# Package 'RCT'

January 20, 2025

**Title** Assign Treatments, Power Calculations, Balances, Impact Evaluation of Experiments

## Version 1.2

**Description** Assists in the whole process of designing and evaluating Randomized Control Trials.

Robust treatment assignment by strata/blocks, that handles misfits;

Power calculations of the minimum detectable treatment effect or minimum populations;

Balance tables of T-test of covariates;

Balance Regression: (treatment ~ all x variables) with F-test of null model;

 $Impact\_evaluation: Impact\ evaluation\ regressions.\ This\ function$ 

gives you the option to include control\_vars, fixed effect variables,

cluster variables (for robust SE), multiple endogenous variables and multiple heterogeneous variables (to test treatment effect heterogeneity)

multiple heterogeneous variables (to test treatment effect heterogeneity)

summary\_statistics: Function that creates a summary statistics table with statistics rank observations in n groups: Creates a factor variable with n groups. Each group has a min and max label attach to each category.

Athey, Susan, and Guido W. Imbens (2017) <arXiv:1607.00698>.

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## **Encoding UTF-8**

**Imports** dplyr, purrr, glue, rlang, tidyr, stringr, MASS, pracma, estimatr, broom (>= 1.0.0), forcats, magrittr, ggplot2, utils, tidyselect (>= 1.0.0)

Suggests knitr, rmarkdown, testthat

RoxygenNote 7.3.1

VignetteBuilder knitr

NeedsCompilation no

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# Description

balance\_regression() Runs a LPM of treatment status against all covariates (treatment~X'B).

## Usage

```
balance_regression(data, treatment)
```

## **Arguments**

data A data.frame, tibble or data.table treatment a string with treatment status column

## **Details**

This functions runs a Linear Probability model of each treatment group & control on all the columns in data. For instance, if treatment column has values of (0,1,2), balance\_regression will run two models: 1) LPM(treatment(0,1)~X'b) and 2) LPM(treatment(0,2)~X'b). The value are the regression tables and details of the F\_test of these models.

# Value

A list: regression\_tables = regression output of treatment against all covariates, F\_test = table with the F tests of each regression

```
data <-data.frame(x1 = rnorm(n = 100, mean = 100, sd = 15), x2= rnorm(n = 100, mean = 65), treat = rep(c(0,1,2,3,4), each = 20)) balance_regression(data = data, treatment = "treat")
```

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balance\_table

Creates balance table for the X variables across treatment status

## **Description**

Creates balance table for the X variables across treatment status

## Usage

```
balance_table(data, treatment)
```

## **Arguments**

data A data.frame, tibble or data.table
treatment a string with treatment status column

#### **Details**

balance\_table() performs t.test(X~treatment) for each X column in data. Every value of treatment i.e 1,2,3,...N is compared against control value (0) or the first value of the treatment column. For instance, If treatment column has values of (0,1,2,3), balance\_table will return: the mean value of each treatment (for all X's), and the p\_values of the t.test of (1,2,3) against treatment = 0.

## Value

A tibble with Mean\_value of each treatment status and p\_values

# **Examples**

impact\_eval

Impact Evaluation of Treatment Effects

## **Description**

Impact Evaluation of Treatment Effects

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## Usage

```
impact_eval(
  data,
  endogenous_vars,
  treatment,
  heterogenous_vars,
  fixed_effect_vars = NULL,
  control_vars,
  cluster_var
)
```

## **Arguments**

data A data.frame, tibble or data.table

endogenous\_vars

Vector of Y's on which treatment effects will be evaluated

treatment Variable indicating the treatment status

heterogenous\_vars

Vector of variables for which you wish to assess treatment distributions/heterogeneities.

fixed\_effect\_vars

Vector of variables to add as fixed effects. Default is without fixed effects

control\_vars Vector of variables to control for in the evaluation. Default is without controls

cluster\_var String of cluster variable the standard errors. Default is without clustered std

errors

## **Details**

This function carries out the evaluation of treatment effects on endogenous variables. It automatically runs the regressions of all the endogenous\_vars supplied & all the combinations of endogenous\_vars and heterogenous\_vars. Additionally, the function has the option of include fixed\_effects, controls and cluster variables for clustered std errors.

