# Package 'TrialEmulation'

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**Title** Causal Analysis of Observational Time-to-Event Data **Version** 0.0.4.0

**Description** Implements target trial emulation methods to apply randomized clinical trial design and analysis in an observational setting. Using marginal structural models, it can estimate intention-to-treat and per-protocol effects in emulated trials using electronic health records. A description and application of the method can be found in Danaei et al (2013) <doi:10.1177/0962280211403603>.

**License** Apache License (>= 2)

URL https://causal-lda.github.io/TrialEmulation/,
 https://github.com/Causal-LDA/TrialEmulation

BugReports https://github.com/Causal-LDA/TrialEmulation/issues

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Collate 'RcppExports.R' 'calculate\_weights.R' 'data.R' 
'data\_extension.R' 'data\_manipulation.R' 'data\_preparation.R' 
'data\_simulation.R' 'data\_utils.R' 'expand\_trials.R' 
'generics.R' 'initiators.R' 'lr\_utils.R' 'te\_model\_fitter.R' 
'te\_outcome\_model.R' 'te\_datastore.R' 'te\_weights.R'

2 Contents

te_data.R' te_expansion.R' trial_sequence.R' modelling.R'
'package.R' 'predict.R' 'robust.R' 'sampling.R'
'te_datastore_csv.R' 'te_datastore_duckdb.R' 'te_parsnip.R'
'te_stats_glm_logit.R' 'utils.R' 'weighting.R'

# NeedsCompilation yes

Author Isaac Gravestock [aut, cre] (<a href="https://orcid.org/0000-0003-0283-2065">https://orcid.org/0000-0003-0283-2065>), Li Su [aut], Page of Page of Cartilla Ca

Roonak Rezvani [aut] (<a href="https://orcid.org/0000-0001-5580-5058">https://orcid.org/0000-0001-5580-5058</a>>, Original package author),

Julia Moesch [aut],

Medical Research Council (MRC) [fnd],

F. Hoffmann-La Roche AG [cph, fnd]

Maintainer Isaac Gravestock <isaac.gravestock@roche.com>

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# **Contents**

calculate_weights	3
case_control_sampling_trials	4
lata_censored	5
lata_preparation	6
expand_trials	9
ît_msm	10
ît_weights_model	11
nitiators	12
pw_data	16
oad_expanded_data	17
outcome_data	18
parsnip_model	
predict_marginal	
print.TE_weight_summary	23
read_expanded_data	23
sample_expanded_data	
save_expanded_data	
save_to_csv	
save_to_datatable	
save_to_duckdb	
set_censor_weight_model	
set_data	
set_expansion_options	
set_outcome_model	
set_switch_weight_model	35
show_weight_models	36
stats_glm_logit	
summary.TE_data_prep	37

calculate\_weights 3

	te_data-class	38
	te_datastore-class	
	te_data_ex	
	te_model_ex	39
	te_model_fitter-class	40
	te_outcome_data-class	<del>1</del> 0
	te_outcome_fitted-class	41
	te_outcome_model-class	41
	trial_example	<del>1</del> 2
	trial_msm	12
	trial_sequence	45
	trial_sequence-class	46
	vignette_switch_data	46
	weight_model_data_indices	17
Index		49

calculate\_weights

Calculate Inverse Probability of Censoring Weights

# **Description**

### [Experimental]

### Usage

```
calculate_weights(object, ...)
## S4 method for signature 'trial_sequence_ITT'
calculate_weights(object, quiet = FALSE)
## S4 method for signature 'trial_sequence_AT'
calculate_weights(object, quiet = FALSE)
## S4 method for signature 'trial_sequence_PP'
calculate_weights(object, quiet = FALSE)
```

# Arguments

object A trial\_sequence object
... Other arguments used by methods.
quiet Prints model summaries is TRUE.

### Value

A trial\_sequence object with updated censor\_weights and/or switch\_weights slots

#### **Examples**

```
save_dir <- file.path(tempdir(), "switch_models")</pre>
ts <- trial_sequence("PP") |>
 set_data(
   data = data_censored,
   id = "id",
   period = "period",
    treatment = "treatment",
    outcome = "outcome",
   eligible = "eligible"
 ) |>
 set_switch_weight_model(
   numerator = \sim age + x1 + x3,
   denominator = ~age,
   model_fitter = stats_glm_logit(save_path = save_dir)
 ) |>
 calculate_weights()
```

case\_control\_sampling\_trials

Case-control sampling of expanded data for the sequence of emulated trials

### **Description**

[Stable]

#### Usage

```
case_control_sampling_trials(
  data_prep,
  p_control = NULL,
  subset_condition,
  sort = FALSE
)
```

### Arguments

data\_prep Result from data\_preparation().

p\_control Control sampling probability for selecting potential controls at each follow-up

time of each trial.

subset\_condition

Expression used to subset() the trial data before case-control sampling.

Sort data before applying case-control sampling to make sure that the resulting data are identical when sampling from the expanded data created with separate\_files

= TRUE or separate\_files = FALSE.

data\_censored 5

#### **Details**

Perform case-control sampling of expanded data to create a data set of reduced size and calculate sampling weights to be used in trial\_msm().

#### Value

A data.frame or a split() data.frame if length(p\_control) > 1. An additional column sample\_weight containing the sample weights will be added to the result. These can be included in the models fit with trial\_msm().

## **Examples**

### **Description**

This data contains data from 89 patients followed for up to 19 periods.

censoring

#### Usage

data\_censored

### Format

A data frame with 725 rows and 12 variables:

id patient identifier

period time period

treatment indicator for receiving treatment in this period, 1=treatment, 0=non-treatment

- **x1** A time-varying categorical variable relating to treatment and the outcome
- x2 A time-varying numeric variable relating to treatment and the outcome
- x3 A fixed categorical variable relating to treatment and the outcome
- x4 A fixed categorical variable relating to treatment and the outcome

age patient age in years

age\_s patient age

outcome indicator for outcome in this period, 1=event occurred, 0=no event

censored indicator for patient being censored in this period, 1=censored, 0=not censored

eligible indicator for eligibility for trial start in this period, 1=yes, 0=no

data\_preparation

data\_preparation

Prepare data for the sequence of emulated target trials

#### **Description**

[Stable]

# Usage

```
data_preparation(
  data,
  id = "id",
 period = "period",
  treatment = "treatment",
 outcome = "outcome",
 eligible = "eligible",
 model_var = NULL,
 outcome_cov = \sim 1,
  estimand_type = c("ITT", "PP", "As-Treated"),
  switch_n_cov = ~1,
  switch_d_cov = ~1,
  first_period = NA,
  last_period = NA,
  use_censor_weights = FALSE,
  cense = NA,
  pool_cense = c("none", "both", "numerator"),
  cense_d_cov = ~1,
  cense_n_cov = \sim1,
  eligible_wts_0 = NA,
  eligible_wts_1 = NA,
 where_var = NULL,
  data_dir,
  save_weight_models = FALSE,
 glm_function = "glm",
  chunk_size = 500,
  separate_files = FALSE,
  quiet = FALSE,
)
```

# Arguments

A data. frame containing all the required variables in the person-time format, i.e., the 'long' format.

Id Name of the variable for identifiers of the individuals. Default is 'id'.

period Name of the variable for the visit/period. Default is 'period'.

data\_preparation 7

Default is 'outcome'.

Name of the variable for the treatment indicator at that visit/period. Default is 'treatment'.

Name of the variable for the indicator of the outcome event at that visit/period.

Name of the variable for the indicator of eligibility for the target trial at that

visit/period. Default is 'eligible'.

Treatment variables to be included in the marginal structural model for the emulated trials. model\_var = "assigned\_treatment" will create a variable assigned\_treatment that is the assigned treatment at the trial baseline, typically used for ITT and per-protocol analyses. model\_var = "dose" will create a variable dose that is the cumulative number of treatments received since the trial baseline, typically used in as-treated analyses.

A RHS formula with baseline covariates to be adjusted for in the marginal structural model for the emulated trials. Note that if a time-varying covariate is specified in outcome\_cov, only its value at each of the trial baselines will be included in the expanded data.

Specify the estimand for the causal analyses in the sequence of emulated trials. estimand\_type = "ITT" will perform intention-to-treat analyses, where treatment switching after trial baselines are ignored. estimand\_type = "PP" will perform per-protocol analyses, where individuals' follow-ups are artificially censored and inverse probability of treatment weighting is applied. estimand\_type = "As-Treated" will fit a standard marginal structural model for all possible treatment sequences, where individuals' follow-ups are not artificially censored but treatment switching after trial baselines are accounted for by applying inverse probability of treatment weighting.

