Package ‘emmeans’

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emmeans-package

Estimated marginal means (aka Least-squares means)

Description

This package provides methods for obtaining estimated marginal means (EMMs, also known as least-squares means) for factor combinations in a variety of models. Supported models include [generalized linear] models, models for counts, multivariate, multinomial and ordinal responses, survival models, GEEs, and Bayesian models. For the latter, posterior samples of EMMs are provided. The package can compute contrasts or linear combinations of these marginal means with various multiplicity adjustments. One can also estimate and contrast slopes of trend lines. Some graphical displays of these results are provided.

Overview

Vignettes A number of vignettes are provided to help the user get acquainted with the emmeans package and see some examples.

Concept Estimated marginal means (see Searle et al. 1980) 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 the marginal means you would have observed that the data arisen from a balanced experiment. Earlier developments regarding these techniques were developed in a least-squares context and are sometimes referred to as “least-squares means”. Since its early development, the concept has expanded far beyond least-squares settings.

Reference grids The implementation in emmeans 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 estimated marginal means (or EMMs) are defined as averages of these predictions over zero or more dimensions of the grid. The function ref_grid explicitly creates a reference grid that can subsequently be used to obtain least-squares means. The object returned by ref_grid is of class "emmGrid", the same class as is used for estimated marginal means (see below). Our reference-grid framework expands slightly upon Searle et al.’s definitions of EMMs, in that it is possible to include multiple levels of covariates in the grid.
Models supported  As is mentioned in the package description, many types of models are supported by the package. See vignette("models", "emmeans") for full details. Some models may require other packages be installed in order to access all of the available features. For models not explicitly supported, it may still be possible to do basic post hoc analyses of them via the `qdr` function.

Estimated marginal means  The `emmeans` function computes EMMs given a fitted model (or a previously constructed `emmGrid` object), using a specification indicating what factors to include. The `emtrends` function creates the same sort of results for estimating and comparing slopes of fitted lines. Both return an `emmGrid` object.

Summaries and analysis  The `summary.emmGrid` method may be used to display an `emmGrid` object. Special-purpose summaries are available via `confint.emmGrid` and `test.emmGrid`, 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 for `emmGrid` objects 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 `emmGrid` object is returned, which can be summarized or further analyzed. For convenience, a `pairs.emmGrid` method is provided for the case of pairwise comparisons.

Graphs  The `plot.emmGrid` method will display side-by-side confidence intervals for the estimates, and/or “comparison arrows” whereby the *P* values of pairwise differences can be observed by how much the arrows overlap. The `emmip` function displays estimates like an interaction plot, multi-paneled if there are by variables. These graphics capabilities require the `lattice` package be installed.

MCMC support  When a model is fitted using MCMC methods, the posterior chains(s) of parameter estimates are retained and converted into posterior samples of EMMs or contrasts thereof. These may then be summarized or plotted like any other MCMC results, using tools in, say `coda` or `bayesplot`.

multcomp interface  The `as.glht` function and `glht` method for `emmGrids` 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 `emm` function that works as an alternative to `mcp` in a call to `glht`.

---

**as.list.emmGrid**

*Convert to and from `emmGrid` objects*

**Description**

These are useful utility functions for creating a compact version of an `emmGrid` object that may be saved and later reconstructed, or for converting old `ref.grid` or `lsmobj` objects into `emmGrid` objects.
Usage

## S3 method for class 'emmGrid'

as.list(x, model.info.slot = FALSE, ...)

as.emm_list(object, ...)

as.emmGrid(object, ...)

Arguments

x
An emmGrid object

model.info.slot
Logical value: Include the model.info slot? Set this to TRUE if you want to preserve the original call and information needed by the submodel option. If FALSE, only the nesting information (if any) is saved

... In as.emmGrid, additional arguments passed to update.emmGrid before returning the object. This argument is ignored in as.list.emmGrid

object
Object to be converted to class emmGrid. It may be a list returned by as.list.emmGrid, or a ref.grid or lsmobj object created by emmeans's predecessor, the lsmeans package. An error is thrown if object cannot be converted.

Details

An emmGrid object is an S4 object, and as such cannot be saved in a text format or saved without a lot of overhead. By using as.list, the essential parts of the object are converted to a list format that can be easily and compactly saved for use, say, in another session or by another user. Providing this list as the arguments for emmobj allows the user to restore a working emmGrid object.

Value

as.list.emmGrid returns an object of class list.

as.emm_list returns an object of class emm_list.

as.emmGrid returns an object of class emmGrid. However, in fact, both as.emmGrid and as.emm_list check for an attribute in object to decide whether to return an emmGrid or emm_list) object.

See Also

emmobj

Examples

pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
pigs.sav <- as.list(ref_grid(pigs.lm))
pigs.anew <- as.emmGrid(pigs.sav)
emmeans(pigs.anew, "source")
Description

When a model is fitted using Markov chain Monte Carlo (MCMC) methods, its reference grid contains a post.beta slot. These functions transform those posterior samples to posterior samples of EMMs or related contrasts. They can then be summarized or plotted using, e.g., functions in the coda package.

Usage

```r
## S3 method for class 'emmGrid'
as.mcmc(x, names = TRUE, sep.chains = TRUE, likelihood,
        NE.include = FALSE, ...)

## S3 method for class 'emm_list'
as.mcmc(x, which = I_bet(1), ...)

## S3 method for class 'emmGrid'
as.mcmc.list(x, names = TRUE, ...)

## S3 method for class 'emm_list'
as.mcmc.list(x, which = I_bet(1), ...)
```

Arguments

- **x** An object of class emmGrid
- **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 EMMs posteriors in each chain.
- **likelihood** Character value or function. If given, simulations are made from the corresponding posterior predictive distribution. If not given, we obtain the posterior distribution of the parameters in object. See Prediction section below.
- **NE.include** Logical value. If TRUE, non-estimable columns are kept but returned as columns of NA values (this may create errors or warnings in subsequent analyses using, say, coda). If FALSE, non-estimable columns are dropped, and a warning is issued. (If all are non-estimable, an error is thrown.)
- **which** item in the emm_list to use
- **...** arguments passed to other methods
as.mcmc.emmGrid

Value

An object of class mcmc or mcmc.list.

Details

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. 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 returned with each chain's results. The as.mcmc.list method is guaranteed to return an mcmc.list, even if it comprises just one chain.

Prediction

When likelihood is specified, it is used to simulate values from the posterior predictive distribution corresponding to the given likelihood and the posterior distribution of parameter values. Denote the likelihood function as $f(y|\theta, \phi)$, where $y$ is a response, $\theta$ is the parameter estimated in object, and $\phi$ comprises zero or more additional parameters to be specified. If likelihood is a function, that function should take as its first argument a vector of $\theta$ values (each corresponding to one row of object@grid). Any $\phi$ values should be specified as additional named function arguments, and passed to likelihood via .... This function should simulate values of $y$.

A few standard likelihoods are available by specifying likelihood as a character value. They are:

"normal" The normal distribution with mean $\theta$ and standard deviation specified by additional argument sigma

"binomial" The binomial distribution with success probability theta, and number of trials specified by trials

"poisson" The Poisson distribution with mean theta (no additional parameters)

"gamma" The gamma distribution with scale parameter $\theta$ and shape parameter specified by shape

Examples

```r
if(requireNamespace("coda")) {
  ### A saved reference grid for a mixed logistic model (see lme4::cbpp)
  cbpp.rg <- do.call(emmobj,
                     readRDS(system.file("extdata", "cbpplist", package = "emmeans")))
  # Predictive distribution for herds of size 20
  # (perhaps a bias adjustment should be applied; see "sophisticated" vignette)
  pred.incidence <- coda::as.mcmc(regrid(cbpp.rg), likelihood = "binomial", trials = 20)
  summary(pred.incidence)
}
```
auto.noise Auto Pollution Filter Noise

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 (but see note) - 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.

Note

While the data source claims that noise is measured in decibels, the values are implausible. I believe that these measurements are actually in tenths of dB (centibels?). Looking at the values in the dataset, note that every measurement ends in 0 or 5, and it is reasonable to believe that measurements are accurate to the nearest half of a decibel.

Source

The dataset was obtained from the Data and Story Library (DASL) at Carnegie-Mellon University. Apparently it has since been removed. The original dataset was altered by assigning meaningful names to the factors and sorting the observations in random order as if this were the run order of the experiment.
Examples

# (Based on belief that noise/10 is in decibel units)
noise.lm <- lm(noise/10 ~ size * type * side, data = auto.noise)

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

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

cld.emmGrid

Compact letter displays

Description

A method for \texttt{multcomp::cld()} is provided for users desiring to produce compact-letter displays (CLDs). This method uses the Piepho (2004) algorithm (as implemented in the \texttt{multcompView} package) to generate a compact letter display of all pairwise comparisons of estimated marginal means. The function obtains (possibly adjusted) P values for all pairwise comparisons of means, using the \texttt{contrast} function with \texttt{method = "pairwise"}. When a P value exceeds \texttt{alpha}, then the two means have at least one letter in common.

Usage

\begin{verbatim}
## S3 method for class 'emmGrid'
cld(object, details = FALSE, sort = TRUE, by, 
a alpha = 0.05, Letters = c("1234567890", LETTERS, letters), 
reversed = FALSE, ...) 

## S3 method for class 'emm_list'
cld(object, ..., which = I_bet(1))
\end{verbatim}

Arguments

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>An object of class \texttt{emmGrid}</td>
</tr>
<tr>
<td>details</td>
<td>Logical value determining whether detailed information on tests of pairwise comparisons is displayed</td>
</tr>
<tr>
<td>sort</td>
<td>Logical value determining whether the EMMs are sorted before the comparisons are produced. When \texttt{TRUE}, the results are displayed according to \texttt{reversed}.</td>
</tr>
<tr>
<td>by</td>
<td>Character value giving the name or names of variables by which separate families of comparisons are tested. If \texttt{NULL}, all means are compared. If \texttt{missing}, the object’s \texttt{by.vars} setting, if any, is used.</td>
</tr>
<tr>
<td>alpha</td>
<td>Numeric value giving the significance level for the comparisons</td>
</tr>
<tr>
<td>Letters</td>
<td>Character vector of letters to use in the display. Any strings of length greater than 1 are expanded into individual characters</td>
</tr>
</tbody>
</table>
reversed Logical value (passed to multcompView::multcompLetters.) 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 Which element of the emm_list object to process (If length exceeds one, only the first one is used)

Note

We warn that such displays encourage a poor practice in interpreting significance tests. CLDs are misleading because they visually group means with comparisons \( P > \alpha \) as though they are equal, when in fact we have only failed to prove that they differ. As alternatives, consider pwpp (graphical display of \( P \) values) or pwpm (matrix display).

References


Examples

```r
if(requireNamespace("multcomp")) {
  pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
  pigs.emm <- emmeans(pigs.lm, "percent", type = "response")
  multcomp::cld(pigs.emm, alpha = 0.10, Letters = LETTERS)
}
```

comb_facs

**Manipulate factors in a reference grid**

**Description**

These functions manipulate the velels of factors comprising a reference grid by combining factor levels, splitting a factor’s levels into combinations of newly-defined factors, or by creating a grouping factor in which factor(s) levels are nested.

**Usage**

```r
comb_facs(object, facs, newname = paste(facs, collapse = "."),
  drop = FALSE, ...)

split_fac(object, fac, newfacs, ...)

add_grouping(object, newname, refname, newlevs, ...)
```
**Arguments**

- **object**: An object of class `emmGrid`
- **facs**: Character vector. The names of the factors to combine
- **newname**: Character name of grouping factor to add (different from any existing factor in the grid)
- **drop**: Logical value. If TRUE, any levels of the new factor that are dropped if all occurrences in the newly reconstructed object have weight zero. If FALSE, all levels are retained. (This argument is ignored if there is no `wgt.` column in `object@grid`.)
- **...**: arguments passed to other methods
- **fac**: The name of a factor that is part of the grid in `object`
- **newfacs**: A named list with the names of new factors and their levels. The names must not already exist in the object, and the product of the lengths of the levels must equal the number of levels of `fac`.
- **refname**: Character name(s) of the reference factor(s)
- **newlevs**: Character vector or factor of the same length as that of the (combined) levels for `refname`. The grouping factor `newname` will have the unique values of `newlevs` as its levels. The order of levels in `newlevs` is the same as the order of the level combinations produced by `expand.grid` applied to the levels of `refname` – that is, the first factor’s levels change the fastest and the last one’s vary the slowest.

**Value**

A modified object of class `emmGrid`

**The `comb_facs` function**

`comb_facs` combines the levels of factors into a single factor in the reference grid (similar to `interaction`). This new factor replaces the factors that comprise it.

Additional note: The choice of whether to drop levels or not can make a profound difference. If the goal is to combine factors for use in `joint_tests`, we advise **against** `drop = TRUE` because that might change the weights used in deriving marginal means. If combining factors in a nested structure, dropping unused cases can considerably reduce the storage required.

**The `split_fac` function**

The levels in `newfacs` are expanded via `expand.grid` into combinations of levels, and the factor `fac` is replaced by those factor combinations. Unlike `add_grouping`, this creates a crossed, rather than a nested structure. Note that the order of factor combinations is systematic with the levels of first factor in `newfacs` varying the fastest; and those factor combinations are assigned respectively to the levels of `fac` as displayed in `str(object)`.

*comb_facs*
The `add_grouping` function

This function adds a grouping factor to an existing reference grid or other `emmGrid` object, such that the levels of one or more existing factors (call them the reference factors) are mapped to a smaller number of levels of the new grouping factor. The reference factors are then nested in a new grouping factor named `newname`, and a new nesting structure `refname %in% newname`. This facilitates obtaining marginal means of the grouping factor, and contrasts thereof.

Additional notes: By default, the levels of `newname` will be ordered alphabetically. To dictate a different ordering of levels, supply `newlevs` as a factor having its levels in the desired order.

When `refname` specifies more than one factor, this can fundamentally (and permanently) change what is meant by the levels of those individual factors. For instance, in the `gwrg` example below, there are two levels of `wool` nested in each `prod`; and that implies that we now regard these as four different kinds of wool. Similarly, there are five different tensions (L, M, H in prod 1, and L, M in prod 2).

Examples

```r
mtcars.lm <- lm(mpg ~ factor(vs)+factor(cyl)*factor(gear), data = mtcars)
  (v.c.g <- ref_grid(mtcars.lm))
  (v.cg <- comb_facs(v.c.g, c("cyl", "gear")))

  # One use is obtaining a single test for the joint contributions of two factors:
  joint_tests(v.c.g)

  joint_tests(v.cg)

  # undo the 'comb_facs' operation:
  split_fac(v.cg, "cyl.gear", list(cyl = c(4, 6, 8), gear = 3:5))

IS.glm <- glm(count ~ spray, data = InsectSprays, family = poisson)
IS.emm <- emmeans(IS.glm, "spray")
IS.new <- split_fac(IS.emm, "spray", list(A = 1:2, B = c("low", "med", "hi")))
str(IS.new)

fiber.lm <- lm(strength ~ diameter + machine, data = fiber)
  ( frg <- ref_grid(fiber.lm) )

  # Suppose the machines are two different brands
  brands <- factor(c("FiberPro", "FiberPro", "Acme"), levels = c("FiberPro", "Acme"))
  ( gfrg <- add_grouping(frg, "brand", "machine", brands ) )

  emmeans(gfrg, "machine")

  emmeans(gfrg, "brand")

  ### More than one reference factor
  warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
  gwrg <- add_grouping(ref_grid(warp.lm),
      "prod", c("tension", "wool"), c(2, 1, 1, 1, 2, 1))
  # level combinations: LA MA HA LB MB HB
```
### Description

These methods provide for follow-up analyses of `emmGrid` objects: Contrasts, pairwise comparisons, tests, and confidence intervals. They may also be used to compute arbitrary linear functions of predictions or EMMs.

