# Package 'tern.mmrm'

July 4, 2025

```
Title Tables and Graphs for Mixed Models for Repeated Measures (MMRM)
Version 0.3.3
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Description Mixed models for repeated measures (MMRM) are a popular
      choice for analyzing longitudinal continuous outcomes in randomized
      clinical trials and beyond; see for example Cnaan, Laird and Slasor
      (1997)
      <doi:10.1002/(SICI)1097-0258(19971030)16:20%3C2349::AID-SIM667%3E3.0.CO;2-E>.
      This package provides an interface for fitting MMRM within the 'tern'
      <a href="https://cran.r-project.org/package=tern">https://cran.r-project.org/package=tern</a>> framework by Zhu et al.
      (2023) and tabulate results easily using 'rtables'
      <a href="https://cran.r-project.org/package=rtables">https://cran.r-project.org/package=rtables</a>> by Becker et al. (2023).
      It builds on 'mmrm' <a href="https://cran.r-project.org/package=mmrm">https://cran.r-project.org/package=mmrm</a> by
      Sabanés Bové et al. (2023) for the actual MMRM computations.
License Apache License 2.0
URL https://github.com/insightsengineering/tern.mmrm,
      https://insightsengineering.github.io/tern.mmrm/
BugReports https://github.com/insightsengineering/tern.mmrm/issues
Depends R (>= 3.6), tern (>= 0.9.9)
Imports checkmate (>= 2.1.0), cowplot, dplyr (>= 1.1.0), emmeans (>=
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      (>= 0.2.0), magrittr, mmrm (>= 0.3.5), parallelly (>= 1.25.0),
      rlang (>= 1.0.1), rtables (>= 0.6.13), stats, tidyr (>= 0.8.3)
Suggests broom (>= 0.7.10), grid, knitr (>= 1.42), maditr (>= 0.8.1),
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Config/Needs/verdepcheck insightsengineering/tern, mllg/checkmate,
      wilkelab/cowplot, tidyverse/dplyr, rvlenth/emmeans,
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      tidyverse/ggplot2, r-lib/lifecycle, tidyverse/magrittr,
      openpharma/mmrm, HenrikBengtsson/parallelly, r-lib/rlang,
```

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| insightsengineering/rtables, tidyverse/tidyr, tidymodels/broom, yihui/knitr, gdemin/maditr, cran/Matrix, rstudio/rmarkdown, r-lib/testthat, r-lib/vdiffr, r-lib/withr  |
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build\_formula

Building Model Formula

# Description

## [Stable]

This builds the model formula which is used inside fit\_mmrm() and provided to mmrm::mmrm() internally. It can be instructive to look at the resulting formula directly sometimes.

## Usage

```
build_formula(
   vars,
   cor_struct = c("unstructured", "toeplitz", "heterogeneous toeplitz", "ante-dependence",
        "heterogeneous ante-dependence", "auto-regressive", "heterogeneous auto-regressive",
        "compound symmetry", "heterogeneous compound symmetry")
)
```

# Arguments

```
vars
(list)
variables to use in the model.

cor_struct
(string)
specify the covariance structure to use.
```

## Value

Formula to use in mmrm::mmrm().

```
vars <- list(
  response = "AVAL", covariates = c("RACE", "SEX"),
  id = "USUBJID", arm = "ARMCD", visit = "AVISIT"
)
build_formula(vars, "auto-regressive")
build_formula(vars)</pre>
```

```
extract_mmrm_subgroups
```

Extraction of MMRM Subgroup Results based on Population Model Definition

## **Description**

## [Experimental]

This prepares LS mean estimates and contrasts for a specific visit and treatment arm relative to the reference arm, along with a list of subgroup variables and corresponding (grouped) factor levels.