A RHS formula to specify the logistic models for estimating the numerator terms of the inverse probability of treatment weights. A derived variable named time\_on\_regime containing the duration of time that the individual has been on the current treatment/non-treatment is available for use in these models.

A RHS formula to specify the logistic models for estimating the denominator terms of the inverse probability of treatment weights.

First time period to be set as trial baseline to start expanding the data.

Last time period to be set as trial baseline to start expanding the data.

use\_censor\_weights

Require the inverse probability of cen

Require the inverse probability of censoring weights. If use\_censor\_weights = TRUE, then the variable name of the censoring indicator needs to be provided in the argument cense.

Variable name for the censoring indicator. Required if use\_censor\_weights = TRUE.

Fit pooled or separate censoring models for those treated and those untreated at the immediately previous visit. Pooling can be specified for the models for the numerator and denominator terms of the inverse probability of censoring weights. One of "none", "numerator", or "both" (default is "none" except when estimand\_type = "ITT" then default is "numerator").

outcome\_cov

outcome

eligible

model\_var

estimand\_type

switch\_n\_cov

switch\_d\_cov

first\_period
last\_period

cense

pool\_cense

8 data\_preparation

cense\_d\_cov A RHS formula to specify the logistic models for estimating the denominator terms of the inverse probability of censoring weights.

cense\_n\_cov A RHS formula to specify the logistic models for estimating the numerator terms of the inverse probability of censoring weights.

eligible\_wts\_0 See definition for eligible\_wts\_1

visit.

eligible\_wts\_1 Exclude some observations when fitting the models for the inverse probability of treatment weights. For example, if it is assumed that an individual will stay on treatment for at least 2 visits, the first 2 visits after treatment initiation by definition have a probability of staying on the treatment of 1.0 and should thus be excluded from the weight models for those who are on treatment at the immediately previous visit. Users can define a variable that indicates that these 2 observations are ineligible for the weight model for those who are on treatment at the immediately previous visit and add the variable name in the argument eligible\_wts\_1. Similar definitions are applied to eligible\_wts\_0 for excluding observations when fitting the models for the inverse probability of treatment weights for those who are not on treatment at the immediately previous

Specify the variable names that will be used to define subgroup conditions when fitting the marginal structural model for a subgroup of individuals. Need to specify jointly with the argument where \_case.

Directory to save model objects when save\_weight\_models=TRUE and expanded data as separate CSV files names as trial\_i.csvs if separate\_files = TRUE. If the specified directory does not exist it will be created. If the directory already contains trial files, an error will occur, other files may be overwritten.

save\_weight\_models

Save model objects for estimating the weights in data\_dir.

Specify which glm function to use for the marginal structural model from the stats or parglm packages. The default function is the glm function in the stats package. Users can also specify glm\_function = "parglm" such that the parglm function in the parglm package can be used for fitting generalized linear models in parallel. The default control setting for parglm is nthreads = 4 and method = "FAST", where four cores and Fisher information are used for faster computation. Users can change the default control setting by passing the arguments nthreads and method in the parglm.control function of the parglm package, or alternatively, by passing a control argument with a list produced by parglm.control(nthreads = , method = ).

Number of individuals whose data to be processed in one chunk when separate\_files = TRUE

separate\_files Save expanded data in separate CSV files for each trial.

quiet Suppress the printing of progress messages and summaries of the fitted models.

Additional arguments passed to glm\_function. This may be used to specify initial values of parameters or arguments to control. See stats::glm, parglm::parglm and parglm::parglm.control() for more information.

where\_var

data\_dir

glm\_function

chunk\_size

uict

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expand\_trials 9

#### **Details**

This function expands observational data in the person-time format (i.e., the 'long' format) to emulate a sequence of target trials and also estimates the inverse probability of treatment and censoring weights as required.

The arguments chunk\_size and separate\_files allow for processing of large datasets that would not fit in memory once expanded. When separate\_files = TRUE, the input data are processed in chunks of individuals and saved into separate files for each emulated trial. These separate files can be sampled by case-control sampling to create a reduced dataset for the modelling.

#### Value

An object of class TE\_data\_prep, which can either be sampled from (case\_control\_sampling\_trials) or directly used in a model (trial\_msm). It contains the elements

data the expanded dataset for all emulated trials. If separate\_files = FALSE, it is a data.table;
 if separate\_files = TRUE, it is a character vector with the file path of the expanded data as
 CSV files.

min\_period index for the first trial in the expanded data

max\_period index for the last trial in the expanded data

N the total number of observations in the expanded data

data\_template a zero-row data. frame with the columns and attributes of the expanded data
switch\_models a list of summaries of the models fitted for inverse probability of treatment weights,
if estimand\_type is "PP" or "As-Treated"

**censor\_models** a list of summaries of the models fitted for inverse probability of censoring weights, if use\_censor\_weights=TRUE

args a list contain the parameters used to prepare the data and fit the weight models

expand\_trials

Expand trials

#### **Description**

#### [Experimental]

### Usage

expand\_trials(object)

#### **Arguments**

object

A trial\_sequence object

#### Value

The trial\_sequence object with a data set containing the full sequence of target trials. The data is stored according to the options set with set\_expansion\_options() and especially the save\_to\_\* function.

10 fit\_msm

fit\_msm

Fit the marginal structural model for the sequence of emulated trials

#### **Description**

### [Experimental]

#### Usage

```
fit_msm(
  object,
 weight_cols = c("weight", "sample_weight"),
 modify_weights = NULL
)
## S4 method for signature 'trial_sequence'
fit_msm(
  object,
 weight_cols = c("weight", "sample_weight"),
 modify_weights = NULL
)
```

#### **Arguments**

object A trial\_sequence object

weight\_cols character vector of column names in expanded outcome dataset, ie outcome\_data(object).

If multiple columns are specified, the element wise product will be used. Specify

NULL if no weight columns should be used.

modify\_weights a function to transform the weights (or NULL for no transformation). Must take a numeric vector of weights and a vector of positive, finite weights of the same length. See examples for some possible function definitions.

> Before the outcome marginal structural model can be fit, the outcome model must be specified with set\_outcome\_model() and the data must be expanded into the trial sequence with expand\_trials().

> The model is fit based on the model\_fitter specified in set\_outcome\_model using the internal fit\_outcome\_model method.

#### Value

A modified trial\_sequence object with updated outcome\_model slot.

#### **Examples**

```
trial_seq_object <- trial_sequence("ITT") |>
 set_data(data_censored) |>
 set_outcome_model(
   adjustment_terms = ~age_s,
```

fit\_weights\_model 11

```
followup_time_terms = ~ stats::poly(followup_time, degree = 2)
  set_expansion_options(output = save_to_datatable(), chunk_size = 500) |>
  expand_trials() |>
  load_expanded_data()
fit_msm(trial_seq_object)
# Using modify_weights functions ----
# returns a function that truncates weights to limits
limit_weight <- function(lower_limit, upper_limit) {</pre>
  function(w) {
    w[w > upper_limit] <- upper_limit</pre>
    w[w < lower_limit] <- lower_limit</pre>
  }
}
# calculate 1st and 99th percentile limits and truncate
p99_weight <- function(w) {
  p99 <- quantile(w, prob = c(0.01, 0.99), type = 1)
  limit_weight(p99[1], p99[2])(w)
}
# set all weights to 1
all_ones <- function(w) {</pre>
  rep(1, length(w))
}
fit_msm(trial_seq_object, modify_weights = limit_weight(0.01, 4))
fit_msm(trial_seq_object, modify_weights = p99_weight)
```

fit\_weights\_model

Method for fitting weight models

### **Description**

Method for fitting weight models

### Usage

```
fit_weights_model(object, data, formula, label)
```

#### **Arguments**

object	The object determining which method should be used, containing any slots containing user defined parameters.
data	data.frame containing outcomes and covariates as defined in formula.
formula	formula describing the model.
label	A short string describing the model.

