# Package: FSM (via r-universe)

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Title Finite Selection Model

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<b>Description</b> Randomized and balanced allocation of units to treatment groups using the Finite Selection Model (FSM). The FSM was originally proposed and developed at the RAND corporation by Carl Morris to enhance the experimental design for the now famous Health Insurance Experiment. See Morris (1979) <doi:10.1016 0304-4076(79)90053-8=""> for details on the original version of the FSM.</doi:10.1016>
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crd

Completely Randomized Design (CRD)

### **Description**

Generates an assignment under completely randomized design (CRD).

### Usage

```
crd(data_frame, n_treat, treat_sizes, control = FALSE)
```

### Arguments

data\_frame A data frame corresponding to the full sample of units.

n\_treat Number of treatment groups.

treat\_sizes A vector of treatment group sizes. If control = TRUE, the first element of treat\_sizes should be the control group size.

control If TRUE, treatments are labeled as 0,1,...,g-1 (0 representing the control group). If FALSE, they are labeled as 1,2,...,g.

The original data frame augmented with the column of the treatment indicator.

### Author(s)

Value

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

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### **Examples**

```
# Consider N = 12, n1 = n2 = n3 = 4. 
df_sample = data.frame(index = 1:12, x = c(20,30,40,40,50,60,20,30,40,40,50,60)) # Draw a random assignment from CRD. 
fc = crd(data_frame = df_sample, n_treat = 3, treat_sizes = c(4,4,4)) # Get vector of treatment assignments. 
Z_crd = fc$Treat
```

ess\_model

Model-based Effective Sample Size (ESS)

### Description

Computes the model-based effective sample size (ESS) of a collection of assignments under a given set of potential outcomes.

### Usage

```
ess_model(X_cov, assign_matrix, Y_mat, contrast = c(1, -1))
```

### **Arguments**

X_cov	A matrix of covariates or transformations thereof that will be used as explanatory variables in the linear outcome models within each treatment group.
assign_matrix	A matrix containing a collection of treatment assignment vectors, each column containing a particular assignment vector.
Y_mat	A matrix of potential outcomes, where rows represent units and columns represent treatment levels (ordered).
contrast	A vector of the coefficients of the treatment contrast of interest. For example, for estimating the average treatment effect of treatment 1 versus treatment 2, $contrast = c(1,-1)$ .

### Value

A vector of effective sample sizes for the given collection of assignments.

#### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

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### **Examples**

```
# Consider the Lalonde dataset.
# Get the full sample size.
N = nrow(Lalonde)
# Get the treatment group sizes.
n1 = floor(N/2)
n2 = N-n1
# Generate an SOM.
som_obs = som(n_treat = 2, treat_sizes = c(n1,n2),include_discard = FALSE,
method = 'SCOMARS', marginal_treat = rep((n2/N),N), control = FALSE)
# Generate a treatment assignment given som_obs.
f = fsm(data_frame = Lalonde, SOM = som_obs, s_function = 'Dopt', eps = 0.0001,
ties = 'random', intercept = TRUE, standardize = TRUE, units_print = FALSE)
# Get assignment vector under the FSM.
Z_fsm_obs = f$data_frame_allocated$Treat
# Draw a random CRD.
Z_crd_obs = crd(data_frame = Lalonde, n_treat = 2, treat_sizes = c(n1, n2),
control = FALSE)$Treat
Z_big = cbind(Z_crd_obs, Z_fsm_obs)
# Generate the potential outcomes.
Y_1 = 100 - Lalonde$Age + 6 * Lalonde$Education - 20 * Lalonde$Black +
20 * Lalonde$Hispanic + 0.003 * Lalonde$Re75 + rnorm(N,0,4)
Y_1 = round(Y_1, 2)
# Set unit level causal effect = tau = 0.
tau = 0
Y_2 = Y_1 + tau
# Get the matrix of potential outcomes.
Y_appended = cbind(Y_1, Y_2)
# Get the matrix of covariates.
X_{cov} = Lalonde[,-1]
ess = ess_model(X_cov = X_cov, assign_matrix = Z_big, Y_mat = Y_appended, contrast = c(1,-1))
```

ess\_rand

Randomization-based Effective Sample Size (ESS)

### Description

Computes the randomization-based effective sample size (ESS) of a collection of assignments under a given set of potential outcomes.