### Usage

```r
contrast(object, ...)  
## S3 method for class 'emmGrid'
contrast(object, method = "eff", interaction = FALSE, by,  
     offset = NULL, scale = NULL, name = "contrast",  
     options = get_emm_option("contrast"), type, adjust, simple,  
     combine = FALSE, ratios = TRUE, parens, enhance.levels = TRUE, ...)

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

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

### Arguments

- **object** 
  An object of class `emmGrid`

- **...** 
  Additional arguments passed to other methods

- **method** 
  Character value giving the root name of a contrast method (e.g. "pairwise" – see `emmc-functions`). Alternatively, a function of the same form, or a named list of coefficients (for a contrast or linear function) that must each conform to the number of results in each by group. In a multi-factor situation, the factor levels are combined and treated like a single factor.

- **interaction** 
  Character vector, logical value, or list. If this is specified, method is ignored. See the “Interaction contrasts” section below for details.

- **by** 
  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.
offset, scale Numeric vectors of the same length as each by group. The scale values, if supplied, multiply their respective linear estimates, and any offset values are added. Scalar values are also allowed. (These arguments are ignored when interaction is specified.)

name Character name to use to override the default label for contrasts used in table headings or subsequent contrasts of the returned object.

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

type Character: prediction type (e.g., "response") – added to options

adjust Character: adjustment method (e.g., "bonferroni") – added to options

simple Character vector or list: Specify the factor(s) not in by, or a list thereof. See the section below on simple contrasts.

combine Logical value that determines what is returned when simple is a list. See the section on simple contrasts.

ratios Logical value determining how log and logit transforms are handled. These transformations are exceptional cases in that there is a valid way to back-transform contrasts: differences of logs are logs of ratios, and differences of logits are odds ratios. If ratios = TRUE and summarized with type = "response", contrast results are back-transformed to ratios whenever we have true contrasts (coefficients sum to zero). For other transformations, there is no natural way to back-transform contrasts, so even when summarized with type = "response", contrasts are computed and displayed on the linear-predictor scale. Similarly, if ratios = FALSE, log and logit transforms are treated in the same way as any other transformation.

parens character or NULL. If a character value, the labels for levels being contrasted are parenthesized if they match the regular expression in parens[1] (via grep). The default is emm_option("parens"). Optionally, parens may contain second and third elements specifying what to use for left and right parentheses (default "(" and ")"). Specify parens = NULL or parens = "a^" (which won’t match anything) to disable all parenthesization.

enhance.levels character or logical. If character, the levels of the named factors that are contrasted are enhanced by appending the name of the factor; e.g., if a factor named "trt" has levels A and codeB, a trt comparison is labeled trtA - trtB. If enhance.levels is logical, then if TRUE (the default), only factors with numeric levels are enhanced; and of course if FALSE, no levels are enhanced. The levels of by variables are not enhanced, and any names of factors that don’t exist are silently ignored. To enhance the labels beyond what is done here, change them directly via levels<-

x An emmGrid object

reverse Logical value - determines whether to use "pairwise" (if TRUE) or "revpairwise" (if FALSE).

Value

contrast and pairs return an object of class emmGrid. Its grid will correspond to the levels of the contrasts and any by variables. The exception is that an emm_list object is returned if simple is a list and complete is FALSE.
coef returns a data.frame containing the object's grid, along with columns named c.1, c.2, ... containing the contrast coefficients. If

**Pairs method**

The call `pairs(object)` is equivalent to `contrast(object, method = "pairwise")`; and `pairs(object, reverse = TRUE)` is the same as `contrast(object, method = "revpairwise")`.

**Interaction contrasts**

When `interaction` is specified, interaction contrasts are computed. Specifically 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. Any named elements of `interaction` are assigned to contrast methods; others are assigned in order of appearance in `object@levels`. The contrast factors in the resulting `emmGrid` object are ordered the same as in `interaction`.

`interaction` may be a character vector or list of valid contrast methods (as documented for the `method` argument). If the vector or list is shorter than the number needed, it is recycled. Alternatively, if the user specifies `contrast = TRUE`, the contrast specified in `method` is used for all factors involved.

**Simple contrasts**

`simple` is essentially the complement of `by`: When `simple` is a character vector, `by` is set to all the factors in the grid except those in `simple`. If `simple` is a list, each element is used in turn as `simple`, and assembled in an "emm_list". To generate all simple main effects, use `simple = "each"` (this works unless there actually is a factor named "each"). Note that a non-missing `simple` will cause `by` to be ignored.

Ordinarily, when `simple` is a list or "each", the return value is an `emm_list` object with each entry in correspondence with the entries of `simple`. However, with `combine = TRUE`, the elements are all combined into one family of contrasts in a single `emmGrid` object using `rbind.emmGrid`. In that case, the `adjust` argument sets the adjustment method for the combined set of contrasts.

**Note**

When `object` has a nesting structure (this can be seen via `str(object)`), then any grouping factors involved are forced into service as `by` variables, and the contrasts are thus computed separately in each nest. This in turn may lead to an irregular grid in the returned `emmGrid` object, which may not be valid for subsequent `emmeans` calls.

**Examples**

```r
warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)
warp.emm <- emmeans(warp.lm, ~ tension | wool)
contrast(warp.emm, "poly")  # inherits 'by = "wool"' from warp.emm
pairs(warp.emm)
```
# Effects (dev from mean) of the 6 factor combs, with enhanced levels:
contrast(warp.emm, "eff", by = NULL, 
enhance.levels = c("wool", "tension"))
pairs(warp.emm, simple = "wool") # same as pairs(warp.emm, by = "tension")

# Do all "simple" comparisons, combined into one family
pairs(warp.emm, simple = "each", combine = TRUE)

## Not run:
## Note that the following are NOT the same:
contrast(warp.emm, simple = c("wool", "tension"))
contrast(warp.emm, simple = list("wool", "tension"))
## The first generates contrasts for combinations of wool and tension
## (same as by = NULL)
## The second generates contrasts for wool by tension, and for
## tension by wool, respectively.

## End(Not run)

# An interaction contrast for tension:wool
tw.emm <- contrast(warp.emm, interaction = c("tension = "poly", "wool = "consec"),
                   by = NULL)
tw.emm       # see the estimates
go(tw.emm)    # see the contrast coefficients

# Use of scale and offset
# an unusual use of the famous stack-loss data...
mod <- lm(Water.Temp ~ poly(stack.loss, degree = 2), data = stackloss)
(emm <- emmeans(mod, "stack.loss", at = list(stack.loss = 10 * (1:4))))
# Convert results from Celsius to Fahrenheit:
confint(contrast(emm, "identity", scale = 9/5, offset = 32))

---

### Contrast families

**Description**

Functions with an extension of .emmc provide for named contrast families. One of the standard ones documented here may be used, or the user may write such a function.

**Usage**

```r
cpyairwise.emmc(levs, exclude = integer(0), include, ...)

revpairwise.emmc(levs, exclude = integer(0), include, ...)
```
tukey.emmc(levs, reverse = FALSE, ...)

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

trt.vs.ctrl1.emmc(levs, ref = 1, reverse = FALSE, exclude = integer(0), include, ...)

trt.vs.ctrl1.emmc(levs, ref = 1, ...)

trt.vs.ctrlk.emmc(levs, ref = length(levs), ...)

dunnett.emmc(levs, ref = 1, ...)

eff.emmc(levs, exclude = integer(0), include, wts = rep(1, length(levs)), ...)

del.eff.emmc(levs, exclude = integer(0), include, wts = rep(1, length(levs)), ...)

consec.emmc(levs, reverse = FALSE, exclude = integer(0), include, ...)

mean_chg.emmc(levs, reverse = FALSE, exclude = integer(0), include, ...)

identity.emmc(levs, exclude = integer(0), include, ...)

**Arguments**

- **levs**  Vector of factor levels
- **exclude**  integer vector of indices, or character vector of levels to exclude from consideration. These levels will receive weight 0 in all contrasts. Character levels must exactly match elements of levs.
- **include**  integer or character vector of levels to include (the complement of exclude). An error will result if the user specifies both exclude and include.
- **reverse**  Logical value to determine the direction of comparisons
- **max.degree**  Integer specifying the maximum degree of polynomial contrasts
- **ref**  Integer(s) or character(s) specifying which level(s) to use as the reference. Character values must exactly match elements of levs (including any enhancements – see examples)
- **wts**  Optional weights to use with eff.emmc and del.eff.emmc contrasts. These default to equal weights. If exclude or include are specified, wts may be either the same length as levs or the length of the included levels. In the former case, weights for any excluded levels are set to zero. wts has no impact on the results unless there are at least three levels included in the contrast.

**Details**

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

poly.emmc 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".

trt.vs.ctrl.emmc 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) (not the labels) of the reference level(s). trt.vs.ctrl1.emmc is the same as trt.vs.ctrl.emmc with a reference value of 1, and trt.vs.ctrlk.emmc is the same as trt.vs.ctrl with a reference value of length(levs). dunnett.emmc is the same as trt.vs.ctrl. The default multiplicity adjustment method is "dunnetttx", a close approximation to the Dunnett adjustment. Note in all of these functions, it is illegal to have any overlap between the ref levels and the exclude levels. If any is found, an error is thrown.

consec.emmc and mean_chg.emmc are useful for contrasting treatments that occur in sequence. For a factor with levels A, B, C, D, E, consec.emmc generates the comparisons B-A, C-B, and D-C, while mean_chg.emmc 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.

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

identity.emmc simply returns the identity matrix (as a data frame), minus any columns specified in exclude. It is potentially useful in cases where a contrast function must be specified, but none is desired.

### Value

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

### Note

Caution is needed in cases where the user alters the ordering of results (e.g., using the the "[...]" operator), because the contrasts generated depend on the order of the levels provided. For example, suppose trt.vs.ctrl1 contrasts are applied to two by groups with levels ordered (Ctrl, T1, T2) and (T1, T2, Ctrl) respectively, then the contrasts generated will be for (T1 - Ctrl, T2 - Ctrl) in the
first group and (T2 - T1, Ctrl - T1) in the second group, because the first level in each group is used as the reference level.

Examples

```r
code
warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)
warp.emm <- emmeans(warp.lm, ~ tension | wool)
contrast(warp.emm, "poly")
contrast(warp.emm, "trt.vs.ctrl", ref = "M")
## Not run:
## Same when enhanced labeling is used:
contrast(warp.emm, "trt.vs.ctrl",
        enhance.levels = "tension", ref = "tensionM")
## End(Not run)

# Comparisons with grand mean
contrast(warp.emm, "eff")
# Comparisons with a weighted grand mean
contrast(warp.emm, "eff", wts = c(2, 5, 3))

# Compare only low and high tensions
# Note pairs(emm, ...) calls contrast(emm, "pairwise", ...)
pairs(warp.emm, exclude = 2)
# (same results using exclude = "M" or include = c("L","H") or include = c(1,3))

### Setting up a custom contrast function
helmert.emmc <- function(levs, ...) {
    M <- as.data.frame(contr.helmert(levs))
    names(M) <- paste(levs[-1],"vs earlier")
    attr(M, "desc") <- "Helmert contrasts"
    M
}
contrast(warp.emm, "helmert")
## Not run:
# See what is used for polynomial contrasts with 6 levels
emmeans:::poly.emmc(1:6)
## End(Not run)
```

---

**eff_size**

*Calculate effect sizes and confidence bounds thereof*

**Description**

Standardized effect sizes are typically calculated using pairwise differences of estimates, divided by the SD of the population providing the context for those effects. This function calculates effect sizes from an `emmGrid` object, and confidence intervals for them, accounting for uncertainty in both the estimated effects and the population SD.
Usage

```r
eff_size(object, sigma, edf, method = "pairwise", ...)
```

Arguments

- `object`: an `emmGrid` object, typically one defining the EMMs to be contrasted. If instead, `class(object) == "emm_list"`, such as is produced by `emmeans(model, pairwise ~ treatment)`, a message is displayed; the contrasts already therein are used; and `method` is replaced by "identity".
- `sigma`: numeric scalar, value of the population SD.
- `edf`: numeric scalar that specifies the equivalent degrees of freedom for the `sigma`. This is a way of specifying the uncertainty in `sigma`, in that we regard our estimate of `sigma^2` as being proportional to a chi-square random variable with `edf` degrees of freedom. (`edf` should not be confused with the `df` argument that may be passed via `...` to specify the degrees of freedom to use in `t` statistics and confidence intervals.)
- `method`: the contrast method to use to define the effects. This is passed to `contrast` after the elements of `object` are scaled.
- `...`: Additional arguments passed to `contrast`

Details

Any by variables specified in `object` will remain in force in the returned effects, unless overridden in the optional arguments.

For models having a single random effect, such as those fitted using `lm`; in that case, the `stats::sigma` and `stats::df.residual` functions may be useful for specifying `sigma` and `edf`. For models with more than one random effect, `sigma` may be based on some combination of the random-effect variances.

Specifying `edf` can be rather unintuitive but is also relatively uncritical; but the smaller the value, the wider the confidence intervals for effect size. The value of `sqrt(2/edf)` can be interpreted as the relative accuracy of `sigma`; for example, with `edf = 50`, `sqrt(2/50) = 0.2`, meaning that `sigma` is accurate to plus or minus 20 percent. Note in an example below, we tried two different `edf` values as kind of a bracketing/sensitivity-analysis strategy. A value of `Inf` is allowable, in which case you are assuming that `sigma` is known exactly. Obviously, this narrows the confidence intervals for the effect sizes – unrealistically if in fact `sigma` is unknown.

Value

- an `emmGrid` object containing the effect sizes

Computation

This function uses calls to `regrid` to put the estimated marginal means (EMMs) on the log scale. Then an extra element is added to this grid for the log of `sigma` and its standard error (where we assume that `sigma` is uncorrelated with the log EMMs). Then a call to `contrast` subtracts `log(sigma)` from each of the log EMMs, yielding values of `log(EMM/sigma)`. Finally, the results are re-gridded back to the original scale and the desired contrasts are computed using `method`. In the log-scaling part, we actually rescale the absolute values and keep track of the signs.
Note

The effects are always computed on the scale of the linear-predictor; any response transformation or link function is completely ignored. If you wish to base the effect sizes on the response scale, it is not enough to replace object with regrid(object), because this back-transformation changes the SD required to compute effect sizes.

Disclaimer: There is substantial disagreement among practitioners on what is the appropriate sigma to use in computing effect sizes; or, indeed, whether any effect-size measure is appropriate for some situations. The user is completely responsible for specifying appropriate parameters (or for failing to do so).

The examples here illustrate a sobering message that effect sizes are often not nearly as accurate as you may think.

Examples

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

emm <- emmeans(fiber.lm, "machine")
eff_size(emm, sigma = sigma(fiber.lm), edf = df.residual(fiber.lm))

# or equivalently:
eff_size(pairs(emm), sigma(fiber.lm), df.residual(fiber.lm), method = "identity")
```

```r
### Mixed model example:
if (require(nlme)) withAutoprint({
  Oats.lme <- lme(yield ~ Variety + factor(nitro),
                  random = ~ 1 | Block / Variety,
                  data = Oats)
  # Combine variance estimates
  VarCorr(Oats.lme)
  (totSD <- sqrt(214.4724 + 109.6931 + 162.5590))
  # I figure edf is somewhere between 5 (Blocks df) and 51 (Resid df)
  emmV <- emmeans(Oats.lme, ~ Variety)
  eff_size(emmV, sigma = totSD, edf = 5)
  eff_size(emmV, sigma = totSD, edf = 51)
}, spaced = TRUE)

# Multivariate model for the same data:
MOats.lm <- lm(yield ~ Variety, data = MOats)
eff_size(emmeans(MOats.lm, "Variety"),
          sigma = sqrt(mean(sigma(MOats.lm)^2)), # RMS of sigma()
          edf = df.residual(MOats.lm))
```

---

Support for multcomp::glht
Description

These functions and methods provide an interface between emmeans and the multcomp::glht function for simultaneous inference provided by the multcomp package.

Usage

```r
emm(...)
```

```r
as.glht(object, ...)
```

## S3 method for class 'emmGrid'
```r
as.glht(object, ...)
```

Arguments

... In emm, the specs, by, and contr arguments you would normally supply to emmeans. Only specs is required. Otherwise, arguments are passed to other methods. You may also include a which argument; see Details.

object An object of class emmGrid or emm_list

Value

emm returns an object of an intermediate class for which there is a multcomp::glht method.

as.glht returns an object of class glht or glht_list according to whether object is of class emmGrid or emm_list. See Details below for more on glht_lists.

Details for emm

emm is meant to be called only from "glht" as its second (linfct) argument. It works similarly to multcomp::mcp, except with specs (and optionally by and contr arguments) provided as in a call to emmeans.

If the specifications in ... would result in a list (i.e., an emm_list object), then by default, only the last element of that list is passed to glht. However, if ... contains a which argument consisting of integer values, the list elements with those indexes are selected and combined and passed on to glht. No checking is done on whether the indexes are valid, and the keyword which must be spelled-out.

Details for as.glht

When no by variable is in force, we obtain a glht object; otherwise it is a glht_list. The latter is defined in emmeans, not multcomp, and is simply a list of glht objects. Appropriate convenience methods coef, confint, plot, summary, and vcov are provided, which simply apply the corresponding glht methods to each member.

Note

The multivariate-t routines used by glht require that all estimates in the family have the same integer degrees of freedom. In cases where that is not true, a message is displayed that shows what df is used. The user may override this via the df argument.
Examples

```r
if(require(multcomp, quietly = TRUE)) withAutoprint({
  # --- multcomp must be installed
  warp.lm <- lm(breaks ~ wool*tension, data = warpbreaks)
  # Using 'emm'
  summary(glht(warp.lm, emm(pairwise ~ tension | wool)))
  # Combine the means and the comparisons into one family
  summary(glht(warp.lm, emm(pairwise ~ tension | wool,
                            which = 1:2, by = "wool")))

  # Same as first example, but use an existing 'emmeans' result
  warp.emm <- emmeans(warp.lm, ~ tension | wool)
  summary(as.glht(pairs(warp.emm)))
  # Same contrasts, but treat as one family
  summary(as.glht(pairs(warp.emm), by = NULL))
}, spaced = TRUE)
```

emmeans

Estimated marginal means (Least-squares means)

Description

Compute estimated marginal means (EMMs) for specified factors or factor combinations in a linear model; and optionally, comparisons or contrasts among them. EMMs are also known as least-squares means.

Usage

```r
emmeans(object, specs, by = NULL, fac.reduce = function(coefs) apply(coefs,
  2, mean), contr, options = get_emm_option("emmeans"), weights, offset, ..., tran)
```

Arguments

- **object**: An object of class `emmGrid`; 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 `vignette("models", "emmeans")` for details.
- **specs**: A character vector specifying the names of the predictors over which EMMs are desired. specs may also be a formula or a list (optionally named) of valid specs. Use of formulas is described in the Overview section below.
- **by**: A character vector specifying the names of predictors to condition on.
- **fac.reduce**: A function that combines the rows of a matrix into a single vector. This implements the “marginal averaging” aspect of EMMs. 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 EMMs will not be interpretable. NOTE: If the weights argument is non-missing, fac.reduce is ignored.
emmeans

contr A character value or list specifying contrasts to be added. See `contrast`. NOTE: `contr` is ignored when `specs` is a formula.

options If non-NULL, a named list of arguments to pass to `update.emmGrid`, just after the object is constructed. (Options may also be included in `...`; see the ‘options’ section below.)

weights Character value, numeric vector, or numeric matrix specifying weights to use in averaging predictions. See “Weights” section below. Also, if object is not already a reference grid, weights (if it is character) is passed to `ref_grid` as `wt.nuis` in case nuisance factors are specified. We can override this by specifying `wt.nuis` explicitly. This more-or-less makes the weighting of nuisance factors consistent with that of primary factors.

offset Numeric vector or scalar. If specified, this adds an offset to the predictions, or overrides any offset in the model or its reference grid. If a vector of length differing from the number of rows in the result, it is subsetted or cyclically recycled.

... When object is not already a "emmGrid" object, these arguments are passed to `ref_grid`. Common examples are `at`, `cov.reduce`, `data`, `codetype`, `regrid`, `df`, `nesting`, and `vcov`. Model-type-specific options (see `vignette("models", "emmeans")`), commonly `mode`, may be used here as well. In addition, if the model formula contains references to variables that are not predictors, you must provide a `params` argument with a list of their names. These arguments may also be used in lieu of `options`. See the ‘Options’ section below.

tran Placeholder to prevent it from being included in `...`. If non-missing, it is added to `options`. See the ‘Options’ section.

Details

Users should also consult the documentation for `ref_grid`, because many important options for EMMs are implemented there, via the `...` argument.