#### Value

An object of class te\_weights\_fitted

#### **Examples**

```
fitter <- stats_glm_logit(tempdir())
data(data_censored)
# Not usually called directly by a user
fitted <- fit_weights_model(
   object = fitter,
   data = data_censored,
   formula = 1 - censored ~ x1 + age_s + treatment,
   label = "Example model for censoring"
)
fitted
unlink(fitted@summary$save_path$path)</pre>
```

initiators

A wrapper function to perform data preparation and model fitting in a sequence of emulated target trials

#### **Description**

[Stable]

### Usage

```
initiators(
 data,
  id = "id",
  period = "period",
  treatment = "treatment",
 outcome = "outcome",
  eligible = "eligible",
  outcome_cov = \sim 1,
  estimand_type = c("ITT", "PP", "As-Treated"),
 model_var = NULL,
  switch_n_cov = ~1,
  switch_d_cov = ~1,
  first_period = NA,
  last_period = NA,
  first_followup = NA,
  last_followup = NA,
  use_censor_weights = FALSE,
  save_weight_models = FALSE,
  analysis_weights = c("asis", "unweighted", "p99", "weight_limits"),
  weight_limits = c(0, Inf),
  cense = NA,
```

```
pool_cense = c("none", "both", "numerator"),
  cense_d_cov = ~1,
  cense_n_cov = ~1,
  include_followup_time = ~followup_time + I(followup_time^2),
  include_trial_period = ~trial_period + I(trial_period^2),
  eligible_wts_0 = NA,
  eligible_wts_1 = NA,
  where_var = NULL,
  where_case = NA,
  data_dir,
  glm_function = "glm",
  quiet = FALSE,
  ...
)
```

#### **Arguments**

data A data. frame containing all the required variables in the person-time format,

i.e., the 'long' format.

id Name of the variable for identifiers of the individuals. Default is 'id'.

period Name of the variable for the visit/period. Default is 'period'.

treatment Name of the variable for the treatment indicator at that visit/period. Default is

'treatment'.

outcome Name of the variable for the indicator of the outcome event at that visit/period.

Default is 'outcome'.

eligible Name of the variable for the indicator of eligibility for the target trial at that

visit/period. Default is 'eligible'.

outcome\_cov A RHS formula with baseline covariates to be adjusted for in the marginal struc-

tural model for the emulated trials. Note that if a time-varying covariate is specified in outcome\_cov, only its value at each of the trial baselines will be included

in the expanded data.

estimand\_type Specify the estimand for the causal analyses in the sequence of emulated trials. estimand\_type = "ITT" will perform intention-to-treat analyses, where

treatment switching after trial baselines are ignored. estimand\_type = "PP" will perform per-protocol analyses, where individuals' follow-ups are artificially censored and inverse probability of treatment weighting is applied. estimand\_type = "As-Treated" will fit a standard marginal structural model for all possible treatment sequences, where individuals' follow-ups are not artificially censored but treatment switching after trial baselines are accounted for by applying in-

verse probability of treatment weighting.

model\_var Treatment variables to be included in the marginal structural model for the

emulated trials. model\_var = "assigned\_treatment" will create a variable assigned\_treatment that is the assigned treatment at the trial baseline, typically used for ITT and per-protocol analyses. model\_var = "dose" will create a variable dose that is the cumulative number of treatments received since the

trial baseline, typically used in as-treated analyses.

switch\_n\_cov A RHS formula to specify the logistic models for estimating the numerator terms of the inverse probability of treatment weights. A derived variable named time\_on\_regime containing the duration of time that the individual has been on

the current treatment/non-treatment is available for use in these models.

A RHS formula to specify the logistic models for estimating the denominator switch\_d\_cov

terms of the inverse probability of treatment weights.

first\_period First time period to be set as trial baseline to start expanding the data.

last\_period Last time period to be set as trial baseline to start expanding the data.

first\_followup First follow-up time/visit in the trials to be included in the marginal structural

model for the outcome event.

last\_followup Last follow-up time/visit in the trials to be included in the marginal structural

model for the outcome event.

use\_censor\_weights

Require the inverse probability of censoring weights. If use\_censor\_weights = TRUE, then the variable name of the censoring indicator needs to be provided in the argument cense.

save\_weight\_models

Save model objects for estimating the weights in data\_dir.

analysis\_weights

Choose which type of weights to be used for fitting the marginal structural model for the outcome event.

- "asis": use the weights as calculated.
- "p99": use weights truncated at the 1st and 99th percentiles (based on the distribution of weights in the entire sample).
- "weight\_limits": use weights truncated at the values specified in weight\_limits.
- "unweighted": set all analysis weights to 1, even if treatment weights or censoring weights were calculated.

Lower and upper limits to truncate weights, given as c(lower, upper) weight\_limits

Variable name for the censoring indicator. Required if use\_censor\_weights = cense

TRUE.

pool\_cense Fit pooled or separate censoring models for those treated and those untreated

at the immediately previous visit. Pooling can be specified for the models for the numerator and denominator terms of the inverse probability of censoring weights. One of "none", "numerator", or "both" (default is "none" except

when estimand\_type = "ITT" then default is "numerator").

A RHS formula to specify the logistic models for estimating the denominator cense\_d\_cov

terms of the inverse probability of censoring weights.

A RHS formula to specify the logistic models for estimating the numerator terms cense\_n\_cov

of the inverse probability of censoring weights.

include\_followup\_time

The model to include the follow up time/visit of the trial (followup\_time) in the marginal structural model, specified as a RHS formula.

include\_trial\_period

The model to include the trial period (trial\_period) in the marginal structural model, specified as a RHS formula.

eligible\_wts\_0 See definition for eligible\_wts\_1

eligible\_wts\_1 Exclude some observations when fitting the models for the inverse probability

of treatment weights. For example, if it is assumed that an individual will stay on treatment for at least 2 visits, the first 2 visits after treatment initiation by definition have a probability of staying on the treatment of 1.0 and should thus be excluded from the weight models for those who are on treatment at the immediately previous visit. Users can define a variable that indicates that these 2 observations are ineligible for the weight model for those who are on treatment at the immediately previous visit and add the variable name in the argument eligible\_wts\_1. Similar definitions are applied to eligible\_wts\_0 for excluding observations when fitting the models for the inverse probability of treatment weights for those who are not on treatment at the immediately previous

visit.

where\_var Specify the variable names that will be used to define subgroup conditions when

fitting the marginal structural model for a subgroup of individuals. Need to

specify jointly with the argument where\_case.

where\_case Define conditions using variables specified in where\_var when fitting a marginal

structural model for a subgroup of the individuals. For example, if where\_var= "age", where\_case = "age >= 30" will only fit the marginal structural model to

the subgroup of individuals. who are 30 years old or above.

data\_dir Directory to save model objects in.

glm\_function Specify which glm function to use for the marginal structural model from the

stats or parglm packages. The default function is the glm function in the stats package. Users can also specify glm\_function = "parglm" such that the parglm function in the parglm package can be used for fitting generalized linear models in parallel. The default control setting for parglm is nthreads = 4 and method = "FAST", where four cores and Fisher information are used for faster computation. Users can change the default control setting by passing the arguments nthreads and method in the parglm.control function of the parglm package, or alternatively, by passing a control argument with a list produced

by parglm.control(nthreads = , method = ).

quiet Suppress the printing of progress messages and summaries of the fitted models.

Additional arguments passed to glm\_function. This may be used to specify initial values of parameters or arguments to control. See stats::glm, par-

glm::parglm and parglm::parglm.control() for more information.

#### Details

An all-in-one analysis using a sequence of emulated target trials. This provides a simplified interface to the main functions data\_preparation() and trial\_msm().

#### Value

Returns the result of trial\_msm() from the expanded data. An object of class TE\_msm containing

model a glm object

robust a list containing a summary table of estimated regression coefficients and the robust covariance matrix

ipw\_data

ipw\_data

IPW Data Accessor and Setter

### **Description**

### [Experimental]

### Usage

```
ipw_data(object)
ipw_data(object) <- value

## S4 method for signature 'trial_sequence'
ipw_data(object)

## S4 replacement method for signature 'trial_sequence'
ipw_data(object) <- value</pre>
```

### **Arguments**

object trial\_sequence object

value data.table to replace and update in @data

#### **Details**

Generic function to access and update the data used for inverse probability weighting.

The setter method ipw\_data(object) <- value does not perform the same checks and manipulations as set\_data(). To completely replace the data please use set\_data(). This ipw\_data<- method allows small changes such as adding a new column.

#### Value

The data from the @data slot of object used for inverse probability weighting.

#### **Examples**

```
ts <- trial_sequence("ITT")
ts <- set_data(ts, data_censored)
ipw_data(ts)
data.table::set(ipw_data(ts), j = "dummy", value = TRUE)
# or with the setter method:
new_data <- ipw_data(ts)
new_data$x2sq <- new_data$x2^2
ipw_data(ts) <- new_data</pre>
```

load\_expanded\_data 17

load\_expanded\_data

Method to read, subset and sample expanded data

#### **Description**

#### [Experimental]

### Usage

```
load_expanded_data(
  object,
  p_control = NULL,
  period = NULL,
  subset_condition = NULL,
  seed = NULL
)

## S4 method for signature 'trial_sequence'
load_expanded_data(
  object,
  p_control = NULL,
  period = NULL,
  subset_condition = NULL,
  seed = NULL
)
```

#### **Arguments**

object An object of class trial\_sequence.

p\_control Probability of selecting a control, NULL for no sampling (default).

period An integerish vector of non-zero length to select trial period(s) or NULL (default)

to select all trial periods.

subset\_condition

A string or NULL (default). subset\_condition will be translated to a call (in case the expanded data is saved as a data.table or in the csv format) or to a SQL-query (in case the expanded data is saved as a duckdb file).