### Usage

```
ess_rand(assign_array, Y_mat, contrast = c(1, -1))
```

#### **Arguments**

assign\_array

A three dimensional array containing a set of independent realizations of a collection the designs. The first coordinate of the array represents the iterations for each design. The second coordinate represents the units. The third coordinate represents the design.

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Y\_mat A matrix of potential outcomes, where rows represent units and columns represent treatment levels (ordered).

contrast A vector of the coefficients of the treatment contrast of interest. For example, for estimating the average treatment effect of treatment 1 versus treatment 2, contrast = c(1,-1).

#### Value

A vector of effective sample sizes for the given collection of assignments.

### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

```
# Consider N = 12, n1 = n2 = 6.
df_{sample} = data.frame(index = 1:12, x = c(20,30,40,40,50,60,20,30,40,40,50,60))
# Generate the potential outcomes.
Y_1 = 100 + (df_sample$x - mean(df_sample$x)) + rnorm(12, 0, 4)
Y_2 = Y_1 + 50
# Create matrix of potential outcomes.
Y_{appended} = cbind(Y_1, Y_2)
# Generate 100 assignments under CRD and the FSM.
Z_{crd_iter} = matrix(rep(0, 100 * 12), nrow = 100)
Z_fsm_iter = matrix(rep(0, 100 * 12), nrow = 100)
for(i in 1:100)
# Generate an assignment vector under CRD.
fc = crd(data_frame = df_sample, n_treat = 2, treat_sizes = c(6,6), control = FALSE)
Z_crd_iter[i,] = fc$Treat
# Generate an assignment vector under the FSM.
som_iter = som(data_frame = NULL, n_treat = 2,
treat_sizes = c(6, 6),include_discard = FALSE,
method = 'SCOMARS', marginal_treat = rep((6/12), 12), control = FALSE)
f = fsm(data_frame = df_sample, SOM = som_iter, s_function = 'Dopt',eps = 0.0001,
ties = 'random', intercept = TRUE, standardize = TRUE, units_print = FALSE)
Z_fsm_iter[i,] = f$data_frame_allocated$Treat
}
# Create a 3-dim array of assignments.
Z_{array} = array(0, dim = c(100, 12, 2))
Z_{array}[,,1] = Z_{crd}iter
Z_{array}[,,2] = Z_{fsm_iter}
# Calculate the ESS.
ess_rand(assign_array = Z_{array}, Y_{mat} = Y_{appended}, contrast = c(1,-1))
```

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fsm

Finite Selection Model (FSM)

### **Description**

Generates a randomized assignment of a group of units to multiple groups of pre-determined sizes using the Finite Selection Model (FSM).

### Usage

```
fsm(
  data_frame,
  SOM,
  s_function = "Dopt",
  Q_initial = NULL,
  eps = 0.001,
  ties = "random",
  intercept = TRUE,
  standardize = TRUE,
  units_print = TRUE,
  index_col = TRUE,
  Pol_mat = NULL,
  w_pol = NULL
)
```

#### **Arguments**

data\_frame

A data frame containing a column of unit indices (optional) and covariates (or transformations thereof).

SOM

A selection order matrix.

s\_function

Specifies a selection function, a string among 'constant', 'Dopt', 'Aopt', 'max pc', 'min pc', 'Dopt pc', 'max average', 'min average', 'Dopt average'. 'constant' selection function puts a constant value on every unselected unit. 'Dopt' use the D-optimality criteria based on the full set of covariates to select units. 'Aopt' uses the A-optimality criteria. 'max pc' (respectively, 'min pc') selects that unit that has the maximum (respectively, minimum) value of the first principal component. 'Dopt pc' uses the D-optimality criteria on the first principal component, 'max average' (respectively, 'min average') selects that unit that has the maximum (respectively, minimum) value of the simple average of the covariates. 'Dopt average' uses the D-optimality criteria on the simple average of the covariates.