## Usage

```
extract_mmrm_subgroups(
   fit,
   visit,
   subgroups = NULL,
   groups_lists = list(),
   treatment_arm = fit$treatment_levels[1L],
   label_all = "All Patients"
)
```

## **Arguments**

| fit           | (tern_mmrm) model fit on the total population.   |
|---------------|--|
| visit         | (string) single visit or name of averages of visits (referring to the averages specified when creating the fit).                   |
| subgroups     | (character or NULL) names of subgroup variables to use in the forest plot, these need to be factors.                               |
| groups_lists  | (named list of list) optionally contains for each subgroups variable a list, which specifies groups of factor levels, see details. |
| treatment_arm | (string) single treatment arm to compare with the reference arm.   |
| label_all     | (string) label for the total population analysis.  |

## **Details**

The groups\_lists argument is handy when you don't want to have subgroups identical to the original levels of the factor variable. This might be the case when you want to merge levels into a single subgroup, define overlapping subgroups or omit levels completely. Then you insert an element into groups\_lists with the name of the subgroups variable and containing as a named list the subgroup definitions. See the example below.

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## Value

A list with two elements:

• estimates: data. frame with columns arm, n, 1smean, subgroup, var, var\_label, row\_type, containing the LS means results for the overall population and the specified subgroups.

• contrasts: data.frame with columns n\_tot, diff, lcl, ucl, pval, subgroup, var, var\_label, row\_type. Note that this has half the number of rows as estimates.

#### Note

If the original model vars include covariates which are used here in subgroups then these are dropped from covariates before the corresponding model is fitted.

```
mmrm_results <- fit_mmrm(</pre>
 vars = list(
   response = "FEV1",
   covariates = "RACE",
   id = "USUBJID",
   arm = "ARMCD",
   visit = "AVISIT"
 ),
 data = mmrm_test_data,
 cor_struct = "compound symmetry",
 weights_emmeans = "equal",
 averages_emmeans = list(
    "VIS1+2" = c("VIS1", "VIS2")
)
extract_mmrm_subgroups(
 fit = mmrm_results,
 visit = "VIS3",
 subgroups = c("RACE", "SEX"),
 groups_lists = list(
   RACE = list(
     A = c("Asian", "White"),
      B = c("Black or African American", "White")
 )
)
```

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## **Description**

#### [Stable]

Does the MMRM analysis. Multiple other functions can be called on the result to produce tables and graphs.

# Usage

```
fit_mmrm(
  vars = list(response = "AVAL", covariates = c(), id = "USUBJID", arm = "ARM", visit =
       "AVISIT"),
  data,
  conf_level = 0.95,
  cor_struct = "unstructured",
  weights_emmeans = "proportional",
  averages_emmeans = list(),
  parallel = FALSE,
  ...
)
```

#### **Arguments**

vars

(named list of string or character)

specifying the variables in the MMRM. The following elements need to be included as character vectors and match corresponding columns in data:

- response: the response variable.
- covariates: the additional covariate terms (might also include interactions).
- id: the subject ID variable.
- arm: the treatment group variable (factor).
- visit: the visit variable (factor).
- weights: optional weights variable (if NULL or omitted then no weights will be used).

Note that the main effects and interaction of arm and visit are by default included in the model.

data

(data.frame)

with all the variables specified in vars. Records with missing values in any independent variables will be excluded.

conf\_level

(proportion)

confidence level of the interval.

cor\_struct

(string)

specifying the covariance structure, defaults to "unstructured". See the details.

weights\_emmeans

(string)

argument from emmeans::emmeans(), "proportional" by default.

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averages\_emmeans

(list)

optional named list of visit levels which should be averaged and reported along side the single visits.

parallel (flag)

controls whether the optimizer search can use available free cores on the ma-

chine (not default).

... additional arguments for mmrm::mmrm(), in particular reml and options listed in

mmrm::mmrm\_control().

#### **Details**

Multiple different degree of freedom adjustments are available via the method argument for mmrm::mmrm(). In addition, covariance matrix adjustments are available via vcov. Please see mmrm::mmrm\_control() for details and additional useful options.

For the covariance structure (cor\_struct), the user can choose among the following options.