The operators "==", "!=", ">", ">=", "<", "<=", %in%", "&", "|" are supported. Numeric vectors can be written as c(1, 2, 3) or 1:3. Variables are not supported.

*Note*: Make sure numeric vectors written as 1:3 are surrounded by spaces, e.g. a %in% c(1:4, 6:9), otherwise the code will fail.

seed An integer seed or NULL (default).

*Note*: The same seed will return a different result depending on the class of the te\_datastore object contained in the trial\_sequence object.

18 outcome\_data

#### **Details**

This method is used on trial\_sequence objects to read, subset and sample expanded data.

#### Value

An updated trial\_sequence object, the data is stored in slot @outcome\_data as a te\_outcome\_data object.

### **Examples**

```
# create a trial_sequence-class object
trial_itt_dir <- file.path(tempdir(), "trial_itt")</pre>
dir.create(trial_itt_dir)
trial_itt <- trial_sequence(estimand = "ITT") |>
  set_data(data = data_censored) |>
  set_outcome_model(adjustment_terms = ~ x1 + x2)
trial_itt_csv <- set_expansion_options(</pre>
  trial_itt,
  output = save_to_csv(file.path(trial_itt_dir, "trial_csvs")),
  chunk\_size = 500
) |>
  expand_trials()
# load_expanded_data default behaviour returns all trial_periods and doesn't sample
load_expanded_data(trial_itt_csv)
# load_expanded_data can subset the data before sampling
load_expanded_data(
  trial_itt_csv,
  p_{control} = 0.2,
  period = 1:20,
  subset_condition = "followup_time %in% 1:20 & x2 < 1",</pre>
# delete after use
unlink(trial_itt_dir, recursive = TRUE)
```

outcome\_data

Outcome Data Accessor and Setter

### **Description**

[Experimental]

parsnip\_model 19

#### Usage

```
outcome_data(object)
outcome_data(object) <- value

## S4 method for signature 'trial_sequence'
outcome_data(object)

## S4 replacement method for signature 'trial_sequence'
outcome_data(object) <- value</pre>
```

### **Arguments**

object trial\_sequence object

value data.table to replace and update in @outcome\_data

#### **Details**

Generic function to outcome data

#### Value

The object with updated outcome data

### **Examples**

```
ts <- trial_sequence("ITT")
new_data <- data.table::data.table(vignette_switch_data[1:200, ])
new_data$weight <- 1
outcome_data(ts) <- new_data</pre>
```

parsnip\_model

Fit outcome models using parsnip models

### **Description**

#### [Experimental]

# Usage

```
parsnip_model(model_spec, save_path)
```

# Arguments

model\_spec A parsnip model definition with mode = "classification".

save\_path Directory to save models. Set to NA if models should not be saved.

20 predict\_marginal

#### **Details**

Specify that the models should be fit using a classification model specified with the parsnip package.

Warning: This functionality is experimental and not recommended for use in analyses. *sqrtn*-consistency estimation and valid inference of the parameters in marginal structural models for emulated trials generally require that the weights for treatment switching and censoring be estimated at parametric rates, which is generally not possible when using data-adaptive estimation of high-dimensional regressions. Therefore, we only recommend using stats\_glm\_logit().

#### Value

An object of class te\_parsnip\_model inheriting from te\_model\_fitter which is used for dispatching methods for the fitting models.

#### See Also

```
Other model_fitter: stats_glm_logit(), te_model_fitter-class
```

# Examples

```
## Not run:
if (
  requireNamespace("parsnip", quietly = TRUE) &&
    requireNamespace("rpart", quietly = TRUE)
) {
  # Use a decision tree model fitted with the rpart package
  parsnip_model(
    model_spec = parsnip::decision_tree(tree_depth = 30) |>
        set_mode("classification") |>
        set_engine("rpart"),
        save_path = tempdir()
)
}
## End(Not run)
```

predict\_marginal

Predict marginal cumulative incidences with confidence intervals for a target trial population

# Description

**[Stable]** This function predicts the marginal cumulative incidences when a target trial population receives either the treatment or non-treatment at baseline (for an intention-to-treat analysis) or either sustained treatment or sustained non-treatment (for a per-protocol analysis). The difference between these cumulative incidences is the estimated causal effect of treatment. Currently, the predict function only provides marginal intention-to-treat and per-protocol effects, therefore it is only valid when estimand\_type = "ITT" or estimand\_type = "PP".

predict\_marginal 21

### Usage

```
predict(object, ...)
## S4 method for signature 'trial_sequence_ITT'
predict(
  object,
  newdata,
  predict_times,
  conf_int = TRUE,
  samples = 100,
  type = c("cum_inc", "survival")
)
## S4 method for signature 'trial_sequence_PP'
predict(
 object,
  newdata,
 predict_times,
  conf_int = TRUE,
  samples = 100,
  type = c("cum_inc", "survival")
)
## S3 method for class 'TE_msm'
predict(
 object,
  newdata,
 predict_times,
 conf_int = TRUE,
  samples = 100,
  type = c("cum_inc", "survival"),
)
```

### **Arguments**

object from trial\_msm() or initiators() or trial\_sequence.

... Further arguments passed to or from other methods.

newdata Baseline trial data that characterise the target trial population that marginal cu-

mulative incidences or survival probabilities are predicted for. newdata must have the same columns and formats of variables as in the fitted marginal structural model specified in trial\_msm() or initiators(). If newdata contains

rows with followup\_time > 0 these will be removed.

predict\_times Specify the follow-up visits/times where the marginal cumulative incidences or

survival probabilities are predicted.

conf\_int Construct the point-wise 95-percent confidence intervals of cumulative inci-

dences for the target trial population under treatment and non-treatment and their

22 predict\_marginal

differences by simulating the parameters in the marginal structural model from a multivariate normal distribution with the mean equal to the marginal structural model parameter estimates and the variance equal to the estimated robust covariance matrix.

samples Number of samples used to construct the simulation-based confidence intervals.

type Specify cumulative incidences or survival probabilities to be predicted. Either cumulative incidence ("cum\_inc") or survival probability ("survival").

#### Value

A list of three data frames containing the cumulative incidences for each of the assigned treatment options (treatment and non-treatment) and the difference between them.

#### **Examples**

```
# Prediction for initiators() or trial_msm() objects ----
# If necessary set the number of `data.table` threads
data.table::setDTthreads(2)
data("te_model_ex")
predicted_ci <- predict(te_model_ex, predict_times = 0:30, samples = 10)</pre>
# Plot the cumulative incidence curves under treatment and non-treatment
plot(predicted_ci[[1]]$followup_time, predicted_ci[[1]]$cum_inc,
 type = "1",
 xlab = "Follow-up Time", ylab = "Cumulative Incidence",
 ylim = c(0, 0.7)
lines(predicted_ci[[1]]$followup_time, predicted_ci[[1]]$^2.5%^, lty = 2)
lines(predicted_ci[[1]]$followup_time, predicted_ci[[1]]$^97.5%^, lty = 2)
lines(predicted_ci[[2]]$followup_time, predicted_ci[[2]]$cum_inc, type = "1", col = 2)
lines(predicted_ci[[2]]$followup_time, predicted_ci[[2]]$^2.5%^, 1ty = 2, col = 2)
lines(predicted_ci[[2]]$followup_time, predicted_ci[[2]]$^97.5%^, lty = 2, col = 2)
legend("topleft", title = "Assigned Treatment", legend = c("0", "1"), col = 1:2, lty = 1)
# Plot the difference in cumulative incidence over follow up
plot(predicted_ci[[3]]$followup_time, predicted_ci[[3]]$cum_inc_diff,
 type = "1",
 xlab = "Follow-up Time", ylab = "Difference in Cumulative Incidence",
 ylim = c(0.0, 0.5)
lines(predicted_ci[[3]]$followup_time, predicted_ci[[3]]$^2.5%^, lty = 2)
lines(predicted_ci[[3]]$followup_time, predicted_ci[[3]]$^97.5%^, 1ty = 2)
```

```
print.TE_weight_summary
```

Print a weight summary object

### **Description**

[Stable]

### Usage

```
## S3 method for class 'TE_weight_summary'
print(x, full = TRUE, ...)
```

#### **Arguments**

x print TE\_weight\_summary object.

full Print full or short summary.