Value

When `specs` is a character vector or one-sided formula, an object of class "emmGrid". A number of methods are provided for further analysis, including `summary.emmGrid`, `confint.emmGrid`, `test.emmGrid`, `contrast.emmGrid`, and `pairs.emmGrid`. When `specs` is a list or a formula having a left-hand side, the return value is an `emm_list` object, which is simply a list of `emmGrid` objects.

Overview

Estimated marginal means or EMMs (sometimes called least-squares means) are predictions from a linear model over a reference grid; or marginal averages thereof. The `ref_grid` function identifies/creates the reference grid upon which emmeans is based.

For those who prefer the terms “least-squares means” or “predicted marginal means”, functions `lsmeans` and `pmmeans` are provided as wrappers. See `wrappers`.
If `specs` is a formula, it should be of the form `~ specs`, `~ specs | by`, `contr ~ specs`, or `contr ~ specs | by`. The formula is parsed and the variables therein are used as the arguments `specs`, `by`, and `contr` as indicated. The left-hand side is optional, but if specified it should be the name of a contrast family (e.g., `pairwise`). Operators like `*` or `:` are needed in the formula to delineate names, but otherwise are ignored.

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

A number of standard contrast families are provided. They can be identified as functions having names ending in `.emmc` – see the documentation for `emmc-functions` for details – including how to write your own `.emmc` function for custom contrasts.

**Weights**

If `weights` is a vector, its length must equal the number of predictions to be averaged to obtain each EMM. 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 `emmeans` with `weights = "show.levels"`.

If `weights` is 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 EMMs, this causes the function to return a table showing the levels that are averaged over, in the order that 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.

**Offsets**

Unlike in `ref_grid`, an offset need not be scalar. If not enough values are supplied, they are cyclically recycled. For a vector of offsets, it is important to understand that the ordering of results goes with the first name in `specs` varying fastest. If there are any by factors, those vary slower than all the primary ones, but the first by variable varies the fastest within that hierarchy. See the examples.
Options and ...

Arguments that could go in options may instead be included in .... typically, arguments such as type, infer, etc. that in essence are passed to `summary.emmGrid`. Arguments in both places are overridden by the ones in ....

There is a danger that ... arguments could partially match those used by both ref_grid and update.emmGrid, creating a conflict. If these occur, usually they can be resolved by providing complete (or at least longer) argument names; or by isolating non-ref_grid arguments in options; or by calling ref_grid separately and passing the result as object. See a not-run example below.

Also, when specs is a two-sided formula, or contr is specified, there is potential confusion concerning which ... arguments apply to the means, and which to the contrasts. When such confusion is possible, we suggest doing things separately (a call to emmeans with no contrasts, followed by a call to contrast). We do treat for adjust as a special case: it is applied to the emmeans results only if there are no contrasts specified, otherwise it is passed to contrast.

See Also

`ref_grid`, `contrast`, vignette("models","emmeans")

Examples

```r
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
emmeans (warp.lm, ~ wool | tension)
# or equivalently emmeans(warp.lm, "wool", by = "tension")

# 'adjust' argument ignored in emmeans, passed to contrast part...
emmeans (warp.lm, poly ~ tension | wool, adjust = "sidak")

## Not run:
# 'adjust' argument NOT ignored ...
emmeans (warp.lm, ~ tension | wool, adjust = "sidak")

## End(Not run)

## Not run:
### Offsets: Consider a silly example:
emmeans(warp.lm, ~ tension | wool, offset = c(17, 23, 47)) @ grid
# note that offsets are recycled so that each level of tension receives
# the same offset for each wool.
# But using the same offsets with ~ wool | tension will probably not
# be what you want because the ordering of combinations is different.

## End(Not run)
```
Description

The `emmGrid` class encapsulates linear functions of regression parameters, defined over a grid of predictors. This includes reference grids and grids of marginal means thereof (aka estimated marginal means). Objects of class `emmGrid` may be used independently of the underlying model object. Instances are created primarily by `ref_grid` and `emmeans`, and several related functions.

Slots

- `model.info` list. Contains the elements `call` (the call that produced the model), `terms` (its terms object), and `xlev` (factor-level information).
- `roles` list. Contains at least the elements `predictors`, `responses`, and `multresp`. Each is a character vector of names of these variables.
- `grid` data.frame. Contains 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` list. Each entry is a character vector with the distinct levels of each variable in the reference grid. Note that grid is obtained by applying the function `expand.grid` to this list.
- `matlevs` list. Like `levels` but has the levels of any matrices in the original dataset. Matrix columns are always concatenated and treated as a single variable for purposes of the reference grid.
- `linfct` matrix. Each row consists of the linear function of the regression coefficients for predicting its corresponding 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` numeric. The regression coefficients. If there is a multivariate response, the matrix of coefficients is flattened to a single vector, and `linfct` and `V` redefined appropriately. Important: `bhat` must include any NA values produced as a result of collinearity in the predictors. These are taken care of later in the estimability check.
- `nbasis` matrix. The basis for the non-estimable functions of the regression coefficients. Every EMM 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. If everything is estimable, `nbasis` should be a 1 x 1 matrix of NA.
- `V` matrix. The symmetric variance-covariance matrix of `bhat`.
- `dffun` function having two arguments. `dffun(k, dfargs)` should return the degrees of freedom for the linear function `sum(k*bhat)`, or NA if unavailable.
- `dfargs` list. Used to hold any additional information needed by `dffun`.
- `misc` list. 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.emmGrid` method. Elements in this slot may be modified if desired using the `update.emmGrid` method.
- `post.beta` matrix. A sample from the posterior distribution of the regression coefficients, if MCMC methods were used; or a 1 x 1 matrix of NA otherwise. When it is non-trivial, the `as.mcmc.emmGrid` method returns `post.beta %*% t(linfct)`, which is a sample from the posterior distribution of the EMMs.
Methods

All methods for these objects are S3 methods except for show. They include [.emmGrid, as.glht.emmGrid, as.mcmc.emmGrid, as.mcmc.list.emmGrid (see coda), cld.emmGrid (see multcomp), coef.emmGrid, confint.emmGrid, contrast.emmGrid, pairs.emmGrid, plot.emmGrid, predict.emmGrid, print.emmGrid, rbind.emmGrid, show.emmGrid, str.emmGrid, summary.emmGrid, test.emmGrid, update.emmGrid, vcov.emmGrid, and xtable.emmGrid

emmpip

Interaction-style plots for estimated marginal means

Description

Creates an interaction plot of EMMs based on a fitted model and a simple formula specification.

Usage

emmpip(object, formula, ...)  
## Default S3 method:
emmpip(object, formula, type, CIs = FALSE, PIs = FALSE, 
    style, engine = get_emm_option("graphics.engine"), plotit = TRUE, 
    nesting.order = FALSE, ...)

e MMPip_g gplot(e mms, style = "factor", dodge = 0.1, xlab = labs$xlab, 
    ylab = labs$ylab, tlab = labs$tlab, facetlab = "label_context", scale, 
    dotarg = list(), linearg = list(), CIarg = list(lwd = 2, alpha = 0.5), 
    PIarg = list(lwd = 1.25, alpha = 0.33), ...)  

emmpip_lattice(emms, style = "factor", xlab = labs$xlab, ylab = labs$ylab, 
    tlab = labs$tlab, pch = c(1, 2, 6, 7, 9, 10, 15:20), lty = 1, 
    col = NULL, ...)

Arguments

object An object of class emmGrid, or a fitted model of a class supported by the emmeans package

formula Formula of the form trace.factors ~ x.factors | by.factors. The EMMs are plotted against x.factor 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.

... Additional arguments passed to emmeans (when object is not already an emmGrid object), predict.emmGrid, emmpip_ggplot, or emmpip_lattice.
As in `predict.emmGrid`, 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 `emm_options`. In addition, the user may specify type = "scale" to create a transformed scale for the vertical axis based on object’s response transformation or link function.

CIs Logical value. If TRUE, confidence intervals (or HPD intervals for Bayesian models) are added to the plot (works only with engine = "ggplot").

PIs Logical value. If TRUE, prediction intervals are added to the plot (works only with engine = "ggplot"). If both CIs and PIs are TRUE, the prediction intervals will be somewhat longer, lighter, and thinner than the confidence intervals. Additional parameters to `predict.emmGrid` (e.g., sigma) may be passed via .... For Bayesian models, PIs require frequentist = TRUE and a value for sigma.

Optional character value. This has an effect only when the horizontal variable is a single numeric variable. If style is unspecified or "numeric", the horizontal scale will be numeric and curves are plotted using lines (and no symbols). With style = "factor", the horizontal variable is treated as the levels of a factor (equally spaced along the horizontal scale), and curves are plotted using lines and symbols. When the horizontal variable is character or factor, or a combination of more than one predictor, "factor" style is always used.

Character value matching "ggplot" (default), "lattice", or "none". The graphics engine to be used to produce the plot. These require, respectively, the `ggplot2` or `lattice` package to be installed. Specifying "none" is equivalent to setting plotit = FALSE.

Logical value. If TRUE, a graphical object is returned; if FALSE, a data.frame is returned containing all the values used to construct the plot.

Logical value. If TRUE, factors that are nested are presented in order according to their nesting factors, even if those nesting factors are not present in formula. If FALSE, only the variables in formula are used to order the variables.

A data.frame created by calling emmip with plotit = FALSE. Certain variables and attributes are expected to exist in this data frame; see the section detailing the rendering functions.

Numerical amount passed to `ggplot2::position_dodge` by which points and intervals are offset so they do not collide.

Character labels for the horizontal axis, vertical axis, and traces (the different curves), respectively. The emmip function generates these automatically and provides them via the labs attribute, but the user may override these if desired.

Labeller for facets (when by variables are in play). Use "label_value" to show just the factor levels, or "label_both" to show both the factor names and factor levels. The default of "label_context" decides which based on how many by factors there are. See the documentation for `ggplot2::label_context`.

If not missing, an object of class `scales::trans` specifying a (usually) non-linear scaling for the vertical axis. For example, `scales = scales::log_trans()` specifies a logarithmic scale. For fine-tuning purposes, additional arguments to `ggplot2::scale_y_continuous` may be included in ....
dotarg  list of arguments passed to `geom_point` to customize appearance of points
linearg list of arguments passed to `geom_line` to customize appearance of lines
CIarg, PIarg lists of arguments passed to `geom_linerange` to customize appearance of intervals
pch The plotting characters to use for each group (i.e., levels of `trace.factors`). They are recycled as needed.
lty The line types to use for each group. Recycled as needed.
col The colors to use for each group, recycled as needed. If not specified, the default trellis colors are used.

Value

If `plotit = FALSE`, a `data.frame` (actually, a `summary_emm` object) with the table of EMMs that would be plotted. The variables plotted are named `xvar` and `yvar`, and the trace factor is named `tvar`. This data frame has an added "labs" attribute containing the labels `xlab`, `ylab`, and `tlab` for these respective variables. The confidence limits are also included, renamed `LCL` and `UCL`.

If `plotit = TRUE`, the function returns an object of class "`ggplot" or a "trellis", depending on engine.

Details

If `object` is a fitted model, `emmeans` is called with an appropriate specification to obtain estimated marginal 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 `emmeans`). Otherwise, if `object` is an `emmGrid` object, its first element is used, and it must contain one estimate for each combination of the factors present in `formula`.

Rendering functions

The functions `emnip_ggplot` and `emnip_lattice` are called when `plotit == TRUE` to render the plots; but they may also be called later on an object saved via `plotit = FALSE` (or `engine = "none"`). The functions require that `emms` contains variables `xvar`, `yvar`, and `tvar`, and attributes "labs" and "vars". Confidence intervals are plotted if variables `LCL` and `UCL` exist; and prediction intervals are plotted if `LPL` and `UPL` exist. Finally, it must contain the variables named in `attr(emms, "vars")`.

Note

Conceptually, this function is equivalent to `interaction.plot` where the summarization function is thought to return the EMMs.

See Also

`emmeans`, `interaction.plot`.
Examples

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

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

# One interaction plot, using combinations of size and side as the x factor
# ... with added confidence intervals and some formatting changes
emmip(noise.lm, type ~ side * size, CIs = TRUE,
linarg = list(linetype = "dashed"), CIarg = list(lwd = 1, alpha = 1))

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

# Individual traces in panels
emmip(noise.lm, ~ size | type * side)

# Example for the 'style' argument
fib.lm = lm(strength ~ machine * sqrt(diameter), data = fiber)
fib.rg = ref_grid(fib.lm, at = list(diameter = c(3.5, 4, 4.5, 5, 5.5, 6)^2))
emmip(fib.rg, machine ~ diameter)  # curves (because diameter is numeric)
emmip(fib.rg, machine ~ diameter, style = "factor")  # points and lines

# For an example using extra ggplot2 code, see 'vignette("messy-data")',
# in the section on nested models.

### Options with transformations or link functions
neuralgia.glm <- glm(Pain ~ Treatment * Sex + Age, family = binomial(),
data = neuralgia)

# On link scale:
emmip(neuralgia.glm, Treatment ~ Sex)

# On response scale:
emmip(neuralgia.glm, Treatment ~ Sex, type = "response")

# With transformed axis scale and custom scale divisions
emmip(neuralgia.glm, Treatment ~ Sex, type = "scale",
breaks = seq(0.10, 0.90, by = 0.10))

emobj

--- Construct an emmGrid object from scratch

Description

This allows the user to incorporate results obtained by some analysis into an emmGrid object, enabling the use of emmGrid methods to perform related follow-up analyses.
Usage

```
emmobj(bhat, V, levels, linfct = diag(length(bhat)), df = NA, dffun,
dfargs = list(), post.beta = matrix(NA), nesting = NULL, ...)
```

Arguments

- **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 value 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 \(\sum(x\times bhat)\), with any additional parameters in dfargs.
- **dffun**: Overrides df if specified. This is a convenience to match the slot names of the returned object.
- **dfargs**: List containing arguments for df. This is ignored if df is numeric.
- **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.
- **nesting**: Nesting specification as in ref_grid. This is ignored if model.info is supplied.
- **...**: Arguments passed to update.emmGrid

Details

The arguments must be conformable. This includes that the length of bhat, the number of columns of linfct, and the number of columns of post.beta must all be equal. And that the product of lengths in levels must be equal to the number of rows of linfct. The grid slot of the returned object is generated by expand.grid using levels as its arguments. So the rows of linfct should be in corresponding order.

The functions qdrg and emmobj are close cousins, in that they both produce emmGrid objects. When starting with summary statistics for an existing grid, emmobj is more useful, while qdrg is more useful when starting from an unsupported fitted model.

Value

An emmGrid object

See Also

qdrg, an alternative that is useful when starting with a fitted model not supported in emmeans.
Examples

# Given summary statistics for 4 cells in a 2 x 2 layout, obtain
# marginal means and comparisons thereof. Assume heteroscedasticity
# and use the Satterthwaite method
levels <- list(trt = c("A", "B"), dose = c("high", "low"))
ybar <- c(57.6, 43.2, 88.9, 69.8)
s <- c(12.1, 19.5, 22.8, 43.2)
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))
expt.rg <- emmobj(bhat = ybar, V = diag(se2),
  levels = levels, linfct = diag(c(1, 1, 1, 1)),
  df = Satt.df, dfargs = list(v = se2, n = n), estName = "mean")
plot(expt.rg)
(trt.emm <- emmeans(expt.rg, "trt") )
(dose.emm <- emmeans(expt.rg, "dose") )
rbind(pairs(trt.emm), pairs(dose.emm), adjust = "mvt")

emmm_list

The emm_list class

Description

An emm_list object is simply a list of emmGrid objects. Such a list is returned, for example, by
emmeans with a two-sided formula or a list as its specs argument. Several methods for this class
are provided, as detailed below.

Usage

## S3 method for class 'emm_list'
contrast(object, ..., which = I_bet(1))

## S3 method for class 'emm_list'
pairs(x, ..., which = I_bet(1))

## S3 method for class 'emm_list'
test(object, ..., which = seq_along(object))

## S3 method for class 'emm_list'
confint(object, ..., which = seq_along(object))

## S3 method for class 'emm_list'
plot(x, ..., which = I_bet(1))
## S3 method for class `emm_list`
coef(object, ..., which = I_bet(1))

## S3 method for class `emm_list`
str(object, ...)

## S3 method for class `emm_list`
summary(object, ..., which = seq_along(object))

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

I_bet(which)

### Arguments

- `object, x` - an object of class `emm_list`
- `...` - additional arguments passed to corresponding `emmGrid` method
- `which` - integer vector specifying which elements to select.

### Value

A list of objects returned by the corresponding `emmGrid` method (thus, often, another `emm_list` object). However, if `which` has length 1, the one result is not wrapped in a list.

### Note

- The `plot` method uses only the first element of `which`; the others are ignored.
- No export option is provided for printing an `emm_list` (see `print.emmGrid`). If you wish to export these objects, you must do so separately for each element in the list.
- `I_bet()` provides a default value for `which` noisily. It is used in cases where users most likely intended to call a method for an `emmGrid` object rather than an `emm_list` object. An informative message is displayed and then the `which`th element is returned.

### Description

Use `emm_options` to set or change various options that are used in the `emmeans` package. These options are set separately for different contexts in which `emmGrid` objects are created, in a named list of option lists.
**Usage**

```r
emm_options(..., disable)

get_emm_option(x, default = emm_defaults[[x]])
```

**Arguments**

- `...` Option names and values (see Details)
- `disable` If non-missing, this will reset all options to their defaults if `disable` tests `TRUE` (but first save them for possible later restoration). Otherwise, all previously saved options are restored. This is important for bug reporting; please see the section below on reproducible bugs. When `disable` is specified, the other arguments are ignored.
- `x` Character value - the name of an option to be queried
- `default` Value to return if `x` is not found

**Format**

An object of class `list` of length 21.

**Details**

`emmeans`'s options are stored as a list in the system option "emmeans". Thus, `emm_options(foo = bar)` is the same as `options(emmeans = list(..., foo = bar))` where `...` represents any previously existing options. The list `emm_defaults` contains the default values in case the corresponding element of system option `emmeans` is `NULL`.