- unstructured: Unstructured covariance matrix. This is the most flexible choice and default. If there are T visits, then T \* (T+1) / 2 variance parameters are used.
- toeplitz: Homogeneous Toeplitz covariance matrix, which uses T variance parameters.
- heterogeneous toeplitz: Heterogeneous Toeplitz covariance matrix, which uses 2 \* T 1 variance parameters.
- ante-dependence: Homogeneous Ante-Dependence covariance matrix, which uses T variance parameters.
- heterogeneous ante-dependence: Heterogeneous Ante-Dependence covariance matrix, which uses 2 \* T 1 variance parameters.
- auto-regressive: Homogeneous Auto-Regressive (order 1) covariance matrix, which uses 2 variance parameters.
- heterogeneous auto-regressive: Heterogeneous Auto-Regressive (order 1) covariance matrix, which uses T + 1 variance parameters.
- compound symmetry: Homogeneous Compound Symmetry covariance matrix, which uses 2 variance parameters.
- heterogeneous compound symmetry: Heterogeneous Compound Symmetry covariance matrix, which uses T + 1 variance parameters.

#### Value

A tern\_mmrm object which is a list with MMRM results:

- fit: The mmrm object which was fitted to the data. Note that via mmrm::component(fit, "optimizer") the finally used optimization algorithm can be obtained, which can be useful for refitting the model later on.
- cov\_estimate: The matrix with the covariance matrix estimate.
- diagnostics: A list with model diagnostic statistics (REML criterion, AIC, corrected AIC, BIC).
- Ismeans: This is a list with data frames estimates and contrasts. The attributes averages and weights save the settings used (averages\_emmeans and weights\_emmeans).

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- vars: The variable list.
- labels: Corresponding list with variable labels extracted from data.
- cor\_struct: input.
- parallel: input.
- ref\_level: The reference level for the arm variable, which is always the first level.
- treatment\_levels: The treatment levels for the arm variable.
- conf\_level: The confidence level which was used to construct the 1smeans confidence intervals.
- additional: List with any additional inputs passed via . . .

## **Examples**

```
library(dplyr)
library(rtables)
mmrm_results <- fit_mmrm(</pre>
  vars = list(
    response = "FEV1",
    covariates = c("RACE", "SEX"),
    id = "USUBJID",
    arm = "ARMCD",
    visit = "AVISIT"
  ),
  data = mmrm_test_data,
  cor_struct = "unstructured",
  weights_emmeans = "equal",
  averages_emmeans = list(
    "VIS1+2" = c("VIS1", "VIS2")
)
```

 ${\tt get\_mmrm\_lsmeans}$ 

Extract Least Square Means from MMRM

# Description

## [Stable]

Extracts the least square means from an MMRM fit.

## Usage

```
get_mmrm_lsmeans(fit, vars, conf_level, weights, averages = list())
```

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## **Arguments**

fit (mmrm)

result of mmrm::mmrm().

vars

(named list of string or character)

specifying the variables in the MMRM. The following elements need to be included as character vectors and match corresponding columns in data:

- response: the response variable.
- covariates: the additional covariate terms (might also include interactions).
- id: the subject ID variable.
- arm: the treatment group variable (factor).
- visit: the visit variable (factor).
- weights: optional weights variable (if NULL or omitted then no weights will be used).

Note that the main effects and interaction of arm and visit are by default included in the model.

conf\_level (proportion)

confidence level of the interval.

weights (string)

type of weights to be used for the least square means, see emmeans::emmeans()

for details.

averages (list)

named list of visit levels which should be averaged and reported along side the

single visits.

#### Value

A list with data frames estimates and contrasts. The attributes averages and weights save the settings used.

g\_covariance

Visualization of Covariance Matrix

## **Description**

## [Experimental]

Plot of covariance (or correlation) matrix as a function of lag or time. The covariance structure is vectorized internally and lag or time distances are computed and can be used for visualization.