Q\_initial

A (optional) non-singular matrix (called 'initial matrix') that is added the  $(X^TX)$  matrix of the choosing treatment group at any stage, when the  $(X^TX)$  matrix of that treatment group at that stage is non-invertible. If FALSE, the  $(X^TX)$  matrix for the full set of observations is used as the non-singular matrix. Applicable if s\_function = 'Dopt' or 'Aopt'.

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eps	Proportionality constant for Q_initial, the default value is 0.001.
ties	Specifies how to deal with ties in the values of the selection function. If ties = 'random', a unit is selected randomly from the set of candidate units. If ties = 'smallest', the unit that appears earlier in the data frame, i.e. the unit with the smallest index gets selected.
intercept	if TRUE, the design matrix of each treatment group includes a column of intercepts.
standardize	if TRUE, the columns of the $X$ matrix other than the column for the intercept (if any), are standardized.
units_print	if TRUE, the function automatically prints the candidate units at each step of selection.
index_col	if TRUE, data_frame contains a column of unit indices.
Pol_mat	Policy matrix. Applicable only when s_function = 'Aopt'.
w_pol	A vector of policy weights. Applicable only when s_function = 'Aopt'.

### Value

A list containing the following items.

data\_frame\_allocated: The original data frame augmented with the column of the treatment indicator.

som\_appended: The SOM with augmented columns for the indices and covariate values for units selected.

som\_split: som\_appended, split by the levels of the treatment.

crit\_print: The value of the objective function, at each stage of build up process. At each stage, the unit that maximizes the objective function is selected.

### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

Morris, C. (1979), "A finite selection model for experimental design of the health insurance study", Journal of Econometrics, 11(1), 43–61.

Morris, C., Hill, J. (2000), "The health insurance experiment: design using the finite selection model", Public policy and statistics: case studies from RAND, Springer Science & Business Media, 29–53.

```
# Load the data. 
 df_{sample} = data.frame(index = 1:12, x = c(20,30,40,40,50,60,20,30,40,40,50,60)) 
 # Generate an SOM with N = 12, n1 = n2 = 6. 
 som_{sample} = som(_{treat} = 2, treat_{sizes} = c(6,6), method = 'SCOMARS', control = TRUE,
```

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```
marginal\_treat = rep(6/12,12))
# Assign units given the SOM.
f = fsm(data_frame = df_sample, SOM = som_sample, s_function = 'Dopt',
eps = 0.001, ties = 'random', intercept = TRUE, standardize = TRUE, units_print = TRUE,
index_col = TRUE)
```

fsm\_batch

Batched FSM for sequential experiments

### **Description**

Extension of the FSM to cases where units arrive sequentially in batches.

### Usage