Currently, the following main list entries are supported:

- `ref_grid` A named list of defaults for objects created by `ref_grid`. This could affect other objects as well. For example, if `emmeans` is called with a fitted model object, it calls `ref_grid` and this option will affect the resulting `emmGrid` object.
- `emmeans` A named list of defaults for objects created by `emmeans` or `emtrends`.
- `contrast` A named list of defaults for objects created by `contrast.emmGrid` or `pairs.emmGrid`.
- `summary` A named list of defaults used by the methods `summary.emmGrid`, `predict.emmGrid`, `test.emmGrid`, `confint.emmGrid`, and `emmip`. The only option that can affect the latter four is "predict.method".
- `sep` A character value to use as a separator in labeling factor combinations. Such labels are potentially used in several places such as `contrast` and `plot.emmGrid` when combinations of factors are compared or plotted. The default is " ".
- `parens` Character vector that determines which labels are parenthesized when they are contrasted. The first element is a regular expression, and the second and third elements are used as left and right parentheses. See details for the `parens` argument in `contrast`. The default will parenthesize labels containing the four arithmetic operators, using round parentheses.
cov.keep The default value of cov.keep in ref.grid. Defaults to "2", i.e., two-level covariates are treated like factors.

graphics.engine A character value matching c("ggplot", "lattice"), setting the default engine to use in emmip and plot.emmGrid. Defaults to "ggplot".

msg.interaction A logical value controlling whether or not a message is displayed when emmeans averages over a factor involved in an interaction. It is probably not appropriate to do this, unless the interaction is weak. Defaults to TRUE.

msg.nesting A logical value controlling whether or not to display a message when a nesting structure is auto-detected. The existence of such a structure affects computations of EMMs. Sometimes, a nesting structure is falsely detected – namely when a user has omitted some main effects but included them in interactions. This does not change the model fit, but it produces a different parameterization that is picked up when the reference grid is constructed. Defaults to TRUE.

rg.limit An integer value setting a limit on the number of rows in a newly constructed reference grid. This is checked based on the number of levels of the factors involved; but it excludes the levels of any multivariate responses because those are not yet known. The reference grid consists of all possible combinations of the predictors, and this can become huge if there are several factors. An error is thrown if this limit is exceeded. One can use the nuisance argument of ref.grid to collapse on nuisance factors, thus making the grid smaller. Defaults to 10,000.

simplify.names A logical value controlling whether to simplify (when possible) names in the model formula that refer to datasets – for example, should we simplify a predictor name like "data$trt" to just "trt"? Defaults to TRUE.

opt.digits A logical value controlling the precision with which summaries are printed. If TRUE (default), the number of digits displayed is just enough to reasonably distinguish estimates from the ends of their confidence intervals; but always at least 3 digits. If FALSE, the system value getOption("digits") is used.

back.bias.adj A logical value controlling whether we try to adjust bias when back-transforming. If FALSE, we use naive back transformation. If TRUE and sigma is available, a second-order adjustment is applied to estimate the mean on the response scale.

enable.submodel A logical value. If TRUE, enables support for selected model classes to implement the submodel option. If FALSE, this support is disabled. Setting this option to FALSE could save excess memory consumption.

Some other options have more specific purposes:

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

save.ref_grid Logical value of TRUE if you wish the latest reference grid created to be saved in .Last.ref_grid. The default is FALSE.

Options for lme4::lmerMod models Options lmer.df, disable.pbkrtest, pbkrtest.limit, disable.lmerTest, and lmerTest.limit options affect how degrees of freedom are computed for lmerMod objects produced by the lme4 package. See that section of the "models" vignette for details.
Value

`emm_options` returns the current options (same as the result of `getOption("emmeans")`) invisibly, unless called with no arguments.

`get_emm_option` returns the currently stored option for `x`, or its default value if not found.

Reproducible bugs

Most options set display attributes and such that are not likely to be associated with bugs in the code. However, some other options (e.g., `cov.keep`) are essentially configuration settings that may affect how/whether the code runs, and the settings for these options may cause subtle effects that may be hard to reproduce. Therefore, when sending a bug report, please create a reproducible example and make sure the bug occurs with all options set at their defaults. This is done by preceding it with `emm_options(disable = TRUE)`.

By the way, `disable` works like a stack (LIFO buffer), in that `disable = TRUE` is equivalent to `emm_options(saved.opts = emm_options())` and `emm_options(disable = FALSE)` is equivalent to `options(emmeans = get_emm_option("saved.opts"))`. To completely erase all options, use `options(emmeans = NULL)`.

See Also

`update.emmGrid`

Examples

```r
## Not run:
emm_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 emmeans 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.

## End(Not run)

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

## End(Not run)

# See tolerance being used for determining estimability
get_emm_option("estble.tol")

## Not run:
# Set all options to their defaults
emm_options(disable = TRUE)
# ... and perhaps follow with code for a minimal reproducible bug,
# which may include emm_options() clls if they are pertinent ...
emtrends

## Description

The emtrends function is useful when a fitted model involves a numerical predictor \( x \) interacting with another predictor \( a \) (typically a factor). Such models specify that \( x \) has a different trend depending on \( a \); thus, it may be of interest to estimate and compare those trends. Analogous to the emmeans setting, we construct a reference grid of these predicted trends, and then possibly average them over some of the predictors in the grid.

## Usage

emtrends(object, specs, var, delta.var = 0.001 * rng, max.degree = 1, ...)

## Arguments

- **object**: A supported model object (not a reference grid)
- **specs**: Specifications for what marginal trends are desired – as in emmeans. If specs is missing or NULL, emmeans is not run and the reference grid for specified trends is returned.
- **var**: Character value 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 EMMs, we compute and compare the slopes of the var trend over levels of the specified other predictor(s). As in EMMs, marginal averages are computed for the predictors in specs and by. See also the “Generalizations” section below.
- **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 1/1000 of the range of var over the dataset.
- **max.degree**: Integer value. The maximum degree of trends to compute (this is capped at 5). If greater than 1, an additional factor degree is added to the grid, with corresponding numerical derivatives of orders 1, 2, \ldots, max.degree as the estimates.
- **...**: Additional arguments passed to ref_grid or emmeans as appropriate. See Details.
Details

The function works by constructing reference grids for object with various values of var, and then calculating difference quotients of predictions from those reference grids. Finally, emmeans is called with the given specs, thus computing marginal averages as needed of the difference quotients. Any ... arguments are passed to the ref_grid and emmeans; examples of such optional arguments include optional arguments (often mode) that apply to specific models; ref_grid options such as data, at, cov.reduce, mult.names, nesting, or transform; and emmeans options such as weights (but please avoid trend or offset).

Value

An emmGrid or emm_list object, according to specs. See emmeans for more details on when a list is returned.

Generalizations

Instead of a single predictor, 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 object 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, making the predicted trends vary depending on dose.

Note

In earlier versions of emtrends, the first argument was named model rather than object. (The name was changed because of potential mis-matching with a mode argument, which is an option for several types of models.) For backward compatibility, model still works provided all arguments are named.

It is important to understand that trends computed by emtrends are not equivalent to polynomial contrasts in a parallel model where var is regarded as a factor. That is because the model object here is assumed to fit a smooth function of var, and the estimated trends reflect local behavior at particular value(s) of var; whereas when var is modeled as a factor and polynomial contrasts are computed, those contrasts represent the global pattern of changes over all levels of var.

See the pigs.poly and pigs.fact examples below for an illustration. The linear and quadratic trends depend on the value of percent, but the cubic trend is constant (because that is true of a cubic polynomial, which is the underlying model). The cubic contrast in the factorial model has the same P value as for the cubic trend, again because the cubic trend is the same everywhere.

See Also

emmeans, ref_grid

Examples

fiber.lm <- lm(strength ~ diameter*machine, data=fiber)
# Obtain slopes for each machine ...
( fiber.emt <- emtrends(fiber.lm, "machine", var = "diameter") )
# ... and pairwise comparisons thereof
pairs(fiber.emt)
# Suppose we want trends relative to sqrt(diameter)...
emtrends(fiber.lm, ~ machine | diameter, var = "sqrt(diameter)",
at = list(diameter = c(20, 30)))

# Obtaining a reference grid
mtcars.lm <- lm(mpg ~ poly(disp, degree = 2) * (factor(cyl) + factor(am)), data = mtcars)

# Center trends at mean disp for each no. of cylinders
mtcTrends.rg <- emtrends(mtcars.lm, var = "disp",
cov.reduce = disp ~ factor(cyl))

summary(mtcTrends.rg) # estimated trends at grid nodes
eemmeans(mtcTrends.rg, "am", weights = "prop")

### Higher-degree trends ...
pigs.poly <- lm(conc ~ poly(percent, degree = 3), data = pigs)
emt <- emtrends(pigs.poly, ~ degree | percent, "percent", max.degree = 3,
at = list(percent = c(9, 13.5, 18))
    # note: 'degree' is an extra factor created by 'emtrends'

summary(emt, infer = c(TRUE, TRUE))

# Compare above results with poly contrasts when 'percent' is modeled as a factor ...
pigs.fact <- lm(conc ~ factor(percent), data = pigs)
emm <- emmeans(pigs.fact, "percent")

contrast(emm, "poly")
# Some P values are comparable, some aren’t! See Note in documentation

---

**extending-emmeans**

Support functions for model extensions

**Description**

This documents the methods that `ref_grid` calls. A user or package developer may add `emmeans` support for a model class by writing `recover_data` and `emm_basis` methods for that class. (Users in need for a quick way to obtain results for a model that is not supported may be better served by the `qdrg` function.)

**Usage**

`recover_data(object, ...)`

## S3 method for class 'call'
`recover_data(object, trms, na.action, data = NULL, params = "pi", frame, ...)`

`emm_basis(object, trms, xlev, grid, ...)`
.recover_data(object, ...)

.emm_basis(object, trms, xlev, grid, ...)

.emm_register(classes, pkgname)

Arguments

object
An object of the same class as is supported by a new method.

... Additional parameters that may be supported by the method.

trms The terms component of object (typically with the response deleted, e.g. via delete.response)

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. For example, a spline model may involve a local variable knots that is not a predictor, but its value is needed to fit the model. Names of parameters not actually used are harmless, and the default value "pi" (the only numeric constant in base R) is provided in case the model involves it. An example involving splines may be found at https://github.com/rvlenth/emmeans/issues/180.

frame Optional data.frame. Many model objects contain the model frame used when fitting the model. In cases where there are no predictor transformations, this model frame has all the original predictor values and so is usable for recovering the data. Thus, if frame is non-missing and data is NULL, a check is made on trms and if there are no function calls, we use data = frame. This can be helpful because it provides a modicum of security against the possibility that the original data used when fitting the model has been altered or removed.

xlev Named list of factor levels (excluding ones coerced to factors in the model formula)

grid A data.frame (provided by ref_grid) containing the predictor settings needed in the reference grid

classes Character names of one or more classes to be registered. The package must contain the functions recover_data.foo and emm_basis.foo for each class foo listed in classes.

pkgname Character name of package providing the methods (usually should be the second argument of .onLoad)

Value

The recover_data method must return a data.frame containing all the variables that appear as predictors in the model, and attributes "call", "terms", "predictors", and "responses". (recover_data.call will provide these attributes.)

The emm_basis method should return a list with the following elements:
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 \( \sum(k \times bhat) \).

dfargs  A list containing additional arguments needed for dffun.

.recover_data and .emm_basis are hidden exported versions of recover_data and emm_basis, respectively. They run in emmeans’s namespace, thus providing access to all existing 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 emm_basis respectively. A vignette giving details and examples is available via vignette("xtending", "emmeans")

To extend emmeans’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 emm_basis method is more involved, but the existing methods (e.g., emmeans:::emm_basis.lm) can serve as models. Certain recover_data and emm_basis methods are exported from emmeans. (To find out, do methods("recover_data").) 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.emmGrid and its relatives check the estimability of each prediction, using the nonest.basis function in the estimability package.

The models already supported are detailed in the "models" vignette. Some packages may provide additional emmeans support for its object classes.

Communication between methods

If the recover_data method generates information needed by emm_basis, that information may be incorporated by creating a “misc” attribute in the returned recovered data. That information is then passed as the misc argument when ref_grid calls emm_basis.
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 \texttt{misc$estHook}, \texttt{misc$vcovHook} and \texttt{misc$postGridHook}. If just the name of the hook function is provided as a character string, then it is retrieved using \texttt{get}.

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

Registering S3 methods for a model class

The \texttt{.emm_register} function is provided as a convenience to conditionally register your S3 methods for a model class, \texttt{recover_data.foo} and \texttt{emm_basis.foo}, where \texttt{foo} is the class name. Your package should implement an \texttt{.onLoad} function and call \texttt{.emm_register} if \texttt{emmeans} is installed. See the example.

Note

Without an explicit data argument, \texttt{recover_data} returns the \texttt{current version} of the dataset. If the dataset has changed since the model was fitted, then this will not be the data used to fit the model. It is especially important to know this in simulation studies where the data are randomly generated or permuted, and in cases where several datasets are processed in one step (e.g., using \texttt{dplyr}). In those cases, users should be careful to provide the actual data used to fit the model in the data argument.

See Also

Vignette on extending emmeans

Examples

```r
# Not run:
#--- If your package provides recover_data and emm_grid methods for class 'mymod',
#--- put something like this in your package code -- say in zzz.R:
.onLoad = function(libname, pkname) {
  if (requireNamespace("emmeans", quietly = TRUE))
    emmeans::.emm_register("mymod", pkname)
}

# End(Not run)
```
**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**

```r
feedlot
```

**Format**

A data frame with 67 observations and 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**

```r
feedlot.lm <- lm(swt ~ ewt + herd*diet, data = feedlot)
# Obtain EMMs with a separate reference value of ewt for each herd. This reproduces the last part of Table 2 in the reference
emmeans(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 and 3 variables:

- **machine**: a factor with levels A B C. This is 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
fiber.lm <- lm(strength ~ diameter + machine, data=fiber)
ref_grid(fiber.lm)
# Covariate-adjusted means and comparisons
ebbeans(fiber.lm, pairwise ~ machine)
```
`hpd.summary`  
*Summarize an emmGrid from a Bayesian model*

**Description**

This function computes point estimates and HPD intervals for each factor combination in `object@emmGrid`. While this function may be called independently, it is called automatically by the S3 method `summary.emmGrid` when the object is based on a Bayesian model. (Note: the `level` argument, or its default, is passed as `prob`).

**Usage**

```r
hpd.summary(object, prob, by, type, point.est = median, bias.adjust = get_emm_option("back.bias.adj"), sigma, ...)
```

**Arguments**

- `object` an `emmGrid` object having a non-missing `post.beta` slot
- `prob` numeric probability content for HPD intervals (note: when not specified, the current `level` option is used; see `emm_options`)
- `by` factors to use as by variables
- `type` prediction type as in `summary.emmGrid`
- `point.est` function to use to compute the point estimates from the posterior sample for each grid point
- `bias.adjust` Logical value for whether to adjust for bias in back-transforming (type = "response"). This requires a value of `sigma` to exist in the object or be specified.
- `sigma` Error SD assumed for bias correction (when type = "response"). If not specified, `object@misc$sigma` is used, and an error is thrown if it is not found. *Note:* `sigma` may be a vector, as long as it conforms to the number of observations in the posterior sample.
- `...` required but not used

**Value**

An object of class `summary_emm`

**See Also**

`summary.emmGrid`
Examples

```r
if(require("coda")) {
  # Create an emmGrid object from a system file
  cbpp.rg <- do.call(emmobj,
    readRDS(system.file("extdata", "cbpplist", package = "emmeans")))
  hpd.summary(emmeans(cbpp.rg, "period"))
}
```

joint_tests

Compute joint tests of the terms in a model

Description

This function produces an analysis-of-variance-like table based on linear functions of predictors in a model or emmGrid object. Specifically, the function constructs, for each combination of factors (or covariates reduced to two or more levels), a set of (interaction) contrasts via contrast, and then tests them using test with joint = TRUE. Optionally, one or more of the predictors may be used as by variable(s), so that separate tables of tests are produced for each combination of them.

Usage

```r
joint_tests(object, by = NULL, show0df = FALSE, showconf = TRUE,
  cov.reduce = meanint, ...)
```

```r
meanint(x)
```

```r
symmint(ctr)
```

Arguments

- **object**: a fitted model or an emmGrid
- **by**: character names of by variables. Separate sets of tests are run for each combination of these.
- **show0df**: logical value; if TRUE, results with zero numerator degrees of freedom are displayed, if FALSE they are skipped
- **showconf**: logical value. When we have models with estimability issues (e.g., missing cells), then with showconf = TRUE, we test any remaining effects that are not purely due to contrasts of a single term. If found, they are labeled (confounded). See vignette("xplanations") for more information.
- **cov.reduce**: a function. If object is a fitted model, it is replaced by ref_grid(object, cov.reduce = cov.reduce, ...). For this purpose, the functions meanint and symmint are available for returning an interval around the mean or around zero, respectively. See the section below on covariates.
- **...**: additional arguments passed to ref_grid and emmeans
- **x, ctr**: arguments for meanint and symmint
Details

In models with only factors, no covariates, these tests correspond to “type III” tests a la SAS, as long as equal-weighted averaging is used and there are no estimability issues. When covariates are present and they interact with factors, the results depend on how the covariate is handled in constructing the reference grid. See the section on covariates below. The point that one must always remember is that joint_tests always tests contrasts among EMMs, in the context of the reference grid, whereas type III tests are tests of model coefficients – which may or may not have anything to do with EMMs or contrasts.

Value

a summary_emm object (same as is produced by summary.emmGrid). All effects for which there are no estimable contrasts are omitted from the results. There may be an additional row named (confounded) which accounts for additional degrees of freedom for effects not accounted for in the preceding rows.

The returned object also includes an "est.fcns" attribute, which is a named list containing the linear functions associated with each joint test. No estimable functions are included for confounded effects.

meanint returns mean(c) + c(-1, 1). symmint(ctr) returns a function that returns ctr + c(-1, 1). These functions are available primarily for use with cov.reduce.

Dealing with covariates

A covariate (or any other predictor) must have more than one value in the reference grid in order to test its effect and be included in the results. Therefore, when object is a model, we default to cov.reduce = meanint which sets each covariate at a symmetric interval about its mean. But when object is an existing reference grid, it often has only one value for covariates, in which case they are excluded from the joint tests.

Covariates present further complications in that their values in the reference grid can affect the joint tests of other effects. When covariates are centered around their means (the default), then the tests we obtain can be described as joint tests of covariate-adjusted means; and that is our intended use here. However, some software such as SAS and car::Anova adopt the convention of centering covariates around zero; and for that purpose, one can use cov.reduce = symmint(0) when calling with a model object (or in constructing a reference grid). However, adjusted means with covariates set at or around zero do not make much sense in the context of interpreting estimated marginal means, unless the covariate means really are zero.

See the examples below with the toy dataset.

See Also
test

Examples

pigs.lm <- lm(log(conc) ~ source * factor(percent), data = pigs)

(jt <- joint_tests(pigs.lm))  # will be same as type III ANOVA
### Estimable functions associated with "percent"