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## Usage

```
g_covariance(
  vcov_matrix,
  time_prefix = NULL,
  x_var = c("lag", "time_diff"),
  xlab = NULL,
  ylab = ""
)
```

## **Arguments**

```
vcov_matrix (matrix)
symmetric covariance matrix with identical row and column names.

time_prefix (string)
string in the names of vcov_matrix that precedes the time point value.

x_var (string)
can be lag or time_diff for lag or time difference, respectively.

xlab (string or NULL)
x-axis label, if NULL then automatically determined from x_var.

ylab (string)
y-axis label.
```

## **Details**

The default time\_prefix value is NULL, which assumes that the names of the input matrix don't have any character string other than time point value. If a time\_prefix is specified, this string should appear in front of all the names in vcov\_matrix.

## Value

The ggplot object.

```
vcov_matrix <- matrix(
    c(49, 12, 12, 23),
    nrow = 2, ncol = 2,
    dimnames = list(
        c(1, 2),
        c(1, 2)
    )
)
g_covariance(vcov_matrix, x_var = "time_diff")</pre>
```

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g\_mmrm\_diagnostic

Diagnostic Plots for MMRM

## **Description**

## [Stable]

This function produces diagnostic plots.

## Usage

```
g_mmrm_diagnostic(
  object,
  type = c("fit-residual", "q-q-residual"),
  z_threshold = NULL
)
```

## **Arguments**

object (tern\_mmrm)

model result produced by fit\_mmrm().

type (string)

specifying the type of diagnostic plot to be produced:

- fit-residual: A fitted vs residuals plot, grouped by visits. This allows to see if there is remaining structure in the residuals that might be captured by adding additional covariates to the model.
- q-q-residual: A Q-Q plot for the residuals (i.e. sorted standardized residuals vs. normal quantiles), grouped by visits. Observations with an absolute standardized residual above z\_threshold will be labeled.

z\_threshold (numeric)

optional number indicating the normal quantile threshold for the Q-Q plot.

## **Details**

Here we use marginal fitted values and residuals. That is, we estimate fitted values, and the difference of those fitted values vs. the observed data are the residuals.

# Value

A ggplot2 plot.

#### See Also

g\_mmrm\_lsmeans() for plotting the least-squares means and contrasts.

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## **Examples**

```
mmrm_results <- fit_mmrm(
  vars = list(
    response = "FEV1",
    covariates = c("RACE", "SEX"),
    id = "USUBJID",
    arm = "ARMCD",
    visit = "AVISIT"
  ),
  data = mmrm_test_data,
    cor_struct = "unstructured",
    weights_emmeans = "equal"
)
g_mmrm_diagnostic(mmrm_results)
g_mmrm_diagnostic(mmrm_results, type = "q-q-residual")</pre>
```

g\_mmrm\_lsmeans

Plot LS means for MMRM

## Description

### [Stable]

This function summarizes adjusted 1smeans and standard error, as well as conducts comparisons between groups' adjusted 1smeans, where the first level of the group is the reference level.

## Usage

```
g_mmrm_lsmeans(
  object,
  select = c("estimates", "contrasts"),
  titles = c(estimates = paste("Adjusted mean of", object$labels$response,
    "by treatment at visits"), contrasts = paste0("Differences of ",
  object$labels$response, "adjusted means vs. control ('", object$ref_level, "')")),
  xlab = object$labels$visit,
 ylab = paste0("Estimates with ", round(object$conf_level * 100), "% CIs"),
  xlimits = NULL,
 ylimits = NULL,
 width = 0.6,
  show_pval = TRUE,
  show_lines = FALSE,
  constant_baseline = NULL,
  n_baseline = NA_integer_,
  table_stats = character(),
 table_formats = c(n = "xx.", estimate = "xx.x", se = "xx.x", ci = "(xx.xx, xx.xx)"),
  table_labels = c(n = "n", estimate = "LS mean", se = "Std. Error", ci =
    paste0(round(object$conf_level * 100), "% CI")),
  table_font_size = 3,
  table_rel_height = 0.5
)
```

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#### **Arguments**

object (tern\_mmrm)

model result produced by fit\_mmrm().

