the other possibilities above refer to elements of misc: hence, you probably don’t want to replace misc itself. The user must be very careful in replacing slots because they are interrelated: for example, the levels and grid slots must involve the same variable names, and the lengths and dimensions of grid, linfct, bhat, and V must conform.

Using lsm.options: In lsm.options, we may set or change the default values for the above attributes in the lsmobj option list(see options). Currently, the following elements of this list are used if specified:

ref.grid A named list of defaults for objects created by ref.grid. This could affect other objects as well. For example, if lsmobj is called with a fitted model object, it calls ref.grid and this option will affect the resulting lsmobj object.

lsm means A named list of defaults for objects created by lsmobj (or lstrends).

contrast A named list of defaults for objects created by contrast (or pairs).

summary A named list of defaults used by the methods summary, predict, and lsmip. The only option that can affect the latter two is "predict.method".

estble.tol Tolerance for determining estimability in rank-deficient cases. If absent, the value in defaults.lsm$estble.tol) is used.

(others) Other options may be accessed by support code for particular model classes (see models).

For example, the lmer.df, disable.pbkrtest, and pbkrtest.limit options affect how degrees of freedom are computed for lmerMod objects (lme4 package).

Value

update returns a copy of object with its "misc" slot modified (and perhaps other slots). lsm.options returns the current options (same as the result of ‘getOption("lsmoptions")’) – invisibly, unless called with no arguments.
Note

If a call to `lsmeans`, `contrast`, or `ref.grid` contains a non-NULL options list, those options are passed in a call to `update` on the constructed object before it is returned. This allows you, for example, to override the defaults used by `summary`. In addition, user defaults may be set using an `link(options)` setting for "lsmeans". It should be a list with one or more named elements `lsmeans`, `contrast`, or `ref.grid`, used for setting the defaults for objects constructed by functions of these same names. Note that options can get “inherited”. See the examples.

Unlike the update method for model classes (lm, glm, etc.), this does not re-fit or re-estimate anything; but it does affect how object is treated by other methods for its class.

Author(s)
Russell V. Lenth

See Also
`summary`, `make.tran`

Examples

```r
# An altered log transformation
warp.lml <- lm(log(breaks + 1) ~ wool*tension, data = warpbreaks)
rg1 <- update(ref.grid(warp.lml),
   tran = list(linkinv = function(eta) exp(eta) - 1,
              mu.eta = function(eta) exp(eta)),
   inv.lbl = "pred.breaks")
summary(rg1, type = "response")

## Not run:
lsm.options(ref.grid = list(level = .90),
    contrast = list(infer = c(TRUE,FALSE)),
    estble.tol = 1e-6)
## Sets default confidence level to .90 for objects created by ref.grid
## AS WELL AS lsmeans called with a model object (since it creates a
## reference grid). In addition, when we call 'contrast', 'pairs', etc.,
## confidence intervals rather than tests are displayed by default.

## Not run

## Not run:
lsm.options(disable.pbkrtest = TRUE)
## This forces use of asymptotic methods for lmerMod objects.
## Set to FALSE or NULL to re-enable using pbkrtest.

## Not run

## See tolerance being used for determining estimability
get.lsm.option("estble.tol")
```
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