```
attr(jt, "est.fcns") $ "percent"
```

```
joint_tests(pigs.lm, weights = "outer")  ## differently weighted

joint_tests(pigs.lm, by = "source")    ## separate joint tests of 'percent'
```

### Comparisons with type III tests in SAS

```
toy = data.frame(
  treat = rep(c("A", "B"), c(4, 6)),
  female = c(1, 0, 0, 1, 0, 1, 1, 0, 0, 0, 1, 0),
  resp = c(17, 12, 14, 19, 28, 26, 26, 34, 33, 27))

toy.fac = lm(resp ~ treat * factor(female), data = toy)
toy.cov = lm(resp ~ treat * female, data = toy)
# (These two models have identical fitted values and residuals)
```

```
# -- SAS output we'd get with toy.fac --
   Source DF Type III SS Mean Square   F Value Pr > F
#   treat  1  488.8928571  488.8928571  404.60 <.0001
#  female  1   78.8928571   78.8928571   65.29 0.0002
#  treat*female  1   1.7500000   1.7500000   1.45 0.2741

# -- SAS output we'd get with toy.cov --
   Source DF Type III SS Mean Square   F Value Pr > F
#   treat  1  252.0833333  252.0833333  208.62 <.0001
#  female  1   78.8928571   78.8928571   65.29 0.0002
# female*treat  1   1.7500000   1.7500000   1.45 0.2741
```

```
joint_tests(toy.fac)
joint_tests(toy.cov)  # female is regarded as a 2-level factor by default
```

```
# Treat 'female' as a numeric covariate (via cov.keep = 0)
# ... then tests depend on where we center things

# Center around the mean
joint_tests(toy.cov, cov.keep = 0, cov.reduce = meanint)

# Center around zero (like SAS's results for toy.cov)
joint_tests(toy.cov, cov.keep = 0, cov.reduce = symmint(0))

# Center around 0.5 (like SAS's results for toy.fac)
joint_tests(toy.cov, cov.keep = 0, cov.reduce = range)
```

### Example with empty cells and confounded effects

```
low3 <- unlist(attr(ubds, "cells")[1:3])
ubds.lm <- lm(y ~ A*B*C, data = ubds, subset = -low3)

# Show overall joint tests by C:
ref_grid(ubds.lm, by = "C") |> contrast("consec") |> test(joint = TRUE)
```

```
# Break each of the above into smaller components:
joint_tests(ubds.lm, by = "C")
```
lsmeans  Wrappers for alternative naming of EMMs

Description
These are wrappers for `emmeans` and related functions to provide backward compatibility, or for users who may prefer to use other terminology than “estimated marginal means” – namely “least-squares means”. These functions also provide the functionality formerly provided by the `lsmeans` package, which is now just a front-end for `emmeans`.

Usage

```r
lsmeans(...)  
listrends(...)  
lsmip(...)  
lsm(...)  
lsmobj(...)  
lsm.options(...)  
get.lsm.option(x, default = emm_defaults[[x]])
```

Arguments

- `...`: Arguments passed to the corresponding `emxxxx` function
- `x`: Character name of desired option
- `default`: default value to return if `x` not found

Details
For each function with `lsxxxx` in its name, the same function named `emxxxx` is called. Any estimator names or list items beginning with “em” are replaced with “ls” before the results are returned.

Value
The result of the call to `emxxxx`, suitably modified.

get.lsm.option and lsm.options remap options from and to corresponding options in the `emmeans` options system.

See Also

`emmeans`, `emtrends`, `emmip`, `emm`, `emobj`, `emm_options`, `get_emm_option`
make.tran

Examples

```r
pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
lsmeans(pigs.lm, "source")
``` 

Description

The `make.tran` function creates 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.emmGrid` (when the given transformation was already applied in an existing model).

Usage

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

Arguments

- **type**: The name of the transformation. See Details.
- **param**: Numeric parameter needed 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.
- **y, ...**: Used only with type = "scale". These parameters are passed to `scale` to determine param.

Details

The functions `emmeans`, `ref_grid`, and related ones automatically detect response transformations that are recognized by examining the model formula. These are `log`, `log2`, `log10`, `log1p`, `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.mult argument in `update.emmGrid`).

A few additional character strings may be supplied as the tran argument in `update.emmGrid`: "identity", "1/mu^2", "inverse", "reciprocal", "log10", "log2", "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.emmGrid`. 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`; in the following discussion, we use `p` to denote this parameter, and `y` to denote the response variable. 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: \( abs(y)^p \times sign(y) \). There are no restrictions on \( y \), but we require \( p > 0 \) in order for the transformation to be monotone and continuous.
- **"asin.sqrt"** Arcsin-square-root transformation: \( \sin^{/(y/p)^{1/2}} \). Typically, the parameter \( p \) is equal to 1 for a fraction, or 100 for a percentage.
- **"bcnPower"** Box-Cox with negatives allowed, as described for the bcnPower function in the `car` package. It is defined as the Box-Cox transformation \( (z^p - 1)/p \) of the variable \( z = y + (y^2 + g^2)^{1/2} \). This requires `param` to have two elements: the power \( p \) and the offset \( g > 0 \).
- **"scale"** This one is a little different than the others, in that `param` is ignored; instead, `param` is determined by calling `scale(y, ...)`. The user should give as `y` the response variable in the model to be fitted to its scaled version.

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. In the "bcnPower" transformation, the second element is required and must be positive.

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 those returned by `make.link`. The `linkfun` component is the transformation itself.

**Note**

The `genlog` transformation is technically unneeded, because a response transformation of the form \( \log(y + c) \) is now auto-detected by `ref_grid`.

We modify certain `make.link` results in transformations where there is a restriction on valid prediction values, so that reasonable inverse predictions are obtained, 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.
Examples

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

### Using a scaled response...
# Case where it is auto-detected:
fib.lm <- lm(scale(strength) ~ diameter + machine, data = fiber)
ref_grid(fib.lm)

# Case where scaling is not auto-detected -- and what to do about it:
fib.aov <- aov(scale(strength) ~ diameter + Error(machine), data = fiber)
fib.rg <- suppressWarnings(ref_grid(fib.aov, at = list(diameter = c(20, 30))))

# Scaling was not retrieved, so we can do:
fib.rg = update(fib.rg, tran = make.tran("scale", y = fiber$strength))
emmeans(fib.rg, "diameter")

## Not run:
### An existing model 'mod' was fitted with a y^(2/3) transformation...
ptran = make.tran("power", 2/3)
emmeans(mod, "treatment", tran = ptran)

## 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

MOats

Format

A data frame with 18 observations and 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

```r
MOats.lm <- lm (yield ~ Block + Variety, data = MOats)
MOats.rg <- ref_grid (MOats.lm, mult.name = "nitro")
emmeans(MOats.rg, ~ nitro | Variety)
```

Description

Documentation for models has been moved to a vignette. To access it, use `vignette("models", "emmeans")`.

```
mvcontrast
```

Multivariate contrasts

Description

This function displays tests of multivariate comparisons or contrasts. The contrasts are constructed at each level of the variable in `mult.name`, and then we do a multivariate test that the vector of estimates is equal to `null` (zero by default). The $F$ statistic and degrees of freedom are determined via the Hotelling distribution. that is, if there are $m$ error degrees of freedom and multivariate dimensionality $d$, then the resulting $F$ statistic has degrees of freedom $(d, m - d + 1)$ as shown in Hotelling (1931).

Usage

```r
mvcontrast(object, method = "eff", mult.name = object@roles$multresp,
  null = 0, by = object@misp$by.vars, adjust = c("sidak",
  p.adjust.methods), show.est = FALSE, ...)
```
**mvcontrast**

**Arguments**

- **object**: An object of class `emmmGrid`
- **method**: A contrast method, per `contrast.emmGrid`
- **mult.name**: Character vector of nNames of the factors whose levels define the multivariate means to contrast. If the model itself has a multivariate response, that is what is used. Otherwise, `mult.name` must be specified.
- **null**: Scalar or conformable vector of null-hypothesis values to test against
- **by**: Any by variable(s). These should not include the primary variables to be contrasted. For convenience, the by variable is nulled-out if it would result in no primary factors being contrasted.
- **adjust**: Character value of a multiplicity adjustment method ("none" for no adjustment). The available adjustment methods are more limited that in `contrast`, and any default adjustment returned via method is ignored.
- **show.ests**: Logical flag determining whether the multivariate means are displayed
- **...**: Additional arguments passed to `contrast`

**Value**

An object of class `summary_emm` containing the multivariate test results; or a list of the estimates and the tests if `show.ests` is `TRUE`. The test results include the Hotelling $T^2$ statistic, $F$ ratios, degrees of freedom, and $P$ values.

**Note**

If some interactions among the primary and `mult.name` factors are absent, the covariance of the multivariate means is singular; this situation is accommodated, but the result has reduced degrees of freedom and a message is displayed. If there are other abnormal conditions such as non-estimable results, estimates are shown as `NA`.

While designed primarily for testing contrasts, multivariate tests of the mean vector itself can be implemented via method = "identity") (see the examples).

**References**


**Examples**

```r
MOats.lm <- lm(yield ~ Variety + Block, data = MOats)
MOats.emm <- emmeans(MOats.lm, ~ Variety | rep.meas)
mvcontrast(MOats.emm, "consec", show.ests = TRUE)  # mult.name defaults to rep.meas

# Test each mean against a specified null vector
mvcontrast(MOats.emm, "identity", name = "Variety",
          null = c(80, 100, 120, 140), adjust = "none")
# (Note 'name' is passed to contrast() and overrides default name "contrast")
```
neuralgia

# 'mult.name' need not refer to a multivariate response
mvcontrast(MOats.emm, "trt.vs.ctrl1", mult.name = "Variety")

 neuralgia                Neuralgia data

Description

These data arise from a study of analgesic effects of treatments of elderly patients who have neuralgia. Two treatments and a placebo are compared. The response variable is whether the patient reported pain or not. Researchers recorded the age and gender of 60 patients along with the duration of complaint before the treatment began.

Usage

neuralgia

Format

A data frame with 60 observations and 5 variables:

- **Treatment**: Factor with 3 levels A, B, and P. The latter is placebo.
- **Sex**: Factor with two levels F and M.
- **Age**: Numeric covariate – patient's age in years.
- **Duration**: Numeric covariate – duration of the condition before beginning treatment.
- **Pain**: Binary response factor with levels No and Yes.

Source


Examples

# Model and analysis shown in the SAS report:
neuralgia.glm <- glm(Pain ~ Treatment * Sex + Age, family = binomial(),
data = neuralgia)
pairs(emmeans(neuralgia.glm, ~ Treatment, at = list(Sex = "F")),
       reverse = TRUE, type = "response", adjust = "bonferroni")
**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 and 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
nutr.aov <- aov(gain ~ (group + age + race)^2, data = nutrition)

# Summarize predictions for age group 3
nutr.emm <- emmeans(nutr.aov, ~ race * group, at = list(age="3"))

em mip(nutr.emm, race ~ group)

# Hispanics seem exceptional; but this doesn't test out due to very sparse data
pairs(nutr.emm, by = "group")
pairs(nutr.emm, by = "race")
```
oranges

Sales of oranges

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

Usage
oranges

Format
A data frame with 36 observations and 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
This is (or once was) available as a SAS sample dataset.

References

Examples
# Example on p.244 of Littell et al.
oranges.lm <- lm(sales1 ~ price1*day, data = oranges)
emmeans(oranges.lm, "day")

# Example on p.246 of Littell et al.
emmeans(oranges.lm, "day", at = list(price1 = 0))

# A more sensible model to consider, IMHO (see vignette("interactions"))
org.mlm <- lm(cbind(sales1, sales2) ~ price1 * price2 + day + store,
               data = oranges)
**pigs**

*Effects of dietary protein on free plasma leucine concentration in pigs*

**Description**

A two-factor experiment with some observations lost

**Usage**

`pigs`

**Format**

A data frame with 29 observations and 3 variables:

- **source**: Source of protein in the diet (factor with 3 levels: fish meal, soybean meal, dried skim milk)
- **percent**: Protein percentage in the diet (numeric with 4 values: 9, 12, 15, and 18)
- **conc**: Concentration of free plasma leucine, in mcg/ml

**Source**


**Examples**

```r
pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
emmeans(pigs.lm, "source")
```

**plot.emmGrid**

*Plot an emmGrid or summary_emm object*

**Description**

Methods are provided to plot EMMs as side-by-side CIs, and optionally to display “comparison arrows” for displaying pairwise comparisons.
Usage

## S3 method for class 'emmGrid'
plot(x, y, type, CIs = TRUE, PIs = FALSE,
     comparisons = FALSE, colors = c("black", "blue", "blue", "red"),
     alpha = 0.05, adjust = "tukey", int.adjust = "none", intervals,
     frequentist, ...)

## S3 method for class 'summary_emm'
plot(x, y, horizontal = TRUE, CIs = TRUE, xlab, ylab,
     layout, scale = NULL, colors = c("black", "blue", "blue", "red"),
     intervals, plotit = TRUE, ...)

Arguments

x  Object of class emmGrid or summary_emm
y  (Required but ignored)
type Character value specifying the type of prediction desired (matching "linear.predictor", "link", or "response"). See details under summary.emmGrid. In addition, the user may specify type = "scale", in which case a transformed scale (e.g., a log scale) is displayed based on the transformation or link function used. Additional customization of this scale is available through including arguments to ggplot2::scale_x_continuous in ...
CIs Logical value. If TRUE, confidence intervals are plotted for each estimate.
PIs Logical value. If TRUE, prediction intervals are plotted for each estimate. If object is a Bayesian model, this requires frequentist = TRUE and sigma = (some value). Note that the PIs option is not available with summary_emm objects – only for emmGrid objects. Also, prediction intervals are not available with engine = "lattice".
comparisons Logical value. 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. (A warning is issued if this can’t be done.) Note that comparison arrows are not available with ‘summary_emm’ objects.
colors Character vector of color names to use for estimates, CIs, PIs, and comparison arrows, respectively. CIs and PIs are rendered with some transparency, and colors are recycled if the length is less than four; so all plot elements are visible even if a single color is specified.
alpha The significance level to use in constructing comparison arrows
adjust Character value: Multiplicity adjustment method for comparison arrows only.
int.adjust Character value: Multiplicity adjustment method for the plotted confidence intervals only.
intervals If specified, it is used to set CIs. This is the previous argument name for CIs and is provided for backward compatibility.
frequentist Logical value. If there is a posterior MCMC sample and frequentist is non-missing and TRUE, a frequentist summary is used for obtaining the plot data,
rather than the posterior point estimate and HPD intervals. This argument is ignored when it is not a Bayesian model.

Additional arguments passed to `update.emmGrid`, `predict.emmGrid`, or `dotplot`

`horizontal` Logical value specifying whether the intervals should be plotted horizontally or vertically

`xlab` Character label for horizontal axis

`ylab` Character label for vertical axis

`layout` Numeric value passed to `dotplot` when `engine == "lattice"`.  

`scale` Object of class `trans` (in the `scales` package) to specify a nonlinear scale. This is used in lieu of `type = "scale"` when plotting a `summary_emm` object created with `type = "response"`. This is ignored with other types of summaries.

`plotit` Logical value. If `TRUE`, a graphical object is returned; if `FALSE`, a data.frame is returned containing all the values used to construct the plot.

### Value

If `plotit = TRUE`, a graphical object is returned.

If `plotit = FALSE`, a data.frame with the table of EMMs that would be plotted. In the latter case, the estimate being plotted is named `the.emmean`, and any factors involved have the same names as in the object. Confidence limits are named `lower.CL` and `upper.CL`, prediction limits are named `lpl` and `upl`, and comparison-arrow limits are named `lcmpl` and `ucmpl`. There is also a variable named `pri.fac` which contains the factor combinations that are not among the groups.

### Details

If any by variables are in force, the plot is divided into separate panels. For "summary_emm" objects, the ... arguments in plot are passed only to dotplot, whereas for "emmGrid" 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. Note that the maximum and minimum estimates have arrows only in one direction, since there is no need to compare them with anything higher or lower, respectively. See the vignette("xplanations", "emmeans") for details on how these are derived.

If `adjust` or `int.adjust` are not supplied, they default to the internal adjust setting saved in `pairs(x)` and `x` respectively (see `update.emmGrid`).

### Note

In order to play nice with the plotting functions, any variable names that are not syntactically correct (e.g., contain spaces) are altered using `make.names`.

### Examples

```r
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.emm <- emmeans(warp.lm, ~ tension | wool)
```
plot(warp.emm)
plot(warp.emm, by = NULL, comparisons = TRUE, adjust = "mvt",
     horizontal = FALSE, colors = "darkgreen")

### Using a transformed scale
pigs.lm <- lm(log(conc + 2) ~ source * factor(percent), data = pigs)
pigs.emm <- emmeans(pigs.lm, ~ percent | source)
plot(pigs.emm, type = "scale", breaks = seq(20, 100, by = 10))

# Based on a summary.
# To get a transformed axis, must specify 'scale'; but it does not necessarily
# have to be the same as the actual response transformation
pigs.ci <- confint(pigs.emm, type = "response")
plot(pigs.ci, scale = scales::log10_trans())

---

**pwpm**  
**Pairwise P-value matrix (plus other statistics)**

### Description

This function presents results from `emmeans` and pairwise comparisons thereof in a compact way. It displays a matrix (or matrices) of estimates, pairwise differences, and P values. The user may opt to exclude any of these via arguments `means`, `difs`, and `pvals`, respectively. To control the direction of the pairwise differences, use `reverse`; and to control what appears in the upper and lower triangle(s), use `flip`. Optional arguments are passed to `contrast.emmGrid` and/or `summary.emmGrid`, making it possible to control what estimates and tests are displayed.