select (character)

to select one or both of "estimates" and "contrasts" plots. Note "contrasts" option

is not applicable to model summaries excluding an arm variable.

titles (character)

with elements estimates and contrasts containing the plot titles.

xlab (string)

the x axis label.

ylab (string)

the y axis label.

xlimits (numeric)

x axis limits.

ylimits (numeric)

y axis limits.

width (numeric)

width of the error bars.

show\_pval (flag)

should the p-values for the differences of LS means vs. control be displayed (not

default)?

show\_lines (flag)

should the visit estimates be connected by lines (not default)?

constant\_baseline

(named number or NULL)

optional constant baseline for the LS mean estimates. If specified then needs to be a named number, and the name will be used to label the corresponding baseline visit. The differences of LS means will always be 0 at this baseline

visit.

n\_baseline (int or named integer)

optional number of patients at baseline. Since this can be visible in the optional table below the estimates plot, and we cannot infer this from object (since that is then only fit without baseline data), we need to allow the user to specify this. If number then assumed constant across potential treatment arms, otherwise one

number per treatment arm can be provided.

table\_stats (character)

names of the statistics that will be displayed in the table below the estimates plot. Note that the table will only be added when selecting only the "estimates"

plot. Available statistics are n, estimate, se, and ci.

table\_formats (named character)

format patterns for descriptive statistics used in the (optional) estimates table

appended to the estimates plot.

table\_labels (named character)

labels for the statistics in the (optional) estimates table.

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#### **Details**

If variable labels are available in the original data set, then these are used. Otherwise the variable names themselves are used for annotating the plot.

The contrast plot is not going to be returned if treatment is not considered in the tern\_mmrm object input, no matter if select argument contains the contrasts value.

## Value

A ggplot2 plot.

```
library(dplyr)
mmrm_results <- fit_mmrm(</pre>
  vars = list(
   response = "FEV1",
   covariates = c("RACE", "SEX"),
   id = "USUBJID",
   arm = "ARMCD",
   visit = "AVISIT"
  data = mmrm_test_data,
  cor_struct = "unstructured",
  weights_emmeans = "equal"
g_mmrm_lsmeans(mmrm_results, constant_baseline = c(BSL = 0))
g_mmrm_lsmeans(
  mmrm_results,
  select = "estimates",
  show_lines = TRUE,
  xlab = "Visit"
)
g_mmrm_lsmeans(
  mmrm_results,
  select = "contrasts",
  titles = c(contrasts = "Contrasts of FKSI-FWB means"),
  show_pval = TRUE,
  show_lines = TRUE,
  width = 0.8
)
```

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```
mmrm_test_data2 <- mmrm_test_data %>%
  filter(ARMCD == "TRT")
mmrm_results_no_arm <- fit_mmrm(</pre>
  vars = list(
   response = "FEV1",
   covariates = c("RACE", "SEX"),
   id = "USUBJID",
   visit = "AVISIT"
  ),
  data = mmrm_test_data2,
  cor_struct = "unstructured",
  weights_emmeans = "equal"
g_mmrm_lsmeans(mmrm_results_no_arm, select = "estimates")
g_mmrm_lsmeans(
  mmrm_results_no_arm,
  select = c("estimates", "contrasts"),
  titles = c(
   estimates = "Adjusted mean of FKSI-FWB",
   contrasts = "it will not be created"
  ),
  show_pval = TRUE,
  width = 0.8
)
g_mmrm_lsmeans(
  mmrm_results_no_arm,
  select = "estimates",
  titles = c(estimates = "Adjusted mean of FKSI-FWB"),
  show_pval = TRUE,
  width = 0.8,
  show\_lines = TRUE
g_mmrm_lsmeans(
  mmrm_results,
  select = "estimates",
  titles = c(estimates = "Adjusted mean of FKSI-FWB"),
  table_stats = c("n", "ci")
)
```

mmrm\_test\_data

Example dataset for tern.mmrm package.

## **Description**

## [Stable]

Measurements of FEV1 (forced expired volume in one second) is a measure of how quickly the lungs can be emptied. Low levels of FEV1 may indicate chronic obstructive pulmonary disease (COPD).

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## Usage