```
fsm_batch(
  data_frame,
  data_frame_past,
  t_ind,
  SOM,
  s_function = "Dopt",
 Q_initial = NULL,
  eps = 0.001,
  ties = "random",
  intercept = TRUE,
  index_col_past = TRUE,
  standardize = TRUE,
  units_print = TRUE,
  index_col = TRUE,
 Pol_mat = NULL,
  w_pol = NULL
)
```

### **Arguments**

data\_frame Data frame containing a column of unit indices (optional) and covariates (or transformations thereof).

data\_frame\_past

A data frame of units already allocated to treatment groups. Data frame contains a column of unit indices (optional), columns of covariates (or transformations thereof), and a column for treatment indicator.

column name containing the treatment indicator in data\_frame\_past. t\_ind

SOM Selection Order Matrix.

s\_function Specifies a selection function, a string among 'constant', 'Dopt', 'Aopt', 'max pc', 'min pc', 'Dopt pc', 'max average', 'min average', 'Dopt average'.

'constant' selection function puts a constant value on every unselected unit.

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'Dopt' use the D-optimality criteria based on the full set of covariates to select units. 'Aopt' uses the A-optimality criteria. 'max pc' (respectively, 'min pc') selects that unit that has the maximum (respectively, minimum) value of the first principal component. 'Dopt pc' uses the D-optimality criteria on the first principal component, 'max average' (respectively, 'min average') selects that unit that has the maximum (respectively, minimum) value of the simple average of the covariates. 'Dopt average' uses the D-optimality criteria on the simple average of the covariates.

Q\_initial A (optional) non-singular matrix (called 'initial matrix') that is added the  $(X^TX)$ 

matrix of the choosing treatment group at any stage, when the  $(X^TX)$  matrix of that treatment group at that stage is non-invertible. If FALSE, the  $(X^TX)$  matrix for the full set of observations is used as the non-singular matrix. Applicable if

s\_function = 'Dopt' or 'Aopt'.

eps Proportionality constant for Q\_initial, the default value is 0.001.

ties Specifies how to deal with ties in the values of the selection function. If ties =

'random', a unit is selected randomly from the set of candidate units. If ties = 'smallest', the unit that appears earlier in the data frame, i.e. the unit with the

smallest index gets selected.

intercept if TRUE, the design matrix of each treatment group includes a column of inter-

cepts.

index\_col\_past TRUE if column of unit indices is present in data\_frame\_past.

standardize if TRUE, the columns of the X matrix other than the column for the intercept (if

any), are standardized.

units\_print if TRUE, the function automatically prints the candidate units at each step of

selection.

index\_col if TRUE, data\_frame contains a column of unit indices.

Pol\_mat Policy matrix. Applicable only when s\_function = 'Aopt'.

w\_pol A vector of policy weights. Applicable only when s\_function = 'Aopt'.

### Value

A list containing the following items.

data\_frame\_allocated: The original data frame augmented with the column of the treatment indicator.

som\_appended: The SOM with augmented columns for the indices and covariate values for units selected.

som\_split: som\_appended, split by the levels of the treatment.

data\_frame\_allocated\_augmented: data frame combining data\_frame\_allocated and data\_frame\_past.

#### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta

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#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R'.

#### **Examples**

```
# Consider N=18, number of treatments = 2, n1 = n2 = 9, batch sizes = 6,6,6.
# Get data frame for the first batch.
df_{sample_1} = data.frame(index = 1:6, age = c(20,30,40,40,50,60))
# Obtain SOM for all the 12 units.
som_gen = som(data_frame = NULL, n_treat = 2, treat_sizes = c(9,9),
include_discard = FALSE, method = 'SCOMARS', marginal_treat = rep((9/18),18), control = FALSE)
# Assign the first batch.
f1 = fsm(data_frame = df_sample_1, SOM = som_gen[1:6,], s_function = 'Dopt',
eps = 0.0001, ties = 'random', intercept = TRUE, standardize = TRUE, units_print = TRUE)
f1_app = f1$data_frame_allocated
# Get data frame for the second batch.
df_{sample_2} = data.frame(index = 7:12, age = c(20,30,40,40,50,60))
# Assign the second batch.
f2 = fsm_batch(data_frame = df_sample_2, SOM = som_gen[7:12,], s_function = 'Dopt',
eps = 0.0001, ties = 'random', intercept = TRUE, standardize = TRUE, units_print = TRUE,
data_frame_past = f1_app, t_ind = 'Treat', index_col_past = TRUE)
f2_app = f2$data_frame_allocated_augmented
# Get data frame for the third batch.
df_{sample_3} = data.frame(index = 13:18, age = c(20,30,40,40,50,60))
# Assign the third batch.
f3 = fsm_batch(data_frame = df_sample_3, SOM = som_gen[13:18,], s_function = 'Dopt',
eps = 0.0001, ties = 'random', intercept = TRUE, standardize = TRUE, units_print = TRUE,
data_frame_past = f2_app, t_ind = 'Treat', index_col_past = TRUE)
f3_app = f3$data_frame_allocated_augmented
```

Lalonde

The Lalonde experimental dataset

### **Description**

The Nationally Supported Work (NSW) experimental data set by Lalonde (1986).

### References

LaLonde, R. J. (1986), "Evaluating the econometric evaluations of training programs with experimental data". The American Economic Review, pp. 604–620.

Dehejia, R., and Wahba, S. (1999), "Causal effects in nonexperimental studies: Reevaluating the evaluation of training programs", Journal of the American Statistical Association, 94(443), 1053–1062.

https://users.nber.org/~rdehejia/nswdata.html

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

love\_plot

### Description

Generates a Love plot of Absolute Standardized Mean Differences (ASMD) or Target Absolute Standardized Differences (TASMD) between two groups under one or two designs.

### Usage