... Arguments passed to print.data.frame.

#### Value

No return value, only for printing.

read\_expanded\_data

Method to read expanded data

# Description

This method is used on te\_datastore objects to read selected data and return one data. table.

#### Usage

```
read_expanded_data(object, period = NULL, subset_condition = NULL)
## S4 method for signature 'te_datastore_datatable'
read_expanded_data(object, period = NULL, subset_condition = NULL)
```

### Arguments

object An object of class te\_datastore.

period An integerish vector of non-zero length to select trial period(s) or NULL (default)

to select all files.

subset\_condition

A string of length 1 or NULL (default).

### Value

A data.frame of class data.table.

# **Examples**

```
# create a te_datastore_csv object and save some data
temp_dir <- tempfile("csv_dir_")
dir.create(temp_dir)
datastore <- save_to_csv(temp_dir)
data(vignette_switch_data)
expanded_csv_data <- save_expanded_data(datastore, vignette_switch_data[1:200, ])
# read expanded data
read_expanded_data(expanded_csv_data)
# delete after use
unlink(temp_dir, recursive = TRUE)</pre>
```

sample\_expanded\_data
Internal method to sample expanded data

### **Description**

Internal method to sample expanded data

#### Usage

```
sample_expanded_data(
  object,
  p_control,
  period = NULL,
  subset_condition = NULL,
  seed
)

## S4 method for signature 'te_datastore'
sample_expanded_data(
  object,
  p_control,
  period = NULL,
  subset_condition = NULL,
  seed
)
```

save\_expanded\_data 25

### **Arguments**

object An object of class te\_datastore.
p\_control Probability of selecting a control.

period An integerish vector of non-zero length to select trial period(s) or NULL (default)

to select all trial periods.

subset\_condition

A string or NULL.

seed An integer seed or NULL (default).

#### Value

A data.frame of class data.table.

### **Examples**

```
# Data object normally created by [expand_trials]
datastore <- new("te_datastore_datatable", data = te_data_ex$data, N = 50139L)
sample_expanded_data(datastore, period = 260:275, p_control = 0.2, seed = 123)</pre>
```

save\_expanded\_data

Method to save expanded data

### Description

This method is used internally by expand\_trials to save the data to the "datastore" defined in set\_expansion\_options.

### Usage

```
save_expanded_data(object, data)
## S4 method for signature 'te_datastore_datatable'
save_expanded_data(object, data)
```

### **Arguments**

object An object of class te\_datastore or a child class.

data A data frame containing the expanded trial data. The columns trial\_period

and id are present, which may be used in methods to save the data in an optimal

way, such as with indexes, keys or separate files.

#### Value

An updated object with the data stored. Notably object@N should be increased

26 save\_to\_csv

#### **Examples**

```
temp_dir <- tempfile("csv_dir_")
dir.create(temp_dir)
datastore <- save_to_csv(temp_dir)
data(vignette_switch_data)
save_expanded_data(datastore, vignette_switch_data[1:200, ])
# delete after use
unlink(temp_dir, recursive = TRUE)</pre>
```

save\_to\_csv

Save expanded data as CSV

#### **Description**

### [Experimental]

### Usage

```
save_to_csv(path)
```

### **Arguments**

path

Directory to save CSV files in. Must be empty.

#### Value

A te\_datastore\_csv object.

### See Also

```
Other save_to: save_to_datatable(), save_to_duckdb(), set_expansion_options()
```

### **Examples**

```
csv_dir <- file.path(tempdir(), "expanded_trials_csv")
dir.create(csv_dir)
csv_datastore <- save_to_csv(path = csv_dir)

trial_to_expand <- trial_sequence("ITT") |>
    set_data(data = data_censored) |>
    set_expansion_options(output = csv_datastore, chunk_size = 500)

# Delete directory after use
unlink(csv_dir)
```

save\_to\_datatable 27

save\_to\_datatable

Save expanded data as a data.table

# Description

### [Experimental]

# Usage

```
save_to_datatable()
```

#### See Also

```
Other save_to: save_to_csv(), save_to_duckdb(), set_expansion_options()
```

### **Examples**

```
trial_to_expand <- trial_sequence("ITT") |>
  set_data(data = data_censored) |>
  set_expansion_options(output = save_to_datatable(), chunk_size = 500)
```

save\_to\_duckdb

Save expanded data to DuckDB

### **Description**

### [Experimental]

# Usage

```
save_to_duckdb(path)
```

# Arguments

path

Directory to save DuckDB database file in.

#### Value

```
A te_datastore_duckdb object.
```

#### See Also

```
Other save_to: save_to_csv(), save_to_datatable(), set_expansion_options()
```

## **Examples**

```
if (require(duckdb)) {
  duckdb_dir <- file.path(tempdir(), "expanded_trials_duckdb")

trial_to_expand <- trial_sequence("ITT") |>
  set_data(data = data_censored) |>
  set_expansion_options(output = save_to_duckdb(path = duckdb_dir), chunk_size = 500)

# Delete directory after use
unlink(duckdb_dir)
}
```

set\_censor\_weight\_model

Set censoring weight model

### **Description**

#### [Experimental]

### Usage

```
set_censor_weight_model(
 object,
  censor_event,
 numerator,
 denominator,
 pool_models = NULL,
 model_fitter
)
## S4 method for signature 'trial_sequence'
set_censor_weight_model(
 object,
 censor_event,
  numerator,
  denominator,
 pool_models = c("none", "both", "numerator"),
 model_fitter = stats_glm_logit()
)
## S4 method for signature 'trial_sequence_PP'
set_censor_weight_model(
 object,
  censor_event,
 numerator,
```

```
denominator,
  pool_models = "none",
 model_fitter = stats_glm_logit()
)
## S4 method for signature 'trial_sequence_ITT'
set_censor_weight_model(
  object,
  censor_event,
 numerator,
  denominator,
  pool_models = "numerator";
 model_fitter = stats_glm_logit()
)
## S4 method for signature 'trial_sequence_AT'
set_censor_weight_model(
  object,
  censor_event,
  numerator,
 denominator,
 pool_models = "none",
 model_fitter = stats_glm_logit()
)
```

# **Arguments**

object trial sequence.

censor\_event string. Name of column containing censoring indicator.

numerator A RHS formula to specify the logistic models for estimating the numerator terms

of the inverse probability of censoring weights.

denominator A RHS formula to specify the logistic models for estimating the denominator

terms of the inverse probability of censoring weights.

pool\_models Fit pooled or separate censoring models for those treated and those untreated

at the immediately previous visit. Pooling can be specified for the models for the numerator and denominator terms of the inverse probability of censoring weights. One of "none", "numerator", or "both" (default is "none" except when

estimand = "ITT" then default is "numerator").

model\_fitter An object of class te\_model\_fitter which determines the method used for

fitting the weight models. For logistic regression use stats\_glm\_logit().

#### Value

object is returned with @censor\_weights set

#### **Examples**

```
trial_sequence("ITT") |>
```

set\_data

```
set_data(data = data_censored) |>
set_censor_weight_model(
  censor_event = "censored",
  numerator = ~ age_s + x1 + x3,
  denominator = ~ x3 + x4,
  pool_models = "both",
  model_fitter = stats_glm_logit(save_path = tempdir())
)
```

set\_data

Set the trial data

### **Description**

### [Experimental]

### Usage

```
set_data(object, data, ...)
## S4 method for signature 'trial_sequence_ITT,data.frame'
set_data(
  object,
  data,
  id = "id",
  period = "period",
  treatment = "treatment",
  outcome = "outcome",
  eligible = "eligible"
)
## S4 method for signature 'trial_sequence_AT,data.frame'
set_data(
 object,
  data,
  id = "id",
  period = "period",
  treatment = "treatment",
  outcome = "outcome",
  eligible = "eligible"
)
## S4 method for signature 'trial_sequence_PP,data.frame'
set_data(
  object,
  data,
  id = "id",
  period = "period",
```

set\_expansion\_options 31

```
treatment = "treatment",
outcome = "outcome",
eligible = "eligible"
)
```

### **Arguments**

object	A trial_sequence object
data	A data.frame containing all the required variables in the person-time format, i.e., the <u+2018>long<u+2019> format.</u+2019></u+2018>
	Other arguments used by methods internally.
id	Name of the variable for identifiers of the individuals. Default is $<$ U+2018 $>$ id $<$ U+2019 $>$ .
period	Name of the variable for the visit/period. Default is <u+2018>period<u+2019>.</u+2019></u+2018>
treatment	Name of the variable for the treatment indicator at that visit/period. Default is <u+2018>treatment<u+2019>.</u+2019></u+2018>
outcome	Name of the variable for the indicator of the outcome event at that visit/period. Default is <u+2018>outcome<u+2019>.</u+2019></u+2018>
eligible	Name of the variable for the indicator of eligibility for the target trial at that visit/period. Default is <u+2018>eligible<u+2019>.</u+2019></u+2018>