### Usage

```
pwpm(emm, by, reverse = FALSE, pvals = TRUE, means = TRUE,
     diffs = TRUE, flip = FALSE, digits, ...)```

### Arguments

- **emm**  
  An `emmGrid` object

- **by**  
  Character vector of variable(s) in the grid to condition on. These will create different matrices, one for each level or level-combination. If missing, `by` is set to `emm@misc$by.vars`. Grid factors not in `by` are the primary factors: whose levels or level combinations are compared pairwise.

- **reverse**  
  Logical value passed to `pairs.emmGrid`. Thus, FALSE specifies "pairwise" comparisons (earlier vs. later), and TRUE specifies "revpairwise" comparisons (later vs. earlier).

- **pvals**  
  Logical value. If TRUE, the pairwise differences of the EMMs are included in each matrix according to `flip`.

- **means**  
  Logical value. If TRUE, the estimated marginal means (EMMs) from `emm` are included in the matrix diagonal(s).
pwpp

<table>
<thead>
<tr>
<th>diffs</th>
<th>Logical value. If TRUE, the pairwise differences of the EMMs are included in each matrix according to flip.</th>
</tr>
</thead>
<tbody>
<tr>
<td>flip</td>
<td>Logical value that determines where P values and differences are placed. FALSE places the P values in the upper triangle and differences in the lower, and TRUE does just the opposite.</td>
</tr>
<tr>
<td>digits</td>
<td>Integer. Number of digits to display. If missing, an optimal number of digits is determined.</td>
</tr>
<tr>
<td>...</td>
<td>Additional arguments passed to <code>contrast.emmGrid</code> and <code>summary.emmGrid</code>. You should not include method here, because pairwise comparisons are always used.</td>
</tr>
</tbody>
</table>

**Value**

A matrix or ‘list’ of matrices, one for each ‘by’ level.

**See Also**

A graphical display of essentially the same results is available from `pwpp`

**Examples**

```r
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.emm <- emmeans(warp.lm, ~ tension | wool)
pwpm(warp.emm)

# use dot options to specify noninferiority tests
pwpm(warp.emm, by = NULL, side = ">", delta = 5, adjust = "none")
```

**Description**

Constructs a plot of P values associated with pairwise comparisons of estimated marginal means.

**Usage**

```r
pwpp(emm, method = "pairwise", by, sort = TRUE, values = TRUE,
     rows = ".", xlab, ylab, xsub = "", plim = numeric(0), add.space = 0,
     aes, ...)```
Arguments

emmm An emmGrid object

method Character or list. Passed to contrast, and defines the contrasts to be displayed. Any contrast method may be used, provided that each contrast includes one coefficient of 1, one coefficient of -1, and the rest 0. That is, calling contrast(object, method) produces a set of comparisons, each with one estimate minus another estimate.

by Character vector of variable(s) in the grid to condition on. These will create different panels, one for each level or level-combination. Grid factors not in by are the primary factors: whose levels or level combinations are compared pairwise.

sort Logical value. If TRUE, levels of the factor combinations are ordered by their marginal means. If FALSE, they appear in order based on the existing ordering of the factor levels involved. Note that the levels are ordered the same way in all panels, and in many cases this implies that the means in any particular panel will not be ordered even when sort = TRUE.

values Logical value. If TRUE, the values of the EMMs are included in the plot. When there are several side-by-side panels due to by variable(s), the labels showing values start stealing a lot of space from the plotting area; in those cases, it may be desirable to specify FALSE or use rows so that some panels are vertically stacked.

rows Character vector of which by variable(s) are used to define rows of the panel layout. Those variables in by not included in rows define columns in the array of panels. A "." indicates that only one row is used, so all panels are stacked side-by-side.

xlab Character label to use in place of the default for the P-value axis.

ylab Character label to use in place of the default for the primary-factor axis.

xsub Character label used as caption at the lower right of the plot.

plim numeric vector of value(s) between 0 and 1. These are included among the observed p values so that the range of tick marks includes at least the range of plim. Choosing plim = c(0, 1) will ensure the widest possible range.

add.space Numeric value to adjust amount of space used for value labels. Positioning of value labels is tricky, and depends on how many panels and the physical size of the plotting region. This parameter allows the user to adjust the position. Changing it by one unit should shift the position by about one character width (right if positive, left if negative). Note that this interacts with aes$label below.

aes optional named list of lists. Entries considered are point, segment, and label, and contents are passed to the respective ggplot2::geom_xxx() functions. These affect rendering of points, line segments joining them, and value labels. Defaults are point = list(size = 2), segment = list(), and label = list(size = 2.5).

... Additional arguments passed to contrast and summary.emmGrid, as well as to geom_segment and geom_label
Details

Factor levels (or combinations thereof) are plotted on the vertical scale, and P values are plotted on the horizontal scale. Each P value is plotted twice – at vertical positions corresponding to the levels being compared – and connected by a line segment. Thus, it is easy to visualize which P values are small and large, and which levels are compared. In addition, factor levels are color-coded, and the points and half-line segments appear in the color of the other level. The P-value scale is nonlinear, so as to stretch-out smaller P values and compress larger ones. P values smaller than 0.0004 are altered and plotted in a way that makes them more distinguishable from one another.

If xlab, ylab, and xsub are not provided, reasonable labels are created. xsub is used to note special features; e.g., equivalence thresholds or one-sided tests.

Note

The ggplot2 and scales packages must be installed in order for pwpp to work.

Additional plot aesthetics are available by adding them to the returned object; see the examples

See Also

A numerical display of essentially the same results is available from pwpm

Examples

```r
pigs.lm <- lm(log(conc) ~ source * factor(percent), data = pigs)
emm = emmeans(pigs.lm, ~ percent | source)
pwpp(emm)
pwpp(emm, method = "trt.vs.ctrl1", type = "response", side = ">")

# custom aesthetics:
my.aes <- list(point = list(shape = "square"),
                segment = list(linetype = "dashed", color = "red"),
                label = list(family = "serif", fontface = "italic"))
my.pal <- c("darkgreen", "blue", "magenta", "orange")
pwpp(emm, aes = my.aes) + ggplot2::scale_color_manual(values = my.pal)
```

qdrg

Quick and dirty reference grid

Description

This function may make it possible to compute a reference grid for a model object that is otherwise not supported.

Usage

`qdrg(formula, data, coef, vcov, df, mcmc, object, subset, weights, contrasts, link, qr, ordinal.dim, ...)`
Arguments

- **formula**: Formula for the fixed effects
- **data**: Dataset containing the variables in the model
- **coef**: Fixed-effect regression coefficients (must conform to formula)
- **vcov**: Variance-covariance matrix of the fixed effects
- **df**: Error degrees of freedom
- **mcmc**: Posterior sample of fixed-effect coefficients
- **object**: Optional model object. *This rarely works!*; but if provided, we try to set other arguments based on an expectation that ‘object’ has a similar structure to ‘lm’ objects. See Details.
- **subset**: Subset of data used in fitting the model
- **weights**: Weights used in fitting the model
- **contrasts**: List of contrasts specified in fitting the model
- **link**: Link function (character or list) used, if a generalized linear model. (Note: response transformations are auto-detected from formula)
- **qr**: QR decomposition of the model matrix; used only if there are NAs in coef.
- **ordinal.dim**: Integer number of levels in an ordinal response. If not missing, the intercept terms are modified appropriate to predicting the latent response (see vignette("models"), Section O, but note that the other modes, e.g., mode = "prob" are not provided). In this case, we expect the first ordinal.dim - 1 elements of coef to be the estimated threshold parameters, followed by the coefficients for the linear predictor.
- ... Optional arguments passed to ref_grid

Details

Usually, you need to provide formula, data, coef, and vcov, and perhaps other parameters. It is usually fairly straightforward to figure out how to get these from the model object; see the documentation for the model class that was fitted. Sometimes one or more of these quantities contains extra parameters, and if so, you may need to subset them to make everything conformable. For a given formula and data, you can find out what is needed via colnames(model.matrix(formula, data)). (However, for an ordinal model, we expect the first ordinal.dim - 1 coefficients to replace (Intercept). And for a multivariate model, we expect coef to be a matrix with these row names, and vcov to have as many rows and columns as the total number of elements of coef.)

If your model object follows fairly closely the conventions of an lm object, you may be able to get by providing the model as object, plus and (probably) data, and perhaps some other parameters to override the defaults. When object is specified, it is used as detailed below to try to obtain the other arguments. The user should ensure that the defaults shown below do indeed work.

The default values for the arguments are as follows:

- **formula**: Required unless obtainable via formula(object)
- **data**: Required if variables are not in parent.frame() or obtainable via object$data
- **coef**: coef(object)
qdrg

- `vcov`: `vcov(object)`
- `df`: Set to Inf if not available in `object$df.residual`
- `mcmc`: `object$sample`
- `subset`: NULL (so that all observations in data are used)
- `contrasts`: `object$contrasts`

The functions `qdrg` and `emmobj` are close cousins, in that they both produce `emmGrid` objects. When starting with summary statistics for an existing grid, `emmobj` is more useful, while `qdrg` is more useful when starting from a fitted model.

Value

An `emmGrid` object constructed from the arguments

Rank deficiencies

Different model-fitting packages take different approaches when the model matrix is singular, but `qdrg` tries to reconcile them by comparing the linear functions created by `formula` to `coefs` and `vcov`. We may then use the `estimability` package to determine what quantities are estimable. For reconciling to work properly, `coef` should be named and `vcov` should have dimnames. To disable this name-matching action, remove the names from `coef`, e.g., by calling `unname()`. No reconciliation is attempted in multivariate-response cases. For more details on estimability, see the documentation in the `estimability` package.

See Also

`emmobj` for an alternative way to construct an `emmGrid`. 

Examples

```r
if (require(biglm, quietly = TRUE)) withAutoprint({
  # Post hoc analysis of a "biglm" object -- not supported by emmeans
  bigmod <- biglm(log(conc) ~ source + factor(percent), data = pigs)
  rg1 <- qdrg(log(conc) ~ source + factor(percent), data = pigs,
              coef = coef(bigmod), vcov = vcov(bigmod), df = bigmod$df.residual)
  emmeans(rg1, "source", type = "response")
  ## But in this particular case, we could have done it the easy way:
  ## rg1 <- qdrg(object = bigmod, data = pigs)
}, spaced = TRUE)

if(require(coda, quietly = TRUE) && require(lme4, quietly = TRUE))
  withAutoprint({
    # Use a stored example having a posterior sample
    # Model is based on the data in lme4::cbpp
    post <- readRDS(system.file("extdata", "cbpplist", package = "emmeans"))$post.beta
    rg2 <- qdrg(~ size + period, data = lme4::cbpp, mcmc = post, link = "logit")
    summary(rg2, type = "response")
  }, spaced = TRUE)

if(require(ordinal, quietly = TRUE)) withAutoprint({
  wine.clm <- clm(rating ~ temp * contact, data = wine)
  ref_grid(wine.clm)
})
```
# verify that we get the same thing via:
qdrg(object = wine.clm, data = wine, ordinal.dim = 5)
}, spaced = TRUE)

---

**rbind.emmGrid**

Combine or subset emmGrid objects

**Description**

These functions provide methods for `rbind` and `[` that may be used to combine emmGrid objects together, or to extract a subset of cases. The primary reason for doing this would be to obtain multiplicity-adjusted results for smaller or larger families of tests or confidence intervals.

**Usage**

```r
## S3 method for class 'emmGrid'
rbind(..., deparse.level = 1, adjust = "bonferroni")

## S3 method for class 'emmGrid'
e1 + e2

## S3 method for class 'emmGrid'
x[i, adjust, drop.levels = TRUE, ...]

## S3 method for class 'emmGrid'
subset(x, subset, ...)

force_regular(object)

## S3 method for class 'emm_list'
rbind(..., which, adjust = " bonferroni")
```

**Arguments**

- `...` Additional arguments: In `rbind`, object(s) of class `emmGrid`. In `"[", it is ignored. In `subset`, it is passed to `[.emmGrid`
- `deparse.level` (required but not used)
- `adjust` Character value passed to `update.emmGrid`
- `e1` An `emmGrid` object
- `e2` Another `emmGrid` object
- `x` An `emmGrid` object to be subsetted
- `i` Integer vector of indexes
- `drop.levels` Logical value. If TRUE, the "levels" slot in the returned object is updated to hold only the predictor levels that actually occur
\texttt{rbind.emmGrid}

\begin{itemize}
  \item \textbf{subset} \quad \text{logical expression indicating which rows of the grid to keep}
  \item \textbf{object} \quad \text{an object of class \texttt{emmGrid}}
  \item \textbf{which} \quad \text{Integer vector of subset of elements to use; if missing, all are combined}
\end{itemize}

\textbf{Value}

A revised object of class \texttt{emmGrid}

The result of \texttt{e1 + e2} is the same as \texttt{rbind(e1, e2)}

\texttt{force_regular} adds extra (invisible) rows to an \texttt{emmGrid} object to make it a regular grid (all combinations of factors). This regular structure is needed by \texttt{emmeans}. An object can become irregular by, for example, subsetting rows, or by obtaining contrasts of a nested structure.

The \texttt{rbind} method for \texttt{emm_list} objects simply combines the \texttt{emmGrid} objects comprising the first element of \ldots

\textbf{Note}

\texttt{rbind} throws an error if there are incompatibilities in the objects' coefficients, covariance structures, etc. But they are allowed to have different factors; a missing level '...' is added to factors as needed.

\textbf{Examples}

\begin{verbatim}
warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.rg <- ref_grid(warp.lm)

  # Do all pairwise comparisons within rows or within columns,
  # all considered as one family of tests:
  w.t <- pairs(emmeans(warp.rg, ~ wool | tension))
  t.w <- pairs(emmeans(warp.rg, ~ tension | wool))
  rbind(w.t, t.w, adjust = "mvt")
  update(w.t + t.w, adjust = "fdr")  ## same as above except for adjustment

  # Show only 3 of the 6 cases
  summary(warp.rg[c(2, 4, 5)])

  # After-the-fact 'at' specification
  subset(warp.rg, wool == "A")  ## or warp.rg |> subset(wool == "A")

  ### Irregular object
  tmp <- warp.rg[-1]
  ## emmeans(tmp, "tension")  ## will fail because tmp is irregular
  emmeans(force_regular(tmp), "tension")  ## will show some results

  ### Working with 'emm_list' objects
  mod <- lm(conc ~ source + factor(percent), data = pigs)
  all <- emmeans(mod, list(src = pairwise ~ source, pct = consec ~ percent))
  rbind(all, which = c(2, 4), adjust = "mvt")
\end{verbatim}
**ref_grid**

Create a reference grid from a fitted model

**Description**

Using a fitted model object, determine a reference grid for which estimated marginal means are defined. The resulting ref_grid object encapsulates all the information needed to calculate EMMs and make inferences on them.

**Usage**

```r
ref_grid(object, at, cov.reduce = mean, cov.keep = get_emm_option("cov.keep"), mult.names, mult.levs, options = get_emm_option("ref_grid"), data, df, type, regrid, nesting, offset, sigma, nuisance = character(0), non.nuisance, wt.nuis = "equal", rg.limit = get_emm_option("rg.limit"), ...)
```

**Arguments**

- `object`: An object produced by a supported model-fitting function, such as `lm`. Many models are supported. See `vignette("models","emmeans")`.
- `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 `cov.keep` or `at` is reduced according to these specifications. See the section below on “Using `cov.reduce` and `cov.keep`”.
- `cov.keep`: Character vector: names of covariates that are *not* to be reduced; these are treated as factors and used in weighting calculations. `cov.keep` may also include integer value(s), and if so, the maximum of these is used to set a threshold such that any covariate having no more than that many unique values is automatically included in `cov.keep`.
- `mult.names`: Character value: the name(s) to give to the pseudo-factor(s) whose levels delineate the elements of a multivariate response. If this is provided, it overrides the default name(s) used for `class(object)` when it has a multivariate response (e.g., the default is "rep.meas" for "mlm" objects).
- `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.emmGrid`, 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. See the note with `recover_data` for an important precaution.
- `df`: Numeric value. This is equivalent to specifying `options(df = df)`. See `update.emmGrid`.
type Character value. If provided, this is saved as the "predict.type" setting. See `update.emmGrid` and the section below on prediction types and transformations.

regrid Character, logical, or list. If non-missing, the reference grid is reconstructed via `regrid` with the argument `transform = regrid`. See the section below on prediction types and transformations. Note: This argument was named `transform` in version 1.7.2 and earlier. For compatibility with old code, `transform` is still accepted if found among ... , as long as it doesn’t match tran.

nesting If the model has nested fixed effects, this may be specified here via a character vector or named list specifying the nesting structure. Specifying nesting overrides any nesting structure that is automatically detected. See the section below on Recovering or Overriding Model Information.

offset Numeric scalar value (if a vector, only the first element is used). This may be used to add an offset, or override offsets based on the model. A common usage would be to specify `offset = 0` for a Poisson regression model, so that predictions from the reference grid become rates relative to the offset that had been specified in the model.

sigma Numeric value to use for subsequent predictions or back-transformation bias adjustments. If not specified, we use `sigma(object)`, if available, and `NULL` otherwise.

nuisance, non.nuisance, wt.nuis If `nuisance` is a vector of predictor names, those predictors are omitted from the reference grid. Instead, the result will be as if we had averaged over the levels of those factors, with either equal or proportional weights as specified in `wt.nuis` (see the weights argument in `emmeans`). The factors in `nuisance` must not interact with other factors, not even other nuisance factors. Specifying nuisance factors can save considerable storage and computation time, and help avoid exceeding the maximum reference-grid size (`get_emm_option("rg.limit")`).

rg.limit Integer limit on the number of reference-grid rows to allow (checked before any multivariate responses are included).

... Optional arguments passed to `summary.emmGrid`, `emm_basis`, and `recover_data`, such as `params`, `vcov.` (see Covariance matrix below), or options such as `mode` for specific model types (see vignette("models", "emmeans")).

Details

To users, the `ref_grid` function itself is important because most of its arguments are in effect arguments of `emmeans` and related functions, in that those functions pass their ... arguments to `ref_grid`

The reference grid consists of combinations of independent variables over which predictions are made. Estimated marginal 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 data. 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. See also the section below on recovering/overriding model information.
Value

An object of the S4 class "emmGrid" (see emmGrid-class). These objects encapsulate everything needed to do calculations and inferences for estimated marginal means, and contain nothing that depends on the model-fitting procedure.

