

Package ‘outstandR’

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Title Model-Based Standardisation for Indirect Treatment Comparison with Limited Subject-Level Data

Version 2.0.0

Description For the problem of indirect treatment comparison with limited subject-level data, this package provides tools for model-based standardisation with several different computation approaches.

See Remiro-Azócar A, Heath A, Baio G (2022) “Parametric G-computation for compatible indirect treatment comparisons with limited individual patient data”, Res. Synth. Methods, 1–31. ISSN 1759-2879, <[doi:10.1002/jrsm.1565](https://doi.org/10.1002/jrsm.1565)>.

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AC_IPD_binY_contX	<i>Individual-level patient data for binary outcome, continuous covariates</i>
-------------------	--

Description

This data set contains simulated patient covariate and outcome values.

Usage

```
data(AC_IPD_binY_contX)
```

Format

```
y ~ PF_cont_1 + PF_cont_2 + trt + trt:(EM_cont_1 + EM_cont_2):
```

id Numeric unique identifier

PF_cont_1 Numeric prognostic factor continuous covariate

PF_cont_2 Numeric prognostic factor continuous covariate

EM_cont_1 Numeric effect modifier continuous covariate

EM_cont_2 Numeric effect modifier continuous covariate

trt Factor treatment identifier. Levels A, C

y Integer binary outcome

true_eta Numeric linear predictor

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

AC_IPD_contY_mixedX	<i>Individual-level patient data for continuous outcome, mixed covariates</i>
---------------------	---

Description

This data set contains simulated patient covariate and outcome values. Corresponds to ALD data set.

Usage

```
data(AC_IPD_contY_mixedX)
```

Format

$y \sim X1 + X3 + X4 + \text{trt} + \text{trt}:(X2 + X3 + X4):$

id Numeric unique identifier

X1 Numeric prognostic factor continuous covariate

X2 Numeric prognostic factor and effect modifier binary covariate

X3 Numeric prognostic factor and effect modifier continuous covariate

X4 Numeric effect modifier binary covariate

trt Factor treatment identifier. Levels A, C

y Integer binary outcome

true_eta Numeric linear predictor

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

AC_IPD_countY_contX *Individual-level patient data for count outcome, continuous covariates*

Description

This data set contains simulated patient covariate and outcome values. Corresponds to ALD data set.

Usage

`data(AC_IPD_countY_contX)`

Format

$y \sim \text{PF_cont_1} + \text{PF_cont_2} + \text{trt} + \text{trt}:(\text{EM_cont_1} + \text{EM_cont_2}):$

id Numeric unique identifier

PF_cont_1 Numeric prognostic factor continuous covariate

PF_cont_2 Numeric prognostic factor continuous covariate

EM_cont_1 Numeric effect modifier continuous covariate

EM_cont_2 Numeric effect modifier continuous covariate

trt Factor treatment identifier. Levels A, C

y Integer non-negative count outcome

true_eta Numeric linear predictor

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

BC_ALD_binY_contX	<i>Aggregate level patient data for binary outcome, continuous covariates</i>
-------------------	---

Description

This data set contains summaries of simulated patient covariate and outcome values.

Usage

```
data(BC_ALD_binY_contX)
```

Format

```
y ~ PF_cont_1 + PF_cont_2 + trt + trt:(EM_cont_1 + EM_cont_2):
```

variable String covariate or outcome name. From EM_cont_1, EM_cont_2, PF_cont_1, PF_cont_2, y.

statistic String summary statistic name. From mean, sd, sum, N

value Numeric value

trt Treatment (arm) name. From B, C

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

BC_ALD_contY_mixedX *Aggregate level patient data for continuous outcome, mixed covariates*

Description

This data set contains summaries of simulated patient covariate and outcome values. Corresponds to IPD data set.

Usage

```
data(BC_ALD_contY_mixedX)
```

Format

$y \sim X1 + X3 + X4 + \text{trt} + \text{trt}:(X2 + X3 + X4)$:

variable String covariate or outcome name. From X1, X2, X3, X4, y.

statistic String summary statistic name. From mean, sd, prob, sum, N

value Numeric value

trt Treatment (arm) name. From B, C

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

BC_ALD_countY_contX *Aggregate level patient data for count outcome, continuous covariates*

Description

This data set contains summaries of simulated patient covariate and outcome values. Corresponds to IPD data set.

Usage

```
data(BC_ALD_countY_contX)
```

Format

$y \sim \text{PF_cont_1} + \text{PF_cont_2} + \text{trt} + \text{trt}:(\text{EM_cont_1} + \text{EM_cont_2}):$

variable String covariate or outcome name. From EM_cont_1, EM_cont_2, PF_cont_1, PF_cont_2, y.

statistic String summary statistic name. From mean, sd, sum, N

value Numeric value

trt Treatment (arm) name. From B, C

Source

Simulated data

References

Remiro-Azocar A, Heath A, Baio G (2022)

calculate_ate	<i>Calculate Average Treatment Effect</i>
---------------	---

Description

Computes the average treatment effect (ATE) based on the specified effect scale.