```
love_plot(
  data_frame,
  index_col = TRUE,
  alloc1,
  alloc2 = NULL,
  imbalance = "TASMD",
  treat_lab = 1,
  vline = "",
  xupper = 1,
  mean_tar = NULL,
  sd_tar = NULL,
  denom = "target",
  legend_text = "FSM",
  legend_position = "topright"
)
```

### Arguments

data_frame	Data frame containing a column of unit indices (optional) and covariates (or transformations thereof).
index_col	if TRUE, data_frame contains a column of unit indices.
alloc1	A vector of treatment assignment.
alloc2	A (optional) vector of treatment assignment.
imbalance	Measure of imbalance used. If imbalance = 'TASMD', imbalance is computed using the Target Absolute Standardized Mean Differences (TASMD). If imbalance = 'ASMD', imbalance is computed using the Absolute Standardized Mean Differences (ASMD)
treat_lab	Label of the treatment group in which the TASMD is computed. Applicable only when imbalance = 'TASMD'.
vline	A (optional) x-coordinate at which a vertical line is drawn.
xupper	Upper limit of the x-axis.
mean_tar	A (optional) vector of target profile of the covariates under consideration, e.g., mean of the covariates in the target population. Applicable only when imbalance = 'TASMD'. If mean_tar = NULL, the full-sample average of the covariates is considered as the target profile.

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sd\_tar A optional vector of the standard deviation of the covariates in the target population. Applicable only when imbalance = 'TASMD'.

denom Specifies the denominator for the computation of TASMD. If denom = 'target',

the standard deviations of the covariates in the target population are used. If denom = 'group', the standard deviations of the covariates in the treatment group given by treat\_lab are used. Applicable only when imbalance = 'TASMD'.

legend\_text Legend of the two designs under consideration.

legend\_position

= Position of the legend in the plot. The default is 'topright'.

#### Value

Love plot of the ASMD/TASMD of the covariates.

#### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

Love, T. (2004), "Graphical display of covariate balance", Presentation, See http://chrp.org/love/JSM2004RoundTableHandor 1364.

```
# Consider the Lalonde dataset.
# Get the full sample size.
N = nrow(Lalonde)
# Get the treatment group sizes.
n1 = floor(N/2)
n2 = N-n1
# Generate an SOM.
som_obs = som(n_treat = 2, treat_sizes = c(n1,n2),include_discard = FALSE,
method = 'SCOMARS', marginal_treat = rep((n2/N),N), control = FALSE)
# Generate a treatment assignment given som_obs.
f = fsm(data_frame = Lalonde, SOM = som_obs, s_function = 'Dopt', eps = 0.0001,
ties = 'random', intercept = TRUE, standardize = TRUE, units_print = FALSE)
# Get assignment vector under the FSM.
Z_fsm_obs = f$data_frame_allocated$Treat
# Draw a random CRD.
Z_crd_obs = crd(data_frame = Lalonde, n_treat = 2, treat_sizes = c(n1, n2),
control = FALSE)$Treat
# Draw Love plot.
love_plot(data_frame = Lalonde, index_col = TRUE, alloc1 = Z_fsm_obs, alloc2 = Z_crd_obs,
imbalance = 'TASMD', treat_lab = 1, mean_tar = NULL, sd_tar = NULL, denom = 'target',
vline = "", legend_text = c("FSM","CRD"), xupper = 0.15, legend_position = 'topright')
```

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make_sq_inter	Squares and two-way interactions of variables	
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### Description

Generates squares and/or two-way interactions (pairwise products) of the columns of a data frame.

### Usage