```
mmrm_test_data
```

#### **Format**

A tibble with 800 rows and 7 variables:

- USUBJID: unique subject identifier.
- AVISIT: visit number.
- ARMCD: treatment, TRT or PBO.
- RACE: 3-category race.
- SEX: sex.
- FEV1\_BL: FEV1 at baseline (%).
- FEV1: FEV1 at study visits.

tabulate\_mmrm

Tabulation of MMRM Results

## **Description**

## [Stable]

These functions can be used to produce tables from a fitted MMRM produced with fit\_mmrm().

# Usage

```
## S3 method for class 'tern_mmrm'
as.rtable(x, type = c("fixed", "cov", "diagnostic"), ...)
h_mmrm_fixed(x, format = "xx.xxxxx")
h_mmrm_cov(x, format = "xx.xxxxx")
h_mmrm_diagnostic(x, format = "xx.xxxxx")

## S3 method for class 'tern_mmrm'
tidy(x, ...)
s_mmrm_lsmeans(df, .in_ref_col, show_relative = c("reduction", "increase"))
a_mmrm_lsmeans(df, .in_ref_col, show_relative = c("reduction", "increase"))
s_mmrm_lsmeans_single(df)
a_mmrm_lsmeans_single(df)
```

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```
summarize_lsmeans(
   lyt,
   arms = TRUE,
   ...,
   table_names = "lsmeans_summary",
   .stats = NULL,
   .formats = NULL,
   .indent_mods = NULL,
   .labels = NULL
)
```

## **Arguments**

x (tern\_mmrm)

the original result from fit\_mmrm().

type (string)

type of table which should be returned.

additional argument format for controlling the numeric format.

format (string)

format for the numbers in the table.

df (data frame)

data set containing all analysis variables.

.in\_ref\_col (logical)

TRUE when working with the reference level, FALSE otherwise.

show\_relative should the "reduction" (control - treatment, default) or the "increase" (treatment

- control) be shown for the relative change from baseline?

lyt (layout)

input layout where analyses will be added to.

arms (flag)

should treatment variable be considered when using summarize\_lsmeans layout

generating function.

table\_names (character)

this can be customized in case that the same vars are analyzed multiple times,

to avoid warnings from rtables.

.stats (character)

statistics to select for the table.

.formats (named character or list)

formats for the statistics.

.indent\_mods (named integer)

indent modifiers for the labels.

.labels (named character)

labels for the statistics (without indent).

## Value

as.rtable.tern\_mmrm() returns the fixed effects, covariance estimate or diagnostic statistics tables.