Usage

```
calculate_ate(mean_comp, mean_ref, effect)
```

Arguments

mean_comp, mean_ref

Mean of the outcome for the comparator and reference / common

effect

A character string specifying the effect scale. Options are:

"log_odds" Log-odds difference.

"risk_difference" Risk difference.

"delta_z" Probit scale difference (z-scores).

"log_relative_risk_rare_events" Log relative risk for rare events.

"log_relative_risk" Log relative risk.

Value

Numeric computed average treatment effect on the specified scale.

Examples

```
calculate_ate(mean_comp = 0.7, mean_ref = 0.5, effect = "log_odds")
calculate_ate(mean_comp = 0.7, mean_ref = 0.5, effect = "risk_difference")
```

calculate_trial_mean *Calculate Trial Mean Wrapper*

Description

Calculate Trial Mean Wrapper

Usage

```
calculate_trial_mean(ald, tid, effect, family)
```

Arguments

ald	Aggregate level data. Data frame in long format.
tid	Treatment ID
effect	Effect name. String.
family	Family distribution

Value

Numeric mean value.

calculate_trial_mean_binary
Calculate Trial Mean Binary Data

Description

Calculate Trial Mean Binary Data

Usage

```
calculate_trial_mean_binary(ald, tid, effect)
```

Arguments

ald	Aggregate level data. Data frame in long format.
tid	Treatment ID
effect	Effect name. String.

Value

Numeric mean value.

`calculate_trial_mean_continuous`*Calculate Trial Mean Continuous Data*

Description

Calculate Trial Mean Continuous Data

Usage

```
calculate_trial_mean_continuous(ald, tid, effect, verbatim = FALSE)
```

Arguments

<code>ald</code>	Aggregate level data. Data frame in long format.
<code>tid</code>	Treatment ID
<code>effect</code>	Effect name. String.
<code>verbatim</code>	Print messages, logical

Value

Numeric mean value.

`calculate_trial_mean_count`*Calculate Trial Mean Count Data*

Description

Calculate Trial Mean Count Data

Usage

```
calculate_trial_mean_count(ald, tid, effect, verbatim = FALSE)
```

Arguments

<code>ald</code>	Aggregate level data. Data frame in long format.
<code>tid</code>	Treatment ID
<code>effect</code>	Effect name. String.
<code>verbatim</code>	Print messages, logical

Value

Numeric mean value.

 calculate_trial_variance

Calculate trial variance

Description

Computes the variance of treatment effects for a trial based on the specified family distribution.

Usage

```
calculate_trial_variance(ald, tid, effect, family)
```

Arguments

ald	Aggregate-level data. Data frame.
tid	Treatment identifier used to extract relevant columns from ald.
effect	A character string specifying the effect scale (e.g., "log_odds", "risk_difference").
family	A character string specifying the model family (e.g., "binomial", "gaussian").

Value

Numeric computed variance of treatment effects.

Examples

```
ald <- data.frame(trt = c("B", "C", "B", "C"),
                 variable = c(NA, NA, "y", "y"),
                 statistic = c("N", "N", "sum", "sum"),
                 value = c(100, 100, 50, 60))

calculate_trial_variance(
  ald, tid = "B", effect = "log_odds", family = "binomial")
```

 calculate_trial_variance_binary

Calculate trial variance binary

Description

Calculate trial variance binary

Usage

```
calculate_trial_variance_binary(ald, tid, effect)
```

Arguments

ald	Aggregate level data
tid	Treatment ID
effect	Effect

Value

Numeric value of total variance.

calculate_trial_variance_continuous
Calculate trial variance continuous

Description

Calculate trial variance continuous

Usage

```
calculate_trial_variance_continuous(ald, tid, effect, verbatim = FALSE)
```

Arguments

ald	Aggregate level data. Data frame in long format.
tid	Treatment ID
effect	Effect name. String.
verbatim	Print messages, logical

Value

Numeric value of total variance.

calculate_trial_variance_count
Calculate trial variance count

Description

Calculate trial variance count

Usage

```
calculate_trial_variance_count(ald, tid, effect)
```

Arguments

ald	Aggregate level data. Data frame in long format.
tid	Treatment ID.
effect	Effect name. String.

Value

Numeric value of total variance.

calc_ALD_stats	<i>Aggregate-level data mean and variance statistics</i>
----------------	--

Description

Computes the mean and variance of marginal treatment effects for aggregate-level trial data.

Usage

```
calc_ALD_stats(strategy, analysis_params)
```

Arguments

strategy	A list containing the strategy details, including the family distribution.
analysis_params	A list containing: <ul style="list-style-type: none"> • ald Aggregate-level trial data • ref_trt Treatment labels reference (common; e.g. placebo) • ald_comp Treatment labels comparator • scale A scaling parameter for the calculation. From "log_odds", "risk_difference", "log_relative_risk".