## Value

impact\_eval() returns a list of regression tables. The names of the list are the same as the endogenous variables. for heterogeneities the names are endogenous\_var\_heterogenous\_var

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```
endogenous_vars = c("y_1", "y_2"),
    treatment = "treat",
    heterogenous_vars = c("heterogenous_var1"),
cluster_var = "cluster_var1", fixed_effect_vars = c("fixed_effect_var1"),
    control_vars = c("control_var1"))
```

ntile\_label

ntile\_label() ranks observations in n groups, with labels

## **Description**

ntile\_label() ranks observations in n groups, with labels

# Usage

```
ntile_label(var, n, digits = 0)
```

# Arguments

rank the variable in n groups

digits

How many digits to include in the label

#### **Details**

 $n\_tile\_label$  is very similar to ntile from dplyr. But  $n\_tile\_label$  creates the n groups and then labels them. For each group i, the value of the  $ntile\_label$  is [min(i) - max(i)].

## Value

A ordered factor vector of each n group. The value has the form of  $[min(n_i) - max(n_i)]$ 

```
data <- data.frame(y_1 = rbinom(n = 100, size = 1, prob = 0.3),

y_2 = rnorm(n = 100, mean = 8, sd = 2))

data$y_1_2 <- ntile_label(data$y_1, n = 2, digits = 0)

data$y_2_4 <- ntile_label(data$y_2, n = 4, digits = 1)
```

6 N\_min

N_min	N_min() computes the minimum population needed to detect difference between control group and each treatment, given a target minimum detectable effect

## **Description**

N\_min() computes the minimum population needed to detect difference between control group and each treatment, given a target minimum detectable effect

## Usage

```
N_min(
  outcome_var,
  tau_min,
  power = 0.8,
  significance = 0.05,
  share_control,
  n_groups = 2
)
```

#### **Arguments**

outcome\_var the variable for which you wish to test the impact of treatment tau\_min the target detectable effect (in outcome\_var units)

power The level of power of the test (1 - Pr(Reject H\_0 | H\_0 True)). Default is 0.8

significance The level of significance of the test Pr(Reject H\_0 | H\_0 False). Default is 0.05

share\_control The share of observations in N assigned to control. This argument allows for sequences (i.e. seq(0,1,0.1))

Number of groups (control + # treatment groups)

## **Details**

n\_groups

This function calculates the minimum experiment's population needed in order to detect at least a difference of tau\_min statistically significantly. This is between any two given groups (e.g. control vs each treatment), given the outcome variable, power and significance

# Value

A tibble with the share\_control and N observations in control group (N\_control), the share and N of each treatment c(share\_ti, N\_ti), total share of treatment rows and N treated (share\_treat, N\_treat), N, the minimum detectable difference between control and all treatments together (tau\_min\_global), the minimum detectable difference between control and each treatment (tau\_min\_each\_treat)

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## **Examples**

```
data <- data.frame(y_1 = rbinom(n = 100, size = 1, prob = 0.3),

y_2 = rnorm(n = 100, mean = 8, sd = 2))

N_min(data$y_1, tau_min = 0.01, share_control = seq(0,1,0.1), n_groups = 3)
```

**RCT** 

Designing, random assigning and evaluating Randomized Control Trials

## **Description**

RCT provides three important group of functions: a) functions for pre-processing the design of the RCT b) Functions for assigning treatment status and checking for balances c) Function for evaluating the impact of the RCT

#### **Details**

RCT helps you focus on the statistics of the randomized control trials, rather than the heavy programming lifting. RCT helps you in the whole process of designing and evaluating a RCT. 1. Clean and summarise the data in which you want to randomly assign treatment 2. Decide the share of observations that will go to control group 3. Decide which variables to use for strata building 4. Robust Random Assignment by strata/blocks 5 Impact evaluation of all y's and heterogeneities To lean more about RCT, start with the vignette: browseVignettes(package = "RCT")

#### **RCT functions**

treatment\_assign: Robust treatment assign by strata/blocks

impact\_eval: Automatized impact evaluation with heterogeneous treatment effects

balance\_table: Balance tables for any length of covariates

balance\_regression: LPM of treatment status against covariates with F-test

tau\_min: Computation of the minimum detectable effect between control & treatment units

tau\_min\_probability: Computation of the minimum detectable effect between control & treatment

units for dichotomous y-vars

summary\_statistics: Summary statistics of all numeric columns in your data

ntile\_label: Rank and divide observations in n groups, with label

## Author(s)

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## References

Athey, Susan, and Guido W. Imbens (2017) "The Econometrics Randomized Experiments". Handbook of economic field experiments. https://arxiv.org/abs/1607.00698

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## See Also

Useful links: https://github.com/isidorogu/RCT Report bugs at https://github.com/isidorogu/RCT/issues

 $summary\_statistics \qquad summary\_statistics() \ Creates \ summary \ statistics \ table \ of \ all \ numeric \\ variables \ in \ data$ 

# Description

summary\_statistics() Creates summary statistics table of all numeric variables in data

## Usage

```
summary_statistics(
  data,
  probs = c(0, 0.05, 0.1, 0.25, 0.5, 0.75, 0.9, 0.95, 1),
  na.rm = T
)
```

# Arguments

data A data.frame, tibble or data.table probs The quantiles to compute. Default is c(0, 0.05, 0.1, 0.25, 0.5, 0.75, 0.9, 0.95, 1) whether to exclude NA's from calculations

# **Details**

This function computes the selected quantiles, mean and N values of all the numeric columns of data.

## Value

A tibble with the Mean, N (not NA) and probs selects for each numeric column