# Value

An updated trial\_sequence object with data

# **Examples**

```
data(trial_example)
trial_sequence("ITT") |>
  set_data(
    data = trial_example,
    id = "id",
    period = "period",
    eligible = "eligible",
    treatment = "treatment"
)
```

set\_expansion\_options Set expansion options

# Description

[Experimental]

#### Usage

```
set_expansion_options(object, ...)
## S4 method for signature 'trial_sequence_ITT'
set_expansion_options(
  object,
  output,
  chunk_size,
  first_period = 0,
 last_period = Inf
)
## S4 method for signature 'trial_sequence_PP'
set_expansion_options(
  object,
  output,
  chunk_size,
  first_period = 0,
  last_period = Inf
)
## S4 method for signature 'trial_sequence_ITT'
set_expansion_options(
  object,
 output,
  chunk_size,
  first_period = 0,
  last_period = Inf
)
```

### **Arguments**

object A trial\_sequence object
... Arguments used in methods

output A te\_datastore object as created by a save\_to\_\* function.

chunk\_size An integer specifying the number of patients to include in each expansion itera-

tion

first\_period An integer specifying the first period to include in the expansion last\_period An integer specifying the last period to include in the expansion

#### Value

object is returned with @expansion set

#### See Also

```
Other save_to: save_to_csv(), save_to_datatable(), save_to_duckdb()
```

set\_outcome\_model 33

#### **Examples**

```
output_dir <- file.path(tempdir(check = TRUE), "expanded_data")
ITT_trial <- trial_sequence("ITT") |>
    set_data(data = data_censored) |>
    set_expansion_options(output = save_to_csv(output_dir), chunk_size = 500)
# Delete directory
unlink(output_dir, recursive = TRUE)
```

set\_outcome\_model

Specify the outcome model

# Description

### [Experimental]

The time-to-event model for outcome is specified with this method. Any adjustment terms can be specified. For ITT and PP estimands the treatment\_var is not specified as it is automatically defined as assigned\_treatment. Importantly, the modelling of "time" is specified in this model with arguments for trial start time and follow up time within the trial.

#### Usage

```
set_outcome_model(object, ...)
## S4 method for signature 'trial_sequence'
set_outcome_model(
  object,
  treatment_var = \sim 0,
  adjustment\_terms = ~1,
  followup_time_terms = ~followup_time + I(followup_time^2),
  trial_period_terms = ~trial_period + I(trial_period^2),
 model_fitter = stats_glm_logit(save_path = NA)
)
## S4 method for signature 'trial_sequence_ITT'
set_outcome_model(
 object,
  adjustment\_terms = \sim 1,
  followup_time_terms = ~followup_time + I(followup_time^2),
  trial_period_terms = ~trial_period + I(trial_period^2),
 model_fitter = stats_glm_logit(save_path = NA)
)
## S4 method for signature 'trial_sequence_PP'
set_outcome_model(
  object,
  adjustment\_terms = ~1,
```

34 set\_outcome\_model

```
followup_time_terms = ~followup_time + I(followup_time^2),
    trial_period_terms = ~trial_period + I(trial_period^2),
    model_fitter = stats_glm_logit(save_path = NA)
)

## S4 method for signature 'trial_sequence_AT'
set_outcome_model(
    object,
    treatment_var = "dose",
    adjustment_terms = ~1,
    followup_time_terms = ~followup_time + I(followup_time^2),
    trial_period_terms = ~trial_period + I(trial_period^2),
    model_fitter = stats_glm_logit(save_path = NA)
)
```

### Arguments

```
object A trial_sequence object

... Parameters used by methods

treatment_var The treatment term, only used for "as treated" estimands. PP and ITT are fixed to use "assigned_treatment".

adjustment_terms
Formula terms for any covariates to adjust the outcome model.

followup_time_terms
Formula terms for followup_time, the time period relative to the start of the trial.

trial_period_terms
Formula terms for trial_period, the time period of the start of the trial.

model_fitter A te_model_fitter object, e.g. from stats_glm_logit().
```

#### Value

A modified object with the outcome\_model slot set

#### **Examples**

```
trial_sequence("ITT") |>
  set_data(data_censored) |>
  set_outcome_model(
   adjustment_terms = ~age_s,
   followup_time_terms = ~ stats::poly(followup_time, degree = 2)
)
```

```
set_switch_weight_model
```

Set switching weight model

#### **Description**

#### [Experimental]

### Usage

```
set_switch_weight_model(object, numerator, denominator, model_fitter, ...)
## S4 method for signature 'trial_sequence'
set_switch_weight_model(
  object,
  numerator,
  denominator,
 model_fitter,
  eligible_wts_0 = NULL,
  eligible_wts_1 = NULL
)
## S4 method for signature 'trial_sequence_ITT'
set_switch_weight_model(object, numerator, denominator, model_fitter)
```

# **Arguments**

object A trial\_sequence object.

Right hand side formula for the numerator model numerator Right hand side formula for the denominator model denominator A te\_model\_fitter object, such as stats\_glm\_logit model\_fitter

Other arguments used by methods.

eligible\_wts\_0 Name of column containing indicator (0/1) for observation to be excluded/included

in weight model.

eligible\_wts\_1 Exclude some observations when fitting the models for the inverse probability of treatment weights. For example, if it is assumed that an individual will stay on treatment for at least 2 visits, the first 2 visits after treatment initiation by definition have a probability of staying on the treatment of 1.0 and should thus be excluded from the weight models for those who are on treatment at the immediately previous visit. Users can define a variable that indicates that these 2 observations are ineligible for the weight model for those who are on treatment at the immediately previous visit and add the variable name in the argument eligible\_wts\_1. Similar definitions are applied to eligible\_wts\_0 for excluding observations when fitting the models for the inverse probability of treatment weights for those who are not on treatment at the immediately previous visit.

36 stats\_glm\_logit

### Value

object is returned with @switch\_weights set

### **Examples**

```
trial_sequence("PP") |>
  set_data(data = data_censored) |>
  set_switch_weight_model(
    numerator = ~ age_s + x1 + x3,
    denominator = ~ x3 + x4,
    model_fitter = stats_glm_logit(tempdir())
)
```

show\_weight\_models

Show Weight Model Summaries

# Description

### [Experimental]

### Usage

```
show_weight_models(object)
```

#### **Arguments**

object

A trial\_sequence object after fitting weight models with calculate\_weights()

#### Value

Prints summaries of the censoring models

 $stats\_glm\_logit$ 

Fit outcome models using stats::glm

# Description

### [Experimental]

### Usage

```
stats_glm_logit(save_path)
```

### **Arguments**

save\_path

Directory to save models. Set to NA if models should not be saved.

#### **Details**

Specify that the pooled logistic regression outcome models should be fit using stats::glm with family = binomial(link = "logit").

Outcome models additional calculate robust variance estimates using sandwich::vcovCL.

#### Value

An object of class te\_stats\_glm\_logit inheriting from te\_model\_fitter which is used for dispatching methods for the fitting models.

#### See Also

```
Other model_fitter: parsnip_model(), te_model_fitter-class
```

## **Examples**

```
stats_glm_logit(save_path = tempdir())
```

```
summary.TE_data_prep Summary methods
```

#### **Description**

[Stable] Print summaries of data and model objects produced by TrialEmulation.

#### Usage

```
## S3 method for class 'TE_data_prep'
summary(object, ...)
## S3 method for class 'TE_data_prep_sep'
summary(object, ...)
## S3 method for class 'TE_data_prep_dt'
summary(object, ...)
## S3 method for class 'TE_msm'
summary(object, ...)
## S3 method for class 'TE_robust'
summary(object, ...)
```

## **Arguments**

```
objectObject to print summaryAdditional arguments passed to print methods.
```

38 te\_datastore-class

# Value

No value, displays summaries of object.

te\_data-class

TrialEmulation Data Class

# **Description**

TrialEmulation Data Class

# **Slots**

data A data.table object with columns "id", "period", "treatment", "outcome", "eligible"

te\_datastore-class

te\_datastore

# Description

This is the parent class for classes which define how the expanded trial data should be stored. To define a new storage type, a new class should be defined which inherits from te\_datastore. In addition, methods save\_expanded\_data and read\_expanded\_data need to be defined for the new class.