Using cov.reduce and cov.keep

The cov.keep argument was not available in emmeans versions 1.4.1 and earlier. Any covariates named in this list are treated as if they are factors: all the unique levels are kept in the reference grid. The user may also specify an integer value, in which case any covariate having no more than that number of unique values is implicitly included in cov.keep. The default for cov.keep is set and retrieved via the emm_options framework, and the system default is "2", meaning that covariates having only two unique values are automatically treated as two-level factors. See also the Note below on backward compatibility.

There is a subtle distinction between including a covariate in cov.keep and specifying its values manually in at: Covariates included in cov.keep are treated as factors for purposes of weighting, while specifying levels in at will not include the covariate in weighting. See the mtcars.lm example below for an illustration.

cov.reduce may be a function, logical value, formula, or a named list of these. 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 including all covariates in cov.keep. Use of 'cov.reduce = FALSE' is inadvisable because it can result in a huge reference grid; it is far better to use cov.keep.

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.

To allow for situations where a simple lm() call as described above won’t be adequate, a formula of the form ext ~ fcnname is also supported, where the left-hand side may be ext, extern, or external (and must not be a predictor name) and the right-hand side is the name of an existing function. The function is called with one argument, a data frame with columns for each variable in the reference grid. The function is expected to use that frame as new data to be used to obtain predictions for one or more models; and it should return a named list or data frame with replacement values for one or more of the covariates.

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 of cov.keep specification for a covariate also named in at is ignored.

Interdependent covariates

Care must be taken when covariate values depend on one another. For example, when a polynomial model was fitted using predictors x, x2 (equal to x^2), and x3 (equal to x^3), the reference grid will by default set x2 and x3 to their means, which is inconsistent. The user should instead use the at argument to set these to the square and cube of mean(x). Better yet, fit the model using a formula involving poly(x, 3) or I(x^2) and I(x^3); then there is only x appearing as a covariate; it will
be set to its mean, and the model matrix will have the correct corresponding quadratic and cubic
terms.

**Matrix covariates**

Support for covariates that appear in the dataset as matrices is very limited. If the matrix has but
one column, it is treated like an ordinary covariate. Otherwise, with more than one column, each
column is reduced to a single reference value – the result of applying `cov.reduce` to each column
(averaged together if that produces more than one value); you may not specify values in `at`; and
they are not treated as variables in the reference grid, except for purposes of obtaining predictions.

**Recovering or overriding model information**

Ability to support a particular class of object depends on the existence of `recover_data` and
`emm_basis` methods – see `extending-emmeans` for details. The call methods("recover_data")
will help identify these.

**Data.** 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.)

**Covariance matrix.** 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.` argu-
ment, 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 arguments (`object, ...`). Be careful with possible unintended conflicts with arguments in
`...`; for example, `sandwich::vcovHAC()` has optional arguments `adjust` and `weights`
that may be intended for `emmeans()` but will also be passed to `vcov()`.

**Nested factors.** Having a nesting structure affects marginal averaging in `emmeans` in that it is done
separately for each level (or combination thereof) of the grouping 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.emmGrid`. The `nesting` argument
may be a character vector, a named list, or NULL. If a list, each name should be the name of a sin-
gle factor in the grid, and its entry a character vector of the name(s) of its grouping factor(s). `nested`
may also be a character value of the form "factor1 %in% (factor2*factor3)" (the parentheses are optional).
If there is more than one such specification, they may be appended separated by
commas, or as separate elements of a character vector. For example, these specifications are equi-
vant: `nesting = list(state = "country", city = c("state", "country"), nesting = "state
%in% country, city %in% (state*country)", and nesting = c("state %in% country", "city
%in% state*country")

**Predictors with subscripts and data-set references**

When the fitted model contains subscripts or explicit references to data sets, the reference grid may
optionally be post-processed to simplify the variable names, depending on the `simplify.names` op-
tion (see `emm_options`), which by default is `TRUE`. For example, if the model formula is `data1$resp`
The simplified predictor names (for use, e.g., in the specs for \texttt{emmeans}) will be `trt`, `data2[[3]]`, and `cov`. Numerical subscripts are not simplified; nor are variables having simplified names that coincide, such as if `data2$trt` were also in the model.

Please note that this simplification is performed after the reference grid is constructed. Thus, non-simplified names must be used in the \texttt{at} argument (e.g., \texttt{at = list(`data2["cov"] = 2:4')}).

If you don’t want names simplified, use \texttt{emm_options(simplify.names = FALSE)}.

### Prediction types and transformations

Transformations can exist because of a link function in a generalized linear model, or as a response transformation, or even both. In many cases, they are auto-detected, for example a model formula of the form `\texttt{sqrt(y)} ~ ...`. Even transformations containing multiplicative or additive constants, such as `\texttt{2*sqrt(y + pi)} ~ ...`, are auto-detected. A response transformation of \(y + 1\) ~ ... is not auto-detected, but \(\text{I}(y + 1) ~ ...\) is interpreted as \(\text{identity}(y + 1) ~ ...\). A warning is issued if it gets too complicated. Complex transformations like the Box-Cox transformation are not auto-detected; but see the help page for \texttt{make.tran} for information on some advanced methods.

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

**Order-of-processing issues:** The \texttt{regrid} argument, if present, is acted on immediately after the reference grid is constructed, while some of the \ldots arguments may be used to update the object at the very end. Thus, code like \texttt{ref_grid(mod, tran = "sqrt", regrid = "response")} will not work correctly if the intention was to specify the response transformation, because the re-grid is done \texttt{before} it processes \texttt{tran = "sqrt"}. To get the intended result, do \texttt{regrid(ref_grid(mod, tran = "sqrt"), transform = "response")}.

### Optional side effect

If the \texttt{save.ref_grid} option is set to \texttt{TRUE} (see \texttt{emm_options}), the most recent result of \texttt{ref_grid}, whether called directly or indirectly via \texttt{emmeans}, \texttt{emtrends}, or some other function that calls one of these, is saved in the user's environment as \texttt{.Last.ref_grid}. This facilitates checking what reference grid was used, or reusing the same reference grid for further calculations. This automatic saving is disabled by default, but may be enabled via \texttt{`emm_options(save.ref_grid = TRUE)`}.

### Note

The system default for \texttt{cov.keep} causes models containing indicator variables to be handled differently than in \texttt{emmeans} version 1.4.1 or earlier. To replicate older analyses, change the default via \texttt{`emm_options(cov.keep = character(0))`}.

Some earlier versions of \texttt{emmeans} offer a \texttt{covnest} argument. This is now obsolete; if \texttt{covnest} is specified, it is harmlessly ignored. Cases where it was needed are now handled appropriately via the code associated with \texttt{cov.keep}.  

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See Also

Reference grids are of class `emmGrid`, and several methods exist for them — for example `summary.emmGrid`. Reference grids are fundamental to `emmeans`. Supported models are detailed in `vignette("models", "emmeans")`. See `update.emmGrid` for details of arguments that can be in `options` (or in `...`).

Examples

```r
fiber.lm <- lm(strength ~ machine*diameter, data = fiber)
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:
summary(ref_grid(fiber.lm, vcov. = sandwich::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)

### Model with indicator variables as predictors:
mtcars.lm <- lm(mpg ~ disp + wt + vs * am, data = mtcars)
(rg.default <- ref_grid(mtcars.lm))
(rg.nokeep <- ref_grid(mtcars.lm, cov.keep = character(0)))
(rg.at <- ref_grid(mtcars.lm, at = list(vs = 0:1, am = 0:1)))

# Two of these have the same grid but different weights:
rg.default@grid
rg.at@grid

### Using cov.reduce formulas...
# Above suggests we can vary disp indep. of other factors - unrealistic
rg.alt <- ref_grid(mtcars.lm, at = list(wt = c(2.5, 3, 3.5)),
                  cov.reduce = disp ~ vs * wt)
rg.alt@grid

# Alternative to above where we model sqrt(disp)
disp.mod <- lm(sqrt(disp) ~ vs * wt, data = mtcars)
disp.fun <- function(dat)
  list(disp = predict(disp.mod, newdata = dat)^2)
rg.alt2 <- ref_grid(mtcars.lm, at = list(wt = c(2.5, 3, 3.5)),
                   cov.reduce = external ~ disp.fun)
rg.alt2@grid

# Multivariate example
MOats.lm = lm(yield ~ Block + Variety, data = MOats)
ref_grid(MOats.lm, mult.names = "nitro")

# Silly illustration of how to use 'mult.levs' to make comb's of two factors
```

ref_grid
ref_grid(MOats.lm, mult.levs = list(T=LETTERS[1:2], U=letters[1:2]))

# Using 'params'
require("splines")
my.knots = c(2.5, 3, 3.5)
mod = lm(Sepal.Length ~ Species * ns(Sepal.Width, knots = my.knots), data = iris)
## my.knots is not a predictor, so need to name it in 'params'
ref_grid(mod, params = "my.knots")

---

**regrid**

*Reconstruct a reference grid with a new transformation or simulations*

**Description**

The typical use of this function is to cause EMMs to be computed on a different scale, e.g., the back-transformed scale rather than the linear-predictor scale. In other words, if you want back-transformed results, do you want to average and then back-transform, or back-transform and then average?

**Usage**

```r
regrid(object, transform = c("response", "mu", "unlink", "none", "pass", 
links), inv.link.lbl = "response", predict.type, 
bias.adjust = get_emm_option("back.bias.adj"), sigma, N.sim, 
sim = mvtnorm::rmvnorm, ...)
```

**Arguments**

- `object`: An object of class `emmGrid`
- `transform`: Character, list, or logical value. If "response", "mu", or TRUE, 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 "none" or FALSE, object is re-gridded so that its bhat slot contains predict(object) and its linfct slot is the identity. Any internal transformation information is preserved. If `transform = "pass"`, the object is not re-gridded in any way (this may be useful in conjunction with `N.sim`). If `transform` is a character value in `links` (which is the set of valid arguments for the `make.link` function, excepting "identity"), or if `transform` is a list of the same form as returned by `make.links` or `make.tran`, the results are formulated as if the response had been transformed with that link function.
- `inv.link.lbl`: Character value. This applies only when `transform` is in `links`, and is used to label the predictions if subsequently summarized with `type = "response"`.
- `predict.type`: Character value. If provided, the returned object is updated with the given type to use by default by `summary.emmGrid` (see `update.emmGrid`). This may be useful if, for example, when one specifies `transform = "log"` but desires summaries to be produced by default on the response scale.
bias.adjust Logical value for whether to adjust for bias in back-transforming (transform = "response"). This requires a value of sigma to exist in the object or be specified.

sigma Error SD assumed for bias correction (when transform = "response" and a transformation is in effect). If not specified, object@misc$sigma is used, and an error is thrown if it is not found.

N.sim Integer value. If specified and object is based on a frequentist model (i.e., does not have a posterior sample), then a fake posterior sample is generated using the function sim.

sim A function of three arguments (no names are assumed). If N.sim is supplied with a frequentist model, this function is called with respective arguments N.sim, object@bhat, and object@V. The default is the multivariate normal distribution.

... Ignored.

Details

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, when transform = "response", the result of summary.emmGrid after applying regrid is identical to the summary of the original object using ‘type="response"’. But subsequent EMMs or contrasts will be conducted on the new scale – which is the reason this function exists.

This function may also be used to simulate a sample of regression coefficients for a frequentist model for subsequent use as though it were a Bayesian model. To do so, specify a value for N.sim and a sample is simulated using the function sim. The grid may be further processed in accordance with the other arguments; or if transform = "pass", it is simply returned with the only change being the addition of the simulated sample.

Value

An emmGrid object with the requested changes

Degrees of freedom

In cases where the degrees of freedom depended on the linear function being estimated (e.g., Satterthwaite method), 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. An annotation is displayed below any subsequent summary results stating that the degrees-of-freedom method is inherited from the previous method at the time of re-gridding.

Note

Another way to use regrid is to supply a regrid argument to ref.grid (either directly or indirectly via emmeans), in which case its value is passed to regrid as transform. This is often a simpler approach if the reference grid has not already been constructed.
Examples

```r
pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
rg <- ref_grid(pigs.lm)
# This will yield EMMs as GEOMETRIC means of concentrations:
(emm1 <- emmeans(rg, "source", type = "response"))
pairs(emm1) ## We obtain RATIOS

# This will yield EMMs as ARITHMETIC means of concentrations:
(emm2 <- emmeans(regrid(rg, transform = "response"), "source"))
pairs(emm2) ## We obtain DIFFERENCES
# Same result, useful if we hadn't already created 'rg'
# emm2 <- emmeans(pigs.lm, "source", regrid = "response")

# Simulate a sample of regression coefficients
set.seed(2.71828)
rgb <- regrid(rg, N.sim = 200, transform = "pass")
emmeans(rgb, "source", type = "response")  ## similar to emm1
```

---

str.emmGrid

**Description**

Miscellaneous methods for emmGrid objects

**Usage**

```r
## S3 method for class 'emmGrid'
str(object, ...)

## S3 method for class 'emmGrid'
print(x, ..., export = FALSE)

## S3 method for class 'emmGrid'
vcov(object, ...)
```

**Arguments**

- `object` An emmGrid object
- `...` (required but not used)
- `x` An emmGrid object
- `export` Logical value. If FALSE, the object is printed. If TRUE, a list is invisibly returned, which contains character elements named summary and annotations that may be saved or displayed as the user sees fit. summary is a character matrix (or list of such matrices, if a by variable is in effect). annotations is a character vector of the annotations that would have been printed below the summary or summaries.
**Value**

The `vcov` method returns a symmetric matrix of variances and covariances for `predict.emmGrid(object, type = "lp")`

---

**summary.emmGrid**

Summaries, predictions, intervals, and tests for `emmGrid` objects

**Description**

These are the primary methods for obtaining numerical or tabular results from an `emmGrid` object. `summary.emmGrid` is the general function for summarizing `emmGrid` objects. It also serves as the print method for these objects; so for convenience, `summary()` arguments may be included in calls to functions such as `emmeans` and `contrast` that construct `emmGrid` objects. Note that by default, summaries for Bayesian models are diverted to `hpd.summary`.

**Usage**

```r
## S3 method for class 'emmGrid'
summary(object, infer, level, adjust, by,
         cross.adjust = "none", type, df, calc, null, delta, side, frequentist,
         bias.adjust = get_emm_option("back.bias.adj"), sigma, ...)

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

test(object, null, ...)

## S3 method for class 'emmGrid'
predict(object, type, interval = c("none", "confidence", "prediction"), level = 0.95,
         bias.adjust = get_emm_option("back.bias.adj"), sigma, ...)

## S3 method for class 'summary_emm'
as.data.frame(x, row.names = NULL, optional,
               check.names = TRUE, destroy.annotations = FALSE, ...)

## S3 method for class 'summary_emm'
x[... , as.df = TRUE]
```

**Arguments**

- `object` An object of class "emmGrid" (see `emmGrid-class`)
infer
A vector of one or 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
Numerical value between 0 and 1. 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 the *P*-value adjustments section
below.

by
Character name(s) of variables to use for grouping into separate tables. This
affects the family of tests considered in adjusted *P* values.

cross.adjust
Character: *P*-value adjustment method to additionally apply across the by groups.
See the section on *P*-value adjustments for details.

type
Character: type of prediction desired. 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.predictor"; these are equiva-
 lent, 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 `emm_options`, and also the section below on transformations and
links.

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

calc
Named list of character value(s) or formula(s). The expressions in char are eval-
uated and appended to the summary, just after the df column. The expression
may include any names up through df in the summary, any additional names in
object@grid (such as .wgt. or .offset.), or any earlier elements of calc.

null
Numeric. Null hypothesis value(s), on the linear-predictor scale, 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 (on the linear-predictor scale). If zero, ordinary tests of signif-
icance are performed. If positive, this specifies a threshold for testing equiva-
 lence (using the TOST or two-one-sided-test method), non-inferiority, or non-
superiority, depending on side. See Details for how the test statistics are de-

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 ":="). See the special section below for more details.

frequentist
Ignored except if a Bayesian model was fitted. If missing or FALSE, the object is
passed to `hpd.summary`. Otherwise, a logical value of TRUE will have it return a
frequentist summary.

bias.adjust
Logical value for whether to adjust for bias in back-transforming (type = "response").
This requires a value of sigma to exist in the object or be specified.
summary.emmGrid

sigma Error SD assumed for bias correction (when type = "response" and a transformation is in effect), or for constructing prediction intervals. If not specified, object@misc$sigma is used, and an error is thrown if it is not found. Note: sigma may be a vector, as long as it conforms to the number of rows of the reference grid.

... Optional arguments such as scheffe.rank (see “P-value adjustments”). In confint.emmGrid, predict.emmGrid, and test.emmGrid, these arguments are passed to summary.emmGrid.

parm (Required argument for confint methods, but not used)

joint Logical value. If FALSE, the arguments are passed to summary.emmGrid with infer=c(FALSE, TRUE). If joint = TRUE, a joint test of the hypothesis L beta = null is performed, where L is object@linfct and beta is the vector of fixed effects estimated by object@betahat. This will be either an F test or a chi-square (Wald) test depending on whether degrees of freedom are available. See also joint_tests.

verbose Logical value. If TRUE and joint = TRUE, a table of the effects being tested is printed.

rows 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.

status logical. If TRUE, a note column showing status flags (for rank deficiencies and estimability issues) is displayed even when empty. If FALSE, the column is included only if there are such issues.

interval Type of interval desired (partial matching is allowed): "none" for no intervals, otherwise confidence or prediction intervals with given arguments, via confint.emmGrid.

x object of the given class

row.names passed to as.data.frame

optional required argument, but ignored in as.data.frame.emmGrid

check.names passed to data.frame

destroy.annotations Logical value. If FALSE, an object of class summary_emm is returned (which inherits from data.frame), but if displayed, details like confidence levels, P-value adjustments, transformations, etc. are also shown. If TRUE (not recommended), a "plain vanilla" data frame is returned, based on row.names and check.names.

as.df Logical value. With x[...], as.df = TRUE], the result is object is coerced to an ordinary data.frame; otherwise, it is left as a summary_emm object.

Details

confint.emmGrid is equivalent to summary.emmGrid with infer = c(TRUE, FALSE). The function test.emmGrid, when called with joint = FALSE, is equivalent to summary.emmGrid with infer = c(FALSE, TRUE).