Value

A list containing:

mean	The marginal treatment effect mean.
var	The marginal treatment effect variance.

See Also

[marginal_treatment_effect\(\)](#) [marginal_variance\(\)](#)

Examples

```
strategy <- list(family = list(family = "binomial")) # basic version

ald <- data.frame(trt = c("B","C","B","C"),
                 variable = c(NA, NA, "y", "y"),
                 statistic = c("N", "N", "sum", "sum"),
                 value = c(100, 100, 50, 60))

calc_ALD_stats(strategy = strategy,
               list(ald = ald,
                   ref_trt = "C",
                   ald_comp = "B",
                   scale = "log_odds"))
```

calc_gcomp_bayes

*Bayesian G-computation using Stan***Description**

Calculate draws of binary responses from posterior predictive distribution from the Bayesian G-computation method using Hamiltonian Monte Carlo.

Usage

```
calc_gcomp_bayes(strategy, analysis_params, ...)
```

Arguments

strategy	A list specifying the model strategy, including: <ul style="list-style-type: none"> • outcome_model: A linear regression formula object. • family: A family object specifying the distribution and link function (e.g., binomial). • iter: Number of iterations for the MCMC sampling. • warmup: Number of warmup iterations for the MCMC sampling. • chains: Number of MCMC chains.
analysis_params	List of analysis parameters. Must contain ipd and ald.
...	Additional arguments passed to <code>rstanarm::stan_glm()</code> .

Value

A list containing:

- means: A list containing:
 - Posterior means for comparator treatment group.
 - Posterior means for reference treatment group.
- model: A list containing the fit object (from `stan_glm`), rho, N, and stan_args.

Examples

```

strategy <- list(
  outcome_model = y ~ trt:X1,
  family = binomial(),
  rho = NA,
  N = 1000L,
  marginal_distns = NA,
  marginal_params = NA,
  trt_var = "trt",
  iter = 2000,
  warmup = 500,
  chains = 4)

ipd <- data.frame(
  trt = sample(c("A", "C"), size = 100, replace = TRUE),
  X1 = rnorm(100, 1, 1),
  y = sample(c(1,0), size = 100, prob = c(0.7, 0.3), replace = TRUE))

ald <- data.frame(
  trt = c(NA, NA, "B", "C", "B", "C"),
  variable = c("X1", "X1", "y", "y", NA, NA),
  statistic = c("mean", "sd", "sum", "sum", "N", "N"),
  value = c(0.5, 0.1, 10, 12, 20, 25))

res <-
  calc_gcomp_bayes(
    strategy,
    analysis_params = list(
      ipd = ipd, ald = ald,
      ref_trt = "C",
      ipd_comp = "A"))

str(res, max.level = 2, list.len = 3, vec.len = 2)

```

calc_gcomp_ml

G-computation Maximum Likelihood Bootstrap

Description

Computes the mean difference in treatment effects using bootstrap resampling.

Usage

```
calc_gcomp_ml(strategy, analysis_params)
```

Arguments

strategy A list specifying the model strategy, including:

- R: Number of bootstrap replications.
- outcome_model: A linear regression formula object for the outcome model.
- family: A family object specifying the distribution and link function (e.g., binomial).
- N: Synthetic sample size for g-computation.

analysis_params

List of analysis parameters.

Value

A list containing:

- means: A list containing:
 - A: Bootstrap estimates for comparator treatment group "A".
 - C: Bootstrap estimates for reference treatment group "C".
- model: A list containing the fit object, rho, and N.

Examples

```
strategy <- list(
  outcome_model = y ~ trt:X1,
  family = binomial(),
  rho = NA,
  N = 1000L,
  n_boot = 100L,
  marginal_distns = NA,
  marginal_params = NA,
  trt_var = "trt")

ipd <- data.frame(trt = sample(c("A", "C"), size = 100, replace = TRUE),
  X1 = rnorm(100, 1, 1),
  y = sample(c(1,0), size = 100, prob = c(0.7,0.3), replace = TRUE))

ald <- data.frame(trt = c(NA, NA, "B", "C", "B", "C"),
  variable = c("X1", "X1", "y", "y", NA, NA),
  statistic = c("mean", "sd", "sum", "sum", "N", "N"),
  value = c(0.5, 0.1, 10, 12, 20, 25))

calc_gcomp_ml(
  strategy,
  analysis_params =
    list(ipd = ipd, ald = ald,
      ref_trt = "C",
      ipd_comp = "A"))
```

calc_IPD_stats	<i>Calculate individual-level patient data statistics</i>
----------------	---

Description

Computes mean and variance statistics for individual-level patient data using various approaches, including Matching-Adjusted Indirect Comparison (MAIC), Simulated Treatment Comparison (STC), and G-computation via Maximum Likelihood Estimation (MLE) or Bayesian inference.