# Value

A 'te\_datastore' object

# **Slots**

N The number of observations in this data. Initially 0.

te\_data\_ex 39

te\_data\_ex

Example of a prepared data object

#### **Description**

A small example object from data\_preparation used in examples. It is created with the following code:

## Usage

```
te_data_ex
```

#### **Format**

An object of class TE\_data\_prep\_dt (inherits from TE\_data\_prep) of length 6.

# **Details**

```
dat <- trial_example[trial_example$id < 200, ]

te_data_ex <- data_preparation(
data = dat,
  outcome_cov = c("nvarA", "catvarA"),
  first_period = 260,
  last_period = 280
)</pre>
```

# See Also

```
te_model_ex
```

te\_model\_ex

Example of a fitted marginal structural model object

# **Description**

A small example object from trial\_msm used in examples. It is created with the following code:

# Usage

```
te_model_ex
```

#### Format

An object of class TE\_msm of length 3.

40 te\_outcome\_data-class

#### **Details**

```
te_model_ex <- trial_msm(
  data = data_subset,
  outcome_cov = c("catvarA", "nvarA"),
  last_followup = 40,
  model_var = "assigned_treatment",
  include_followup_time = ~followup_time,
  include_trial_period = ~trial_period,
  use_sample_weights = FALSE,
  quiet = TRUE,
  glm_function = "glm"
)</pre>
```

#### See Also

```
te_data_ex
```

#### Description

This is a virtual class which other outcome model fitter classes should inherit from. Objects of these class exist to define how the outcome models are fit. They are used for the dispatch of the internal methods fit\_outcome\_model, fit\_weights\_model and predict.

## See Also

```
Other model_fitter: parsnip_model(), stats_glm_logit()
```

# **Description**

TrialEmulation Outcome Data Class

#### **Slots**

```
data A data.table object with columns "id", "period",
n_rows Number of rows
n_ids Number of IDs
periods Vector of periods "treatment", "outcome", "eligible"
```

te\_outcome\_fitted-class

Fitted Outcome Model Object

#### **Description**

Fitted Outcome Model Object

#### **Slots**

```
model list containing fitted model objects.

summary list of data.frames. Tidy model summaries a la broom() and glance()
```

te\_outcome\_model-class

Fitted Outcome Model Object

# Description

Fitted Outcome Model Object

#### **Slots**

formula formula object for the model fitting

adjustment\_vars character. Adjustment variables

treatment\_var Variable used for treatment

stabilised\_weights\_terms formula. Adjustment terms from numerator models of stabilised weights. These must be included in the outcome model.

adjustment\_terms formula. User specified terms to include in the outcome model

treatment\_terms formula. Estimand defined treatment term

followup\_time\_terms formula. Terms to model follow up time within an emulated trial

trial\_period\_terms formula. Terms to model start time ("trial\_period") of an emulated trial

model\_fitter Model fitter object

fitted list. Saves the model objects

42 trial\_msm

trial\_example

Example of longitudinal data for sequential trial emulation

# **Description**

A dataset containing the treatment, outcomes and other attributes of 503 patients for sequential trial emulation. See vignette("Getting-Started").

# Usage

```
trial_example
```

#### **Format**

A data frame with 48400 rows and 11 variables:

id patient identifier

eligible eligible for trial start in this period, 1=yes, 0=no

period time period

outcome indicator for outcome in this period, 1=event occurred, 0=no event

**treatment** indicator for receiving treatment in this period, 1=treatment, 0=no treatment

catvarA A categorical variable relating to treatment and the outcome

catvarB A categorical variable relating to treatment and the outcome

catvarC A categorical variable relating to treatment and the outcome

nvarA A numerical variable relating to treatment and the outcome

**nvarB** A numerical variable relating to treatment and the outcome

nvarC A numerical variable relating to treatment and the outcome

trial\_msm

Fit the marginal structural model for the sequence of emulated trials

# **Description**

[Stable]

trial\_msm 43

## Usage

```
trial_msm(
  data,
  outcome\_cov = ~1,
  estimand_type = c("ITT", "PP", "As-Treated"),
 model_var = NULL,
  first_followup = NA,
  last_followup = NA,
  analysis_weights = c("asis", "unweighted", "p99", "weight_limits"),
  weight_limits = c(0, Inf),
  include_followup_time = ~followup_time + I(followup_time^2),
  include_trial_period = ~trial_period + I(trial_period^2),
  where_case = NA,
  glm_function = c("glm", "parglm"),
  use_sample_weights = TRUE,
  quiet = FALSE,
)
```

#### **Arguments**

data

A data.frame containing all the required variables in the person-time format, i.e., the 'long' format.

outcome\_cov

A RHS formula with baseline covariates to be adjusted for in the marginal structural model for the emulated trials. Note that if a time-varying covariate is specified in outcome\_cov, only its value at each of the trial baselines will be included in the expanded data.

estimand\_type

Specify the estimand for the causal analyses in the sequence of emulated trials. estimand\_type = "ITT" will perform intention-to-treat analyses, where treatment switching after trial baselines are ignored. estimand\_type = "PP" will perform per-protocol analyses, where individuals' follow-ups are artificially censored and inverse probability of treatment weighting is applied. estimand\_type = "As-Treated" will fit a standard marginal structural model for all possible treatment sequences, where individuals' follow-ups are not artificially censored but treatment switching after trial baselines are accounted for by applying inverse probability of treatment weighting.

model\_var

Treatment variables to be included in the marginal structural model for the emulated trials. model\_var = "assigned\_treatment" will create a variable assigned\_treatment that is the assigned treatment at the trial baseline, typically used for ITT and per-protocol analyses. model\_var = "dose" will create a variable dose that is the cumulative number of treatments received since the trial baseline, typically used in as-treated analyses.

first\_followup

First follow-up time/visit in the trials to be included in the marginal structural model for the outcome event.

last\_followup

Last follow-up time/visit in the trials to be included in the marginal structural model for the outcome event.

44 trial\_msm

analysis\_weights

Choose which type of weights to be used for fitting the marginal structural model for the outcome event.

- "asis": use the weights as calculated.
- "p99": use weights truncated at the 1st and 99th percentiles (based on the distribution of weights in the entire sample).
- "weight\_limits": use weights truncated at the values specified in weight\_limits.
- "unweighted": set all analysis weights to 1, even if treatment weights or censoring weights were calculated.

weight\_limits Lower and upper limits to truncate weights, given as c(lower, upper)
include\_followup\_time

The model to include the follow up time/visit of the trial (followup\_time) in the marginal structural model, specified as a RHS formula.

include\_trial\_period

The model to include the trial period (trial\_period) in the marginal structural model, specified as a RHS formula.

where\_case

Define conditions using variables specified in where\_var when fitting a marginal structural model for a subgroup of the individuals. For example, if where\_var= "age", where\_case = "age >= 30" will only fit the marginal structural model to the subgroup of individuals. who are 30 years old or above.

glm\_function

Specify which glm function to use for the marginal structural model from the stats or parglm packages. The default function is the glm function in the stats package. Users can also specify glm\_function = "parglm" such that the parglm function in the parglm package can be used for fitting generalized linear models in parallel. The default control setting for parglm is nthreads = 4 and method = "FAST", where four cores and Fisher information are used for faster computation. Users can change the default control setting by passing the arguments nthreads and method in the parglm.control function of the parglm package, or alternatively, by passing a control argument with a list produced by parglm.control(nthreads = , method = ).

use\_sample\_weights

Use case-control sampling weights in addition to inverse probability weights for treatment and censoring. data must contain a column sample\_weight. The final weights used in the pooled logistic regression are calculated as weight = weight \* sample\_weight.

quiet

Suppress the printing of progress messages and summaries of the fitted models.

. . .

Additional arguments passed to glm\_function. This may be used to specify initial values of parameters or arguments to control. See stats::glm, parglm::parglm and parglm::parglm.control() for more information.

#### **Details**

Apply a weighted pooled logistic regression to fit the marginal structural model for the sequence of emulated trials and calculates the robust covariance matrix of parameter using the sandwich estimator.

The model formula is constructed by combining the arguments outcome\_cov, model\_var, include\_followup\_time, and include\_trial\_period.

trial\_sequence 45

# Value

Object of class TE\_msm containing

```
model a glm object
```

robust a list containing a summary table of estimated regression coefficients and the robust covariance matrix

args a list contain the parameters used to prepare and fit the model

trial\_sequence

Create a sequence of emulated target trials object

# Description

# [Experimental]

#### Usage

```
trial_sequence(estimand, ...)
```

# **Arguments**

estimand The name of the estimand for this analysis, either one of "ITT", "PP", "AT" for intention-to-treat, per-protocol, as-treated estimands respectively, or the name

of a class extending trial\_sequence

... Other parameters used when creating object

#### Value

An estimand specific trial sequence object

# **Examples**

```
trial_sequence("ITT")
```

46 vignette\_switch\_data

#### **Description**

Trial Sequence class

#### Slots

```
data te_data.

estimand character. Descriptive name of estimand.

expansion te_expansion

outcome_model te_outcome_model.

outcome_data te_outcome_data.

censor_weight te_weight. Object to define weighting to account for informative censoring censor_weight te_weight. Object to define weighting to account for informative censoring due to treatment switching
```

## **Description**

This is the expanded dataset created in the vignette ("Getting-Started") known as switch\_data.