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#### **Functions**

- as.rtable(tern\_mmrm): Produce simple MMRM tables via the generic as.rtable().
- h\_mmrm\_fixed(): Helper function to produce fixed effects table.
- h\_mmrm\_cov(): Helper function to produce a covariance matrix table.
- h\_mmrm\_diagnostic(): Helper function to produce a diagnostic statistics table.
- tidy(tern\_mmrm): Helper method (for broom::tidy()) to prepare a data.frame from an tern\_mmrm object containing the least-squares means and contrasts.
- s\_mmrm\_lsmeans(): Statistics function which is extracting estimates from a tidied least-squares means data frame.
- a\_mmrm\_lsmeans(): Formatted Analysis function which can be further customized by calling rtables::make\_afun() on it. It is used as afun in rtables::analyze().
- s\_mmrm\_lsmeans\_single(): Statistics function which is extracting estimates from a tidied least-squares means data frame when ARM is not considered in the model.
- a\_mmrm\_lsmeans\_single(): Formatted Analysis function (when ARM is not considered in the model) which can be further customized by calling rtables::make\_afun() on it. It is used as afun in rtables::analyze().
- summarize\_1smeans(): Analyze function for tabulating least-squares means estimates from tidied mmrm results.

```
result <- fit_mmrm(</pre>
 vars = list(
   response = "FEV1",
   covariates = c("RACE", "SEX"),
   id = "USUBJID",
   arm = "ARMCD",
    visit = "AVISIT"
 ),
 data = mmrm_test_data,
 cor_struct = "unstructured",
 weights_emmeans = "equal"
)
as.rtable(result, type = "cov", format = "xx.x")
result_no_arm <- fit_mmrm(</pre>
 vars = list(
    response = "FEV1",
   covariates = c("RACE", "SEX"),
   id = "USUBJID",
   visit = "AVISIT"
 data = mmrm_test_data,
 cor_struct = "unstructured",
 weights_emmeans = "equal"
)
as.rtable(result_no_arm, type = "cov", format = "xx.x")
df <- broom::tidy(result)</pre>
```

```
df_no_arm <- broom::tidy(result_no_arm)</pre>
s_mmrm_lsmeans(df[8, ], .in_ref_col = FALSE)
s_mmrm_lsmeans_single(df_no_arm[4, ])
library(dplyr)
dat_adsl <- mmrm_test_data %>%
 select(USUBJID, ARMCD) %>%
 unique()
basic_table() %>%
 split_cols_by("ARMCD", ref_group = result$ref_level) %>%
 add_colcounts() %>%
 split_rows_by("AVISIT") %>%
 summarize_lsmeans(
    .stats = c("n", "adj_mean_se", "adj_mean_ci", "diff_mean_se", "diff_mean_ci"),
    .labels = c(adj_mean_se = "Adj. LS Mean (Std. Error)"),
    .formats = c(adj_mean_se = sprintf_format("%.1f (%.2f)"))
 ) %>%
 build_table(
   df = broom::tidy(result),
   alt_counts_df = dat_adsl
basic_table() %>%
 split_rows_by("AVISIT") %>%
 summarize_lsmeans(arms = FALSE) %>%
 build_table(
   df = broom::tidy(result_no_arm),
   alt_counts_df = dat_adsl
```

tabulate\_mmrm\_subgroups

Tabulation of MMRM Subgroups Results

## Description

## [Experimental]

This function tabulates the results from extract\_mmrm\_subgroups().

## Usage

## **Arguments**

lyt (layout) input layout where analyses will be added to. df of data frames containing all analysis variables, is the result from extract\_mmrm\_subgroups(). (character) vars the name of statistics to be reported among n\_tot (total number of patients per group), n (number of patients per treatment arm and group), 1smean (least square mean point estimate), diff (difference of least square mean estimates between treatment and reference arm), ci (confidence interval of difference) and pval (p value of the diff, not adjusted for multiple comparisons). Note, the statistics n\_tot, diff and ci are required. .formats (named list) containing the formats for the statistics. .labels (named list) containing the labels for the statistics.

#### Value

The rtables object.

```
mmrm_results <- fit_mmrm(</pre>
  vars = list(
    response = "FEV1",
    covariates = "RACE",
    id = "USUBJID",
    arm = "ARMCD",
    visit = "AVISIT"
  ),
  data = mmrm_test_data,
  cor_struct = "compound symmetry",
  weights_emmeans = "equal",
  averages_emmeans = list(
    "VIS1+2" = c("VIS1", "VIS2")
  )
)
df <- extract_mmrm_subgroups(</pre>
  fit = mmrm_results,
  visit = "VIS3",
  subgroups = c("RACE", "SEX")
## Table with default columns.
basic_table() %>%
  tabulate_mmrm_subgroups(df)
```

```
## Table with selected columns.
tab <- basic_table() %>%
  tabulate_mmrm_subgroups(
    df = df,
    vars = c("n_tot", "diff", "ci", "pval")
  )
tab

## Forest plot can be produced based on this very easily.
g_forest(tab, logx = FALSE, xlim = c(-10, 10), x_at = c(-10, -5, 0, 5, 10), vline = 0)
```

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