Usage

```
calc_IPD_stats(strategy, analysis_params, ...)

## Default S3 method:
calc_IPD_stats(...)

## S3 method for class 'stc'
calc_IPD_stats(strategy, analysis_params, var_method = NULL, ...)

## S3 method for class 'maic'
calc_IPD_stats(strategy, analysis_params, var_method = NULL, ...)

## S3 method for class 'gcomp_ml'
calc_IPD_stats(strategy, analysis_params, var_method = NULL, ...)

## S3 method for class 'gcomp_bayes'
calc_IPD_stats(strategy, analysis_params, var_method = NULL, ...)

## S3 method for class 'mim'
calc_IPD_stats(strategy, analysis_params, var_method = NULL, ...)
```

Arguments

strategy	A list corresponding to different modelling approaches
analysis_params	A list containing: <ul style="list-style-type: none"> • ipd: Individual-level patient data (data frame) • ald: Aggregate-level trial data (data frame) • ref_trt: Treatment label for the reference arm (common; e.g., "C") • ipd_comp: Treatment label for the comparator arm in the IPD (e.g., "A") • scale: Scaling parameter ("log_odds", "risk_difference", "log_relative_risk")
...	Additional arguments
var_method	A string specifying the variance estimation method, either "sample" (default) or "sandwich".

Value

A list containing:

- contrasts: A list with elements mean and var.
- absolute: A list with elements mean and var.

Simulated treatment comparison statistics

IPD for reference "C" and comparator "A" trial arms are used to fit a regression model describing the observed outcomes y in terms of the relevant baseline characteristics x and the treatment variable z .

Matching-adjusted indirect comparison statistics

Marginal IPD comparator treatment "A" vs reference treatment "C" treatment effect estimates using bootstrapping sampling.

G-computation maximum likelihood statistics

Compute a non-parametric bootstrap with default $R = 1000$ resamples.

G-computation Bayesian statistics

Using Stan, compute marginal relative effects for IPD comparator "A" vs reference "C" treatment arms for each MCMC sample by transforming from probability to linear predictor scale.

Multiple imputation marginalisation

Using Stan, compute marginal relative treatment effect for IPD comparator "A" vs reference "C" arms for each MCMC sample by transforming from probability to linear predictor scale. Approximate by using imputation and combining estimates using pooling.

Examples

```
strategy <- strategy_maic(
  formula = list(outcome_model = y ~ trt, # default
                 balance_model = ~ X1),
  family = binomial())

ipd <- data.frame(trt = sample(c("A", "C"), size = 100, replace = TRUE),
                 X1 = rnorm(100, 1, 1),
                 y = sample(c(1,0), size = 100, prob = c(0.7,0.3), replace = TRUE))

ald <- data.frame(trt = c(NA, "B", "C", "B", "C"),
                 variable = c("X1", "y", "y", NA, NA),
                 statistic = c("mean", "sum", "sum", "N", "N"),
                 value = c(0.5, 10, 12, 20, 25))

calc_IPD_stats(strategy,
  analysis_params = list(ipd = ipd, ald = ald, scale = "log_odds"))
```

estimate_var_sandwich *Estimate Variance Sandwich Estimator*

Description

Computes the robust (sandwich) variance estimator for the treatment effect.

Usage

```
estimate_var_sandwich(strategy, analysis_params, ...)
```

Arguments

strategy	An object of class <code>strategy</code> created by functions such as <code>strategy_maic()</code> , <code>strategy_stc()</code> , or <code>strategy_mim()</code> . Contains modelling details like the formula and family.
analysis_params	List of analysis parameters (ipd, ald, etc.)
...	Additional arguments

Value

Numeric variance estimate for the treatment contrast

get_treatment_effect *Get treatment effect scale corresponding to a link function*

Description

Maps a given link function to its corresponding treatment effect scale.

Usage

```
get_treatment_effect(link)
```

Arguments

link	A character string specifying the link function. Options are: "logit" Log-odds scale. "identity" Risk difference. "probit" Probit scale. "cloglog" Log relative risk for rare events. "log" Log relative risk.
------	--

Value

A character string representing the treatment effect scale.

Examples

```
get_treatment_effect(link = "logit")
get_treatment_effect(link = "identity")
```

marginal_treatment_effect

Marginal treatment effect from reported event counts

Description

Computes the relative treatment effect from aggregate-level data using event counts.

Usage

```
marginal_treatment_effect(ald, ref_trt = NA, comp_trt = NA, scale, family)
```

Arguments

ald	Aggregate-level data
ref_trt	Treatment labels reference (common; e.g. placebo)
comp_trt	Treatment labels comparator
scale	A scaling parameter for the calculation.
family	A character string specifying the family distribution (e.g., "binomial").