```
data <-data.frame(x = c(1:5), y = c(100, 200, 300, 410, 540), z = rep("c", 5)) summary_statistics(data)
```

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tau_min	tau_min() computes the minimum detectable difference between control group and each treatment

## **Description**

tau\_min() computes the minimum detectable difference between control group and each treatment

## Usage

```
tau_min(
  outcome_var,
  N,
  power = 0.8,
  significance = 0.05,
  share_control,
  n_groups = 2
)
```

## **Arguments**

outcome_var	the variable for which you wish to test the impact of treatment
N	number of observations in the RCT, usually nrow(data)
power	The level of power of the test (1 - Pr(Reject H_0   H_0 True) ). Default is 0.8
significance	The level of significance of the test Pr(Reject H_0   H_0 False). Default is 0.05
share_control	The share of observations in N assigned to control. This argument allows for sequences (i.e. $seq(0,1,0.1)$ )
n_groups	Number of groups (control + # treatment groups)

#### **Details**

This function calculates the minimum difference that could show significant  $E[Y(1)-Y(0)] = \tan$ , between any two given groups (e.g. control vs each treatment), given the population size (N), the outcome variable, power and significance

#### Value

A tibble with the share\_control and N observations in control group (N\_control), the share and N of each treatment c(share\_ti, N\_ti), total share of treatment rows and N treated (share\_treat, N\_treat), N, the minimum detectable difference between control and all treatments together (tau\_min\_global), the minimum detectable difference between control and each treatment (tau\_min\_each\_treat)

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tau\_min\_probability tau\_min\_probability() computes the minimum detectable difference between control group and each treatment for a dichotomous variable

# Description

tau\_min\_probability() computes the minimum detectable difference between control group and each treatment for a dichotomous variable

## Usage

```
tau_min_probability(
  prior,
  N,
  power = 0.8,
  significance = 0.05,
  share_control,
  n_groups = 2
)
```

## **Arguments**

prior Pr(Y=1).

N number of observations in the RCT, usually nrow(data)

power The level of power of the test (1 - Pr(Reject H\_0 | H\_0 True)). Default is 0.8

significance The level of significance of the test Pr(Reject H\_0 | H\_0 False). Default is 0.05

share\_control The share of observations in N assigned to control. This argument allows for sequences (i.e. seq(0,1,0.1))

n\_groups Number of groups (control + # treatment groups)

#### Details

This function calculates the minimum difference that could show significant Pr[Y(1)-Y(0)] = tau, between any two given groups (e.g. control vs each treatment), given the population size (N), the outcome variable, power and significance

#### Value

A tibble with the share\_control and N observations in control group (N\_control), the share and N of each treatment c(share\_ti, N\_ti), total share of treatment rows and N treated (share\_treat, N\_treat), N, the minimum detectable difference between control and all treatments together (tau\_min\_global), the minimum detectable difference between control and each treatment (tau\_min\_each\_treat)

```
tau_min_probability(0.4, N = 1000, share_control = seq(0,1,0.1), n_groups = 3)
```

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treatment_assign	treatment_assign()	carries	out	robust	treatment	assignment	by
	strata/blocks						

## **Description**

treatment\_assign() carries out robust treatment assignment by strata/blocks

# Usage

```
treatment_assign(
  data,
  share_control,
  n_t = 2,
  strata_varlist,
  missfits = c("global", "NA", "strata"),
  seed = 1990,
  share_ti = rep(1/n_t - share_control/n_t, times = n_t),
  key
)
```

## Arguments

data A data.frame, tibble or data.table share of the observations assigned to control group share\_control Number of treatments groups n t strata\_varlist vector of categorical variables to form the strata/blocks for random assignment. Should be in the form of vars(var1, var2, ...) missfits How to handle the misfits. Default is "global". See Carril (2016) for details. seed A number used to set.seed(). The share of each treatment group. If NULL (Default), each treatment group share\_ti will have equal share. The key identifier column of data. key

#### **Details**

This function creates a variable that indicates the treatment status. The random assignment is made by strata/blocks. It can handle equal or unequal treatment shares. Finally, it has three methods available to handle misfits (same as randtreat in STATA): "global": assigning the observations that couldn't be randomly assigned globally, "strata": assigning the observations that couldn't be randomly assigned by strata, "NA": set the treat observations that couldn't be randomly assigned to NA.

#### Value

A list: "data" = the data with key, treat, strata, misfit column., "summary\_strata" = A summary tibble with the membership of each strata and its size.

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