# Usage

```
vignette_switch_data
```

#### **Format**

A data frame with 1939053 rows and 7 variables:

id patient identifier

trial\_period trial start time period

followup\_time follow up time within trial

outcome indicator for outcome in this period, 1=event occurred, 0=no event

treatment indicator for receiving treatment in this period, 1=treatment, 0=non-treatment

assigned\_treatment indicator for assigned treatment at baseline of the trial, 1=treatment, 0=nontreatment

weight weights for use with model fitting

catvarA A categorical variable relating to treatment and the outcome

catvarB A categorical variable relating to treatment and the outcome

catvarC A categorical variable relating to treatment and the outcome

nvarA A numerical variable relating to treatment and the outcome

nvarB A numerical variable relating to treatment and the outcome

nvarC A numerical variable relating to treatment and the outcome

```
weight_model_data_indices
```

Data used in weight model fitting

# Description

## [Experimental]

## Usage

```
weight_model_data_indices(
  object,
  type = c("switch", "censor"),
  model,
  set_col = NULL
)
```

#### **Arguments**

object A trial\_sequence object

type Select a censoring or switching model

model The model name

set\_col A character string to specifying a new column to contain indicators for observa-

tions used in fitting this model.

#### Value

If set\_col is not specified a logical data.table column is returned. Otherwise

# Examples

```
trial_pp <- trial_sequence("PP") |>
  set_data(data_censored) |>
  set_switch_weight_model(
    numerator = ~age,
    denominator = ~ age + x1 + x3,
    model_fitter = stats_glm_logit(tempdir())
  ) |>
  calculate_weights()
ipw_data(trial_pp)
```

```
show_weight_models(trial_pp)

# get logical column for own processing
i <- weight_model_data_indices(trial_pp, "switch", "d0")

# set column in data
weight_model_data_indices(trial_pp, "switch", "d0", set_col = "sw_d0")
weight_model_data_indices(trial_pp, "switch", "d1", set_col = "sw_d1")
ipw_data(trial_pp)</pre>
```

# **Index**

* datasets	ipw_data, 16
data_censored, 5	ipw_data,trial_sequence-method
te_data_ex, 39	(ipw_data), 16
te_model_ex, 39	ipw_data<- (ipw_data), 16
trial_example,42	ipw_data<-,trial_sequence-method
vignette_switch_data,46	(ipw_data), 16
* model_fitter	
parsnip_model, 19	load_expanded_data, 17
stats_glm_logit, 36	<pre>load_expanded_data,trial_sequence-method</pre>
te_model_fitter-class,40	(load_expanded_data), 17
* save_to	
save_to_csv, 26	outcome_data, 18
save_to_datatable, 27	outcome_data,trial_sequence-method
save_to_duckdb, 27	(outcome_data), 18
set_expansion_options, 31	outcome_data<- (outcome_data), 18
	outcome_data<-,trial_sequence-method
calculate_weights, 3	(outcome_data), 18
calculate_weights(), 36	
calculate_weights,trial_sequence_AT-method	parglm::parglm, 8, 15, 44
(calculate_weights), 3	parglm::parglm.control(), 8, 15, 44
<pre>calculate_weights,trial_sequence_ITT-method</pre>	parsnip_model, 19, 37, 40
(calculate_weights), 3	predict, 40
calculate_weights,trial_sequence_PP-method	<pre>predict (predict_marginal), 20</pre>
(calculate_weights), 3	<pre>predict,trial_sequence_ITT-method</pre>
<pre>case_control_sampling_trials, 4, 9</pre>	(predict_marginal), $20$
data_censored, 5	<pre>predict,trial_sequence_PP-method</pre>
data_preparation, 6, 39	(predict_marginal), $20$
data_preparation(), $4$ , $15$	<pre>predict.TE_msm(predict_marginal), 20</pre>
$data_preparation(j, 4, 13)$	predict_marginal, 20
expand_trials, 9, 25	print.data.frame, 23
expand_trials(), 10	print.TE_weight_summary, 23
$fit_msm, 10$	read_expanded_data, 23
<pre>fit_msm,trial_sequence-method</pre>	read_expanded_data,te_datastore_datatable-method
(fit_msm), 10	<pre>(read_expanded_data), 23</pre>
<pre>fit_outcome_model, 40</pre>	
fit_weights_model, 11, 40	sample_expanded_data, 24
	<pre>sample_expanded_data,te_datastore-method</pre>
initiators, 12	<pre>(sample_expanded_data), 24</pre>
initiators(), 21	save_expanded_data, 25, 38

50 INDEX

```
save_expanded_data,te_datastore_datatable-metsodset(),4
        (save_expanded_data), 25
                                                  summary.TE_data_prep, 37
save_to_csv, 26, 27, 32
                                                  summary.TE_data_prep_dt
save_to_datatable, 26, 27, 27, 32
                                                           (summary.TE_data_prep), 37
save_to_duckdb, 26, 27, 27, 32
                                                  summary.TE_data_prep_sep
                                                           (summary.TE_data_prep), 37
set_censor_weight_model, 28
\verb|set_censor_weight_model,trial_sequence-methods ummary.TE_msm (summary.TE_data_prep)|, \\
        (set_censor_weight_model), 28
\verb|set_censor_weight_model,trial_sequence_AT-met| \verb|Mwammary.TE_robust| \\
                                                           (summary.TE_data_prep), 37
        (set_censor_weight_model), 28
set_censor_weight_model,trial_sequence_ITT-method
                                                  te_data-class, 38
        (set_censor_weight_model), 28
\verb|set_censor_weight_model|, \verb|trial_sequence_PP-method| \\ |data_ex, 39, 40| \\
                                                  te_datastore, 17, 23, 25, 32
        (set_censor_weight_model), 28
                                                  te_datastore-class, 38
set_data, 30
                                                  te_datastore_csv, 26
set_data(), 16
                                                  te_datastore_duckdb, 27
set_data,trial_sequence_AT,data.frame-method
                                                  te_model_ex, 39, 39
        (set_data), 30
\verb|set_data|, \verb|trial_sequence_ITT|, \verb|data.frame-method| \\ \texttt{te_model_fitter}, \verb|20|, 35|, 37| \\
        (set_data), 30
                                                  te_outcome_data, 18
set\_data, trial\_sequence\_PP, data.frame-method
                                                  te_outcome_data-class, 40
        (set_data), 30
                                                  te_outcome_fitted-class, 41
set_expansion_options, 25-27, 31
                                                  te_outcome_model-class, 41
set_expansion_options(), 9
                                                  trial_example, 42
set_expansion_options,trial_sequence_ITT-method trial_msm, 9, 39, 42
        (set_expansion_options), 31
                                                  trial_msm(), 5, 15, 21
set_expansion_options,trial_sequence_PP-method trial_sequence, 3, 9, 17, 18, 21, 31, 32, 35,
        (set_expansion_options), 31
                                                           36, 45, 45, 47
set_outcome_model, 10, 33
                                                  trial_sequence-class, 46
set_outcome_model(), 10
                                                  trial_sequence_AT-class
set_outcome_model,trial_sequence-method
                                                           (trial_sequence-class), 46
        (set_outcome_model), 33
                                                  trial_sequence_ITT-class
set_outcome_model,trial_sequence_AT-method
                                                           (trial_sequence-class), 46
        (set_outcome_model), 33
                                                  trial_sequence_PP-class
set_outcome_model,trial_sequence_ITT-method
                                                           (trial_sequence-class), 46
        (set_outcome_model), 33
set_outcome_model,trial_sequence_PP-method
                                                  vignette_switch_data, 46
        (set_outcome_model), 33
                                                  weight_model_data_indices, 47
set_switch_weight_model, 35
set_switch_weight_model,trial_sequence-method
        (set_switch_weight_model), 35
set_switch_weight_model, trial_sequence_ITT-method
        (set_switch_weight_model), 35
show_weight_models, 36
split(), 5
stats::glm, 8, 15, 37, 44
stats_glm_logit, 20, 35, 36, 40
stats_glm_logit(), 20, 29
```