Value

Numeric relative treatment effect.

Examples

```
ald <- data.frame(trt = c("B", "C", "B", "C"),
                 variable = c(NA, NA, "y", "y"),
                 statistic = c("N", "N", "sum", "sum"),
                 value = c(100, 100, 50, 60))

marginal_treatment_effect(ald, ref_trt = "C", comp_trt = "B",
                          scale = "log_odds", family = "binomial")
```

marginal_variance	<i>Marginal effect variance using the delta method</i>
-------------------	--

Description

Computes the total variance of marginal treatment effects using the delta method.

Usage

```
marginal_variance(ald, ref_trt = NA, comp_trt = NA, scale, family)
```

Arguments

ald	Aggregate-level data
ref_trt	Treatment labels reference (common; e.g. placebo)
comp_trt	Treatment labels comparator
scale	A scaling parameter for the calculation.
family	A character string specifying the family distribution (e.g., "binomial").

Value

Numeric total variance of marginal treatment effects.

Examples

```
ald <- data.frame(trt = c("B", "C", "B", "C"),
                 variable = c(NA, NA, "y", "y"),
                 statistic = c("N", "N", "sum", "sum"),
                 value = c(100, 100, 50, 60))

marginal_variance(ald, ref_trt = "C", comp_trt = "B",
                 scale = "log_odds", family = "binomial")
```

outstandR	<i>Calculate the difference between treatments using all evidence</i>
-----------	---

Description

This is the main, top-level wrapper for {outstandR}. Methods taken from (Remiro-Azócar et al. 2022).

Usage

```

outstandR(
  ipd_trial,
  ald_trial,
  strategy,
  ref_trt = NA,
  CI = 0.95,
  scale = NULL,
  var_method = NULL,
  seed = NULL,
  verbose = TRUE,
  ...
)

```

Arguments

ipd_trial	Individual-level patient data. For example, suppose between studies <i>A</i> and <i>C</i> . In a long format and must contain a treatment column and outcome column consistent with the formula object. The labels in the treatment are used internally so there must be a common treatment with the aggregate-level data trial.
ald_trial	Aggregate-level data. For example, suppose between studies <i>B</i> and <i>C</i> . The column names are <ul style="list-style-type: none"> • <code>variable</code>: Covariate name. In the case of treatment arm sample size this is NA, • <code>statistic</code>: Summary statistic name from "mean", standard deviation "sd", probability "prop", or "sum", • <code>value</code>: Numerical value of summary statistic, • <code>trt</code>: Treatment label. Because we assume a common covariate distribution between treatment arms this is NA.
strategy	Computation strategy function. These can be <code>strategy_maic()</code> , <code>strategy_stc()</code> , <code>strategy_gcomp_ml()</code> and <code>strategy_gcomp_bayes()</code> .
ref_trt	Reference / common / anchoring treatment name.
CI	Confidence interval level; between 0,1 with default 0.95.
scale	Relative treatment effect scale. If NULL, the scale is automatically determined from the model. Choose from "log-odds", "log_relative_risk", "risk_difference", "delta_z", "mean_difference", "rate_difference" depending on the data type.
var_method	Variance estimation method.
seed	Random seed.
verbose	Logical. If TRUE, prints progress messages and warnings.
...	Additional arguments. Currently, can pass named arguments to <code>rstanarm::stan_glm()</code> via <code>strategy_gcomp_bayes()</code> .

Value

List of length 11 of statistics as a `outstandR` class object. Containing statistics between each pair of treatments. These are the mean, variances and confidence intervals, for contrasts and absolute values.

References

Remiro-Azócar A, Heath A, Baio G (2022). “Parametric G-computation for compatible indirect treatment comparisons with limited individual patient data.” *Res. Synth. Methods*, 1–31. ISSN 1759-2879. doi:10.1002/jrsm.1565. 2108.12208.

See Also

[strategy_maic\(\)](#) [strategy_stc\(\)](#) [strategy_gcomp_ml\(\)](#) [strategy_gcomp_bayes\(\)](#)

Examples

```
data(AC_IPD_binY_contX) # A vs C individual patient-level data
data(BC_ALD_binY_contX) # B vs C aggregate-level data

# linear formula
lin_form <- as.formula("y ~ PF_cont_1 + PF_cont_2 + trt*EM_cont_1 + trt*EM_cont_2")

# sampling values of additional arguments picked for speed
# select appropriate to specific analysis

# matching-adjusted indirect comparison
outstandR_maic <- outstandR(
  AC_IPD_binY_contX, BC_ALD_binY_contX,
  strategy = strategy_maic(formula = lin_form, n_boot = 100))

# simulated treatment comparison
outstandR_stc <- outstandR(
  AC_IPD_binY_contX, BC_ALD_binY_contX,
  strategy = strategy_stc(lin_form))

# G-computation with maximum likelihood
outstandR_gcomp_ml <- outstandR(
  AC_IPD_binY_contX, BC_ALD_binY_contX,
  strategy = strategy_gcomp_ml(lin_form, n_boot = 100, N = 100))

# G-computation with Bayesian inference
outstandR_gcomp_bayes <- outstandR(
  AC_IPD_binY_contX, BC_ALD_binY_contX,
  strategy = strategy_gcomp_bayes(lin_form),
  chains = 1, iter = 1000, warmup = 20)

# Multiple imputation marginalization
outstandR_mim <- outstandR(
  AC_IPD_binY_contX, BC_ALD_binY_contX,
```

```
strategy = strategy_mim(lin_form,
                        N = 100), # size of pseudo-population
chains = 1, iter = 1000, warmup = 20)
```