With joint = TRUE, test.emmGrid calculates the Wald test of the hypothesis linfct %*% bhat = null, where linfct and bhat refer to slots in object (possibly subsetted according to by or
rows). An error is thrown if any row of linfct is non-estimable. It is permissible for the rows of linfct to be linearly dependent, as long as null == 0, in which case a reduced set of contrasts is tested. Linear dependence and nonzero null cause an error. The returned object has an additional "est.fcns" attribute, which is a list of the linear functions associated with the joint test.

Value

summary.emmGrid, confint.emmGrid, and test.emmGrid return an object of class "summary_emm", which is an extension of data.frame but with a special print method that displays it with custom formatting. For models fitted using MCMC methods, the call is diverted to hpd.summary (with prob set to level, if specified); one may alternatively use general MCMC summarization tools with the results of as.mcmc.

predict returns a vector of predictions for each row of object@grid.

The as.data.frame method returns an object that inherits from "data.frame".

Defaults

The misc slot in object may contain default values for by, calc, 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 'emmeans_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.

When bias.adjust is TRUE, then back-transformed estimates are adjusted by adding $0.5h''(u)\sigma^2$, where h is the inverse transformation and u is the linear predictor. This is based on a second-order Taylor expansion. There are better or exact adjustments for certain specific cases, and these may be incorporated in future updates.

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 rbind.emmGrid 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 \(\sqrt{rF(\alpha; r, d)}\), where d is the error degrees of freedom and r is the rank of the set of linear functions under consideration. By default, the value of r is computed from object@linfct for each by group; however, if the user specifies an argument matching scheffe.rank, its value will be used instead. Ordinarily, if there are k means involved, then r = k − 1 for a full set of contrasts involving all k means, and r = k for the means themselves. (The Scheffe adjustment is 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.

"dunnettx" Uses our own ad hoc 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 same 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", "dunnettx", or others more attractive.

"none" Makes no adjustments to the $p$ values.

For tests, not confidence intervals, 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.

Warning: Non-estimable cases are included in the family to which adjustments are applied. You may wish to subset the object using the [] operator to work around this problem.

The cross.adjust argument is a way of specifying a multiplicity adjustment across the by groups (otherwise by default, each group is treated as a separate family in regards to multiplicity adjustments). It applies only to $p$ values. Valid options are one of the `p.adjust.methods` or "sidak". This argument is ignored unless it is other than "none", there is more than one by group, and they are all the same size. Under those conditions, we first use `adjust` to determine the within-group adjusted $p$ values. Imagine each group’s adjusted $p$ values arranged in side-by-side columns, thus forming a matrix with the number of columns equal to the number of by groups. Then we use the cross.adjust method to further adjust the adjusted $p$ values in each row of this matrix. Note that an overall Bonferroni (or Sidak) adjustment is obtainable by specifying both `adjust` and `cross.adjust` as "bonferroni" (or "sidak"). However, less conservative (but yet conservative) overall adjustments are available when it is possible to use an “exact” within-group method (e.g., `adjust = "tukey"` for pairwise comparisons) and `cross.adjust` as a conservative adjustment. `cross.adjust` methods other than "none", "bonferroni", or "sidak" do not seem advisable, but other `p.adjust` methods are available if you can make sense of them.]
Tests of significance, nonsuperiority, noninferiority, or equivalence

When \( \delta = 0 \), test statistics are the usual tests of significance. They are of the form \( \frac{\text{estimate} - \text{null}}{\text{SE}} \). Notationally:

Significance \( H_0 : \theta = \theta_0 \) versus

- \( H_1 : \theta < \theta_0 \) (left-sided), or
- \( H_1 : \theta > \theta_0 \) (right-sided), or
- \( H_1 : \theta \neq \theta_0 \) (two-sided)

The test statistic is

\[
t = \frac{(Q - \theta_0)}{SE}
\]

where \( Q \) is our estimate of \( \theta \); then left, right, or two-sided \( p \) values are produced, depending on side.

When \( \delta \) is positive, the test statistic depends on \( \text{side} \) as follows.

**Left-sided (nonsuperiority)** \( H_0 : \theta \geq \theta_0 + \delta \) versus \( H_1 : \theta < \theta_0 + \delta \)

\[
t = \frac{(Q - \theta_0 - \delta)}{SE}
\]

The \( p \) value is the lower-tail probability.

**Right-sided (noninferiority)** \( H_0 : \theta \leq \theta_0 - \delta \) versus \( H_1 : \theta > \theta_0 - \delta \)

\[
t = \frac{(Q - \theta_0 + \delta)}{SE}
\]

The \( p \) value is the upper-tail probability.

**Two-sided (equivalence)** \( H_0 : |\theta - \theta_0| \geq \delta \) versus \( H_1 : |\theta - \theta_0| < \delta \)

\[
t = \frac{|Q - \theta_0| - \delta}{SE}
\]

The \( p \) value is the lower-tail probability.

Note that \( t \) is the maximum of \( t_{\text{nonsup}} \) and \( -t_{\text{noninf}} \). This is equivalent to choosing the less significant result in the two-one-sided-test (TOST) procedure.

Non-estimable cases

When the model is rank-deficient, each row \( x \) of object’s \( \text{linfct} \) slot is checked for estimability. If \( \sum(x \times \text{bhat}) \) is found to be non-estimable, then the string \( \text{NonEst} \) is displayed for the estimate, and associated statistics are set to \( \text{NA} \). The estimability check is performed using the orthonormal basis \( N \) in the \( \text{nbasis} \) slot for the null space of the rows of the model matrix. Estimability fails when \( \frac{||Nx||^2}{||x||^2} \) exceeds \( \text{tol} \), which by default is \( 1e-8 \). You may change it via \( \text{emm_options} \) by setting \( \text{estble.tol} \) to the desired value.

See the warning above that non-estimable cases are still included when determining the family size for \( P \)-value adjustments.

Warning about potential misuse of \( P \) values

Some in the statistical and scientific community argue that the term “statistical significance” should be completely abandoned, and that criteria such as “\( p < 0.05 \)” never be used to assess the importance of an effect. These practices can be too misleading and are prone to abuse. See the “basics” vignette for more discussion.

Note

In doing testing and a transformation and/or link is in force, any \( \text{null} \) and/or \( \text{delta} \) values specified must always be on the scale of the linear predictor, regardless of the setting for ‘\( \text{type} \)’. If \( \text{type} = \)
"response", the null value displayed in the summary table will be back-transformed from the value supplied by the user. But the displayed delta will not be changed, because there (often) is not a natural way to back-transform it.

When we have type = "response", and bias.adj = TRUE, the null value displayed in the output is both back-transformed and bias-adjusted, leading to a rather non-intuitive-looking null value. However, since the tests themselves are performed on the link scale, this is the response value at which a *P* value of 1 would be obtained.

The default show method for emmGrid objects (with the exception of newly created reference grids) is print(summary()). Thus, with ordinary usage of emmeans and such, it is unnecessary to call summary unless there is a need to specify other than its default options.

If a data frame is needed, summary, confint, and test serve this need. as.data.frame routes to summary by default; calling it with destroy.annotations = TRUE is not recommended for exactly that reason. If you want to see more digits in the output, use print(summary(object), digits = ...); and if you always want to see more digits, use emm_options(opt.digits = FALSE).

See Also

hpd.summary

Examples

warp.lm <- lm(breaks ~ wool * tension, data = warpbreaks)
warp.emm <- emmeans(warp.lm, ~ tension | wool)
warp.emm  # implicitly runs 'summary'

confint(warp.emm, by = NULL, level = .90)

# --------------------------------------------------------------
pigs.lm <- lm(log(conc) ~ source + factor(percent), data = pigs)
pigs.emm <- emmeans(pigs.lm, "percent", type = "response")
summary(pigs.emm)  # (inherits type = "response")
summary(pigs.emm, calc = c(n = ".wgt."))  # Show sample size

# For which percents is EMM non-inferior to 35, based on a 10% threshold?
# Note the test is done on the log scale even though we have type = "response"
test(pigs.emm, null = log(35), delta = log(1.10), side = ">")

con <- contrast(pigs.emm, "consec")
test(con)

test(con, joint = TRUE)

# default Scheffe adjustment - rank = 3
summary(con, infer = c(TRUE, TRUE), adjust = "scheffe")

# Consider as some of many possible contrasts among the six cell means
summary(con, infer = c(TRUE, TRUE), adjust = "scheffe", scheffe.rank = 5)

# Show estimates to more digits
print(test(con), digits = 7)
# Cross-adjusting P values
prs <- pairs(warp.emm)  # pairwise comparisons of tension, by wool
 test(prs, adjust = "tukey", cross.adjust = "bonferroni")

# Same comparisons taken as one big family (more conservative)
 test(prs, adjust = "bonferroni", by = NULL)

ubds

Description

This is a simulated unbalanced dataset with three factors and two numeric variables. There are true relationships among these variables. This dataset can be useful in testing or illustrating messy-data situations. There are no missing data, and there is at least one observation for every factor combination; however, the "cells" attribute makes it simple to construct subsets that have empty cells.

Usage

ubds

Format

A data frame with 100 observations, 5 variables, and a special "cells" attribute:

A  Factor with levels 1, 2, and 3
B  Factor with levels 1, 2, and 3
C  Factor with levels 1, 2, and 3
x  A numeric variable
y  A numeric variable

In addition, attr(ubds, "cells") consists of a named list of length 27 with the row numbers for each combination of A, B, C. For example, attr(ubds, "cells")["213"] has the row numbers corresponding to levels A == 2, B == 1, C == 3. The entries are ordered by length, so the first entry is the cell with the lowest frequency.

Examples

# Omit the three lowest-frequency cells
low3 <- unlist(attr(ubds, "cells")[1:3])
messy.lm <- lm(y ~ (x + A + B + C)^3, data = ubds, subset = -low3)
update.emmGrid

Update an emmGrid object

Description

Objects of class emmGrid contain several settings that affect such things as what arguments to pass to `summary.emmGrid`. The update method allows safer management of these settings than by direct modification of its slots.

Usage

```r
## S3 method for class 'emmGrid'
update(object, ..., silent = FALSE)

## S3 replacement method for class 'emmGrid'
levels(x) <- value

## S3 method for class 'summary_emm'
update(object, by.vars, mesg, ...)
```

Arguments

- `object` An emmGrid object
- `...` Options to be set. These must match a list of known options (see Details)
- `silent` Logical value. If FALSE (the default), a message is displayed if any options are not matched. If TRUE, no messages are shown.
- `x` an emmGrid object
- `value` list or replacement levels. See the documentation for `update.emmGrid` with the `levels` argument, as well as the section below on “Replacing levels”
- `by.vars, mesg` Attributes that can be altered in `update.summary_emm`

Value

- an updated emmGrid object.
- `levels<-` replaces the levels of the object in-place. See the section on replacing levels for details.

Details

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

- `tran, tran2` (list or character) specifies the transformation which, when inverted, determines the results displayed by `summary.emmGrid, predict.emmGrid`, or `emmip` 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.

`tran.offset` Additive constant before a transformation is applied. For example, a response transformation of `log(y + pi)` has `tran.offset = pi`. If no value is present, an offset of 0 is assumed.

`estName` (character) is the column label used for displaying predictions or EMMs.

`inv.lbl` (character) is the column label to use for predictions or EMMs 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.emmGrid`.

`alpha` (numeric) is the default significance level for tests, in `summary.emmGrid` as well as `plot.emmGrid` when ‘CIs = TRUE’. Be cautious that methods that depend on specifying `alpha` are prone to abuse. See the discussion in vignette("basics", "emmeans").

`adjust` (character) is the default for the `adjust` argument in `summary.emmGrid`.

`famSize` (integer) is the number of means involved in a family of inferences; used in Tukey adjustment.

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

`level` (numeric) is the default confidence level, level, in `summary.emmGrid`. Note: You must specify all five letters of 'level' to distinguish it from the slot name 'levels'.

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

`calc` (list) additional calculated columns. See `summary.emmGrid`.

`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).

`sigma` (numeric) Error SD to use in predictions and for bias-adjusted back-transformations

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

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

`bias.adjust`, `frequentist` (character) These are used by `summary` if the value of these arguments are not specified.

`estType` (character) is used internally to determine what adjust methods are appropriate. It should match one of c("prediction", "contrast", "pairs"). As an example of why this is needed, the Tukey adjustment should only be used for pairwise comparisons (`estType = "pairs"`); if `estType` is some other string, Tukey adjustments are not allowed.
avgd.over (character) vector) are the names of the variables whose levels are averaged over in obtaining marginal averages of predictions, i.e., estimated marginal 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.emmGrid display.

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

nesting (Character or named list) specifies the nesting structure. See “Recovering or overriding model information” in the documentation for ref_grid. The current nesting structure is displayed by str.emmGrid.

levels named list of new levels for the elements of the current emmGrid. The list name(s) are used as new variable names, and if needed, the list is expanded using expand.grid. These results replace current variable names and levels. This specification changes the levels, grid, roles, and misc slots in the updated emmGrid, and resets pri.vars, by.vars, adjust, famSize, and avgd.over. In addition, if there is nesting of factors, that may be altered; a warning is issued if it involves something other than mere name changes. Note: All six letters of levels is needed in order to distinguish it from level.

submodel formula or character value specifying a submodel (requires this feature being supported by underlying methods for the model class). When specified, the linfct slot is replaced by its aliases for the specified sub-model. Any factors in the sub-model that do not appear in the model matrix are ignored, as are any interactions that are not in the main model, and any factors associate with multivariate responses. The estimates displayed are then computed as if the sub-model had been fitted. (However, the standard errors will be based on the error variance(s) of the full model.) Note: The formula should refer only to predictor names, excluding any function calls (such as factor or poly) that appear in the original model formula. See the example. The character values allowed should partially match "minimal" or "type2". With "minimal", the sub-model is taken to be the one only involving the surviving factors in object (the ones averaged over being omitted). Specifying "type2" is the same as "minimal" except only the highest-order term in the sub-model is retained, and all effects not containing it are orthogonalized-out. Thus, in a purely linear situation such as an lm model, the joint test of the modified object is in essence a type-2 test as in car::Anova.

For some objects such as generalized linear models, specifying submodel will typically not produce the same estimates or type-2 tests as would be obtained by actually fitting a separate model with those specifications. The reason is that those models are fitted by iterative-reweighting methods, whereas the submodel calculations preserve the final weights used in fitting the full model.

(any other slot name) If the name matches an element of slotNames(object) other than levels, that slot is replaced by the supplied value, if it is of the required class (otherwise an error occurs).

The user must be very careful in replacing slots because they are interrelated; for example, the lengths and dimensions of grid, linfct, bhat, and V must conform.

Replacing levels

The levels<- method uses update.emmGrid to replace the levels of one or more factors. This method allows selectively replacing the levels of just one factor (via subsetting operators), whereas
update(x, levels = list(...)) requires a list of all factors and their levels. If any factors are to be renamed, we must replace all levels and include the new names in the replacements. See the examples.

**Method for summary_emm objects**

This method exists so that we can change the way a summary is displayed, by changing the by variables or the annotations.

**Note**

When it makes sense, an option set by update will persist into future results based on that object. But some options are disabled as well. For example, a calc option will be nulled-out if contrast is called, because it probably will not make sense to do the same calculations on the contrast results, and in fact the variable(s) needed may not even still exist. factor(percent).

**See Also**

emm_options

**Examples**

```r
# Using an already-transformed response:
pigs.lm <- lm(log(conc) ~ source * factor(percent), data = pigs)

# Reference grid that knows about the transformation
# and asks to include the sample size in any summaries:
pigs.rg <- update(ref_grid(pigs.lm), tran = "log",
                  predict.type = "response",
                  calc = c(n = ~.wgt.))
emmeans(pigs.rg, "source")

# Obtain estimates for the additive model
# [Note that the submodel refers to 'percent', not 'factor(percent)']
emmeans(pigs.rg, "source", submodel = ~ source + percent)

# Type II ANOVA
joint_tests(pigs.rg, submodel = "type2")

## Changing levels of one factor
newrg <- pigs.rg
levels(newrg)$source <- 1:3
newrg

## Unraveling a previously standardized covariate
zd = scale(fiber$diameter)
fibz.lm <- lm(strength ~ machine * zd, data = fiber)
(fibz.rg <- ref_grid(fibz.lm, at = list(zd = -2:2))) ### 2*SD range
lev <- levels(fibz.rg)
levels(fibz.rg) <- list (  
  machine = lev$machine,  
  diameter = with(attributes(zd),
```
### Compactify results with a by variable

```r
update(joint_tests(pigs.rg, by = "source"), by = NULL)
```

---

**xtable.emmGrid**

`xtable.emmGrid`  
*Using *xtable* for EMMs*

**Description**

These methods provide support for the *xtable* package, enabling polished presentations of tabular output from *emmeans* and other functions.

**Usage**

```r
## S3 method for class 'emmGrid'
xtable(x, caption = NULL, label = NULL, align = NULL,
       digits = 4, display = NULL, auto = FALSE, ...)
## S3 method for class 'summary_emm'
xtable(x, caption = NULL, label = NULL,
       align = NULL, digits = 4, display = NULL, auto = FALSE, ...)
## S3 method for class 'xtable_emm'
print(x, type = getOption("xtable.type", "latex"),
       include.rownames = FALSE, sanitize.message.function = footnotesize, ...)
```

**Arguments**

- `x` Object of class emmGrid
- `caption` Passed to *xtableList*
- `label` Passed to *xtableList*
- `align` Passed to *xtableList*
- `digits` Passed to *xtableList*
- `display` Passed to *xtableList*
- `auto` Passed to *xtableList*
- `...` Arguments passed to *summary_emmGrid*
- `type` Passed to *print.xtable*
- `include.rownames` Passed to *print.xtable*
- `sanitize.message.function` Passed to *print.xtable*
Details

The methods actually use `xtableList`, because of its ability to display messages such as those for P-value adjustments. These methods return an object of class "xtable_emm" – 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.

Value

The `xtable` methods return an `xtable_emm` object, for which its print method is `print.xtable_emm`.

Examples

```r
pigsint.lm <- lm(log(conc) ~ source * factor(percent), data = pigs)
pigsint.emm <- emmeans(pigsint.lm, ~ percent | source)
xtable::xtable(pigsint.emm, type = "response")
```
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