outstandR-class	<i>outstandR class</i>
-----------------	------------------------

Description

The `outstandR` class contains the results from running a model with the function `outstandR()`.

Details

Objects of class `outstandR` have the following

contrasts A list containing statistics for relative treatment effects:

- `means`: Estimated relative effects (e.g., log-odds ratios, risk differences).
- `variances`: Variance-covariance matrix of the relative effects.
- `contrast_ci`: Confidence intervals for the relative effects.

absolute A list containing statistics for absolute treatment outcomes:

- `means`: Estimated absolute outcomes (e.g., probabilities, mean response).
- `variances`: Variance-covariance matrix of the absolute outcomes.
- `ci`: Confidence intervals for the absolute outcomes.

CI The confidence level used (e.g., 0.95).

ref_trt The name of the reference treatment.

scale The scale of the outcome (e.g., "log odds", "probability").

model A list containing details of the underlying statistical model. Contents vary by strategy:

- `family`: The error distribution and link function.
- `fit`: The underlying model object (e.g., for STC, G-Comp ML, or Bayesian G-Comp).
- `weights`, `ESS`: (MAIC only) The estimated weights and Effective Sample Size.
- `stan_args`: (Bayesian G-Comp, MIM) Arguments passed to Stan.
- `rho`: (G-Comp ML, MIM, Bayesian G-Comp) Correlation coefficient.
- `N`: (G-Comp ML, MIM, Bayesian G-Comp) Number of iterations.
- `nu`, `hats.v`, `M`: (MIM only) Imputation parameters and matrices.

plot.outstandR	<i>Default Plot Method for outstandR Objects</i>
----------------	--

Description

Default Plot Method for outstandR Objects

Usage

```
## S3 method for class 'outstandR'
plot(x, ..., type = c("both", "contrasts", "absolute"), labels = NULL)
```

Arguments

x	An object of class 'outstandR' or a list of 'outstandR' objects.
...	Additional 'outstandR' objects for comparison.
type	Character, one of "both" (default), "contrasts", or "absolute".
labels	Optional character vector of names for the models.

Value

A `ggplot2::ggplot()` object representing the forest plot of the results.

print.outstandR	<i>Print a Summary of a outstandR Object</i>
-----------------	--

Description

This is a method for the function `print()` for objects of the class "outstandR" created by a call to `outstandR()`

Usage

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

Arguments

x	Objects of the class "outstandR"
...	Additional arguments passed to other methods

Value

No return value, called for side effects

See Also

[outstandR\(\)](#)

reshape_ald_to_long *Convert aggregate data from wide to long format*

Description

Convert aggregate data from wide to long format

Usage

```
reshape_ald_to_long(df)
```

Arguments

df A dataframe of ALD

Value

Data frame in long format

reshape_ald_to_wide *Convert aggregate data from long to wide format*

Description

Convert aggregate data from long to wide format

Usage

```
reshape_ald_to_wide(df)
```

Arguments

df A Dataframe of ALD

Value

Data frame in wide format

Examples

```
df <-
  data.frame(
    variable = c("age", "age", "y", "y", "y", "y", "y", "y", "y", "y"),
    statistic = c("mean", "sd", "sum", "bar", "sd", "N", "sum", "bar", "sd", "N"),
    trt = c(NA, NA, "B", "B", "B", "B", "C", "C", "C", "C"),
    value = c(1,1,1,1,1,1,1,1,1,1))
```

strategy-class

Strategy class and subclasses

Description

The strategy class is a virtual class that defines the statistical approach for population adjustment in indirect treatment comparisons. These objects are constructors that validate hyperparameters and encapsulate modelling settings before execution by `outstandR()`.

Details

Objects of class `strategy` have a common structure but carry different subclasses to trigger specific S3 method dispatch.

General fields Shared by all strategies:

- `formula`: The linear regression formula for the outcome model.
- `family`: A base R family object specifying the distribution and link.
- `trt_var`: The name of the treatment variable.

maic subclass Additional fields for Matching-Adjusted Indirect Comparison:

- `n_boot`: Number of bootstrap resamples for variance estimation.

stc subclass Additional fields for Simulated Treatment Comparison:

- `N`: Synthetic sample size for the target population.

gcomp_ml subclass Additional fields for Maximum Likelihood G-computation:

- `rho`: Named square matrix of covariate correlations.
- `marginal_distns`: Names of the marginal distributions for covariates.
- `marginal_params`: Parameters for the marginal distributions.
- `N`: Synthetic sample size for the pseudo-population.
- `n_boot`: Number of bootstrap resamples.

gcomp_bayes subclass Additional fields for Bayesian G-computation:

- `rho`, `marginal_distns`, `marginal_params`, `N`: Same as `gcomp_ml`.
- `...`: Additional arguments passed to the Stan engine via `rstanarm::stan_glm()`.

mim subclass Additional fields for Multiple Imputation Marginalization:

- `rho`: Correlation matrix.
- `N`: Number of iterations/simulated individuals.

strategy_maic	<i>New strategy objects</i>
---------------	-----------------------------

Description

Create a type of strategy class for each modelling approach.

Usage

```
strategy_maic(  
  formula = NULL,  
  family = gaussian(link = "identity"),  
  trt_var = NULL,  
  n_boot = 1000L,  
  moments = 1,  
  int = FALSE,  
  verbatim = TRUE  
)  
  
strategy_stc(  
  formula = NULL,  
  family = gaussian(link = "identity"),  
  trt_var = NULL  
)  
  
strategy_gcomp_ml(  
  formula = NULL,  
  family = gaussian(link = "identity"),  
  trt_var = NULL,  
  rho = NA,  
  marginal_distns = NA,  
  marginal_params = NA,  
  n_boot = 1000L,  
  N = 1000L  
)  
  
strategy_gcomp_bayes(  
  formula = NULL,  
  family = gaussian(link = "identity"),  
  trt_var = NULL,  
  rho = NA,  
  marginal_distns = NA,  
  marginal_params = NA,  
  N = 1000L  
)  
  
strategy_mim(  

```

```

    formula = NULL,
    family = gaussian(link = "identity"),
    trt_var = NULL,
    rho = NA,
    marginal_distns = NA,
    marginal_params = NA,
    N = 1000L
  )
new_strategy(strategy, ...)

```

Arguments

formula	Linear regression formula object. Prognostic factors (PF) are main effects and effect modifiers (EM) are interactions with the treatment variable, e.g., $y \sim X1 + trt + trt:X2$. For covariates as both PF and EM use * syntax.
family	A 'family' object specifying the distribution and link function (e.g., 'binomial'). See <code>stats::family()</code> for more details.
trt_var	Treatment variable name; string
n_boot	The number of resamples used for the non-parametric bootstrap; integer
moments	Integer. The number of moments of the covariates to balance. Setting <code>moments = 2</code> includes both the original variables and their squared terms, which effectively balances both the means and the variances. Default to 1.
int	Logical. If TRUE, includes two-way interactions between all covariates in the balancing model to effectively balance their covariances. Default FALSE
verbatim	Logical. Output to console during running.
rho	A named square matrix of covariate correlations; default NA
marginal_distns	Marginal distributions names; vector default NA. Available distributions are given in <code>stats::Distributions</code> . See <code>copula::Mvdc()</code> for details
marginal_params	Marginal distributions parameters; list of lists, default NA. See <code>copula::Mvdc()</code> for details
N	Synthetic sample size for g-computation
strategy	Class name from <code>strategy_maic</code> , <code>strategy_stc</code> , <code>strategy_gcomp_ml</code> , <code>strategy_gcomp_bayes</code> , <code>strategy_mim</code>
...	Additional arguments

Value

maic class object
 stc class object
 gcomp_ml class object
 gcomp_bayes class object
 mim class object
 Strategy list object

Matching-adjusted indirect comparison (MAIC)

MAIC is a form of non-parametric likelihood reweighting method which allows the propensity score logistic regression model to be estimated without IPD in the *AC* population. The mean outcomes $\mu_{t(AC)}$ on treatment $t = A, B$ in the *AC* target population are estimated by taking a weighted average of the outcomes Y of the N individuals in arm t of the *AB* population.

Used to compare marginal treatment effects where there are cross-trial differences in effect modifiers and limited patient-level data.

$$\hat{Y} = \frac{\sum_{i=1}^N Y_{it(AB)} w_{it}}{\sum_{i=1}^N w_{it}}$$

where the weight w_{it} assigned to the i -th individual receiving treatment t is equal to the odds of being enrolled in the *AC* trial vs the *AB* trial.

Simulated treatment comparison (STC)

[Deprecated]

`strategy_stc()` was deprecated in `outstandR` version 1.X.X. We recommend using G-computation (`strategy_gcomp_ml()`) as a more robust alternative for this type of analysis.

Outcome regression-based method which targets a conditional treatment effect. STC is a modification of the covariate adjustment method. An outcome model is fitted using IPD in the *AB* trial. For example,

$$g(\mu_{t(AB)}(X)) = \beta_0 + \beta_1^T X + (\beta_B + \beta_2^T X^{EM}) I(t = B)$$

where β_0 is an intercept term, β_1 is a vector of coefficients for prognostic variables, β_B is the relative effect of treatment *B* compared to *A* at $X = 0$, β_2 is a vector of coefficients for effect modifiers X^{EM} subvector of the full covariate vector X , and $\mu_{t(AB)}(X)$ is the expected outcome of an individual assigned treatment t with covariate values X which is transformed onto a chosen linear predictor scale with link function $g(\cdot)$.

G-computation maximum likelihood

G-computation marginalizes the conditional estimates by separating the regression modelling from the estimation of the marginal treatment effect for *A* versus *C*. For example, a regression model of the observed outcome y on the covariates x and treatment z is fitted to the *AC* IPD:

$$g(\mu_n) = \beta_0 + \mathbf{x}_n \boldsymbol{\beta}_1 + (\beta_z + \mathbf{x}_n^{EM} \boldsymbol{\beta}_2) I(z_n = 1)$$

In the context of G-computation, this regression model is called the “Q-model”. Having fitted the Q-model, the regression coefficients are treated as nuisance parameters. The parameters are applied to the simulated covariates x_* to predict hypothetical outcomes for each subject under both possible treatments. Namely, a pair of predicted outcomes, also called potential outcomes, under *A* and under *C*, is generated for each subject.

By plugging treatment *C* into the regression fit for every simulated observation, we predict the marginal outcome mean in the hypothetical scenario in which all units are under treatment *C*:

$$\hat{\mu}_0 = \int_{x^*} g^{-1}(\hat{\beta}_0 + x^* \hat{\beta}_1) p(x^*) dx^*$$

To estimate the marginal or population-average treatment effect for *A* versus *C* in the linear predictor scale, one back-transforms to this scale the average predictions, taken over all subjects on the natural outcome scale, and calculates the difference between the average linear predictions:

$$\hat{\Delta}_{10}^{(2)} = g(\hat{\mu}_1) - g(\hat{\mu}_0)$$

G-computation Bayesian

The difference between Bayesian G-computation and its maximum-likelihood counterpart is in the estimated distribution of the predicted outcomes. The Bayesian approach also marginalizes, integrates or standardizes over the joint posterior distribution of the conditional nuisance parameters of the outcome regression, as well as the joint covariate distribution.

Draw a vector of size N^* of predicted outcomes $y^* z$ under each set intervention $z^* \in \{0, 1\}$ from its posterior predictive distribution under the specific treatment. This is defined as $p(y^* z^* | \mathcal{D}_{AC}) = \int_{\beta} p(y^* z^* | \beta) p(\beta | \mathcal{D}_{AC}) d\beta$ where $p(\beta | \mathcal{D}_{AC})$ is the posterior distribution of the outcome regression coefficients β , which encode the predictor-outcome relationships observed in the *AC* trial IPD.

This is given by:

$$\begin{aligned} p(y^* z^* | \mathcal{D}_{AC}) &= \int_{x^*} p(y^* | z^*, x^*, \mathcal{D}_{AC}) p(x^* | \mathcal{D}_{AC}) dx^* \\ &= \int_{x^*} \int_{\beta} p(y^* | z^*, x^*, \beta) p(x^* | \beta) p(\beta | \mathcal{D}_{AC}) d\beta dx^* \end{aligned}$$

In practice, the integrals above can be approximated numerically, using full Bayesian estimation via Markov chain Monte Carlo (MCMC) sampling.

Multiple imputation marginalization (MIM)

MIM targets a marginal treatment effect by using parametric G-computation within a multiple imputation framework. This approach views the covariate adjustment regression as a nuisance model and separates its estimation from the evaluation of the marginal treatment effect of interest. It is particularly useful for ensuring compatibility in indirect comparisons when adjusting for effect modifiers.

Note

While current implementations focus on binary, continuous, and count outcomes, support for survival data (using the `survival` package) is under active development and scheduled for a future version.

See Also

[strategy_gcomp_bayes\(\)](#)
[strategy_gcomp_ml\(\)](#) `copula::Mvdc()`

summary.outstandR	<i>Summary method for outstandR</i>
-------------------	-------------------------------------

Description

Summary method for outstandR

Usage

```
## S3 method for class 'outstandR'  
summary(object, CI = NA, ...)  
  
## S3 method for class 'summary.outstandR'  
print(x, digits = 3, ...)
```

Arguments

object	outstandR() output object.
CI	Confidence interval level.
...	Additional arguments.
x	An object used to select a method.
digits	Minimal number of significant digits, see <code>print.default</code> .

Value

List of class `summary.outstandR`
Original argument, but mainly called for side effects

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