```
make_sq_inter(
  data_frame,
  is_square = TRUE,
  is_inter = TRUE,
  keep_marginal = TRUE
)
```

### Arguments

data_frame	Data frame containing the	variables whose squares	and interactions	are to be

created.

is\_square If TRUE, square of each column of data\_frame is created.

is\_inter If TRUE, product of every pair of columns of data\_frame is created.

keep\_marginal If TRUE, the original columns of data\_frame are retained in the resulting data

frame.

### Value

A data frame containing the squares and/or pairwise products of data\_frame.

### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

```
# Consider a data frame with N = 12 units and 2 covariates. data_frame_sample = data.frame(male = c(rep(1,6),rep(0,6)), age = c(20,30,40,40,50,60,20,30,40,40,50,60))
# Get a data frame with all possible squares and first order interactions. make_sq_inter(data_frame = data_frame_sample, is_square = TRUE, is_inter = TRUE, keep_marginal = FALSE)
```

perm\_test

perm\_test

Fisher's randomization test for sharp null hypothesis.

### Description

Performs Fisher's randomization test for sharp null hypotheses of the form  $H_0: c_1Y_i(1)+c_2Y_i(2)-\tau=0$ , for a vector of contrasts  $(c_1,c_2)$ .

### Usage

```
perm_test(
   Y_obs,
   alloc_obs,
   alloc,
   contrast = c(1, -1),
   tau = 0,
   method = "marginal mean",
   alternative = "not equal"
)
```

### Arguments

Y_obs	Vector of observed outcome.
alloc_obs	Vector of observed treatment assignment.
alloc	A matrix of treatment assignments over which the randomization distribution of the test statistic is computed. Each row of alloc should correspond to an assignment vector.
contrast	A vector of the coefficients of the treatment contrast of interest. For example, for estimating the average treatment effect of treatment 1 versus treatment 2, $contrast = c(1,-1)$ .
tau	The value of the treatment contrast specified by the sharp null hypothesis.
method	The method of computing the test statistic. If method = 'marginal mean', the test statistic is $c_1\hat{Y}_i(1)+c_2\hat{Y}_i(2)$ , where $\hat{Y}(z)$ is the mean of the observed outcome in the group $Z=z$ , for $z=0,1$ . If method = 'marginal rank', the test statistic is $c_1\hat{Y}_i(1)+c_2\hat{Y}_i(2)$ , where $\hat{Y}(z)$ is the mean of the rank of the observed outcome in the group $Z=z$ , for $z=0,1$
alternative	The type of alternative hypothesis used. For right-sided test, alternative = 'greater'. For left-sided test, alternative = 'less'. For both-sided test, alternative = 'not equal'.

### Value

A list containing the following items.

test\_stat\_obs: The observed value of the test statistic.

test\_stat\_iter: A vector of values of the test statistic across repeated randomizations.

p\_value: p-value of the test.

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#### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

### **Examples**

```
# Consider N = 12, n1 = n2 = 6.
# We test the sharp null of no treatment effect under CRD.
df_{sample} = data.frame(index = 1:12, x = c(20,30,40,40,50,60,20,30,40,40,50,60))
# True potential outcomes.
Y_1_{true} = 100 + (df_{sample} - mean(df_{sample})) + rnorm(12, 0, 4)
Y_2_true = Y_1_true + 50
# Generate the realized assignment under CRD.
fc = crd(data_frame = df_sample, n_treat = 2, treat_sizes = c(6,6), control = FALSE)
Z_crd_obs = fc$Treat
# Get the observed outcomes
Y_{obs} = Y_{1_{true}}
Y_obs[Z_crd_obs == 2] = Y_2_true[Z_crd_obs == 2]
# Generate 1000 assignments under CRD.
Z_{crd_iter} = matrix(rep(0, 1000 * 12), nrow = 1000)
for(i in 1:1000)
fc = crd(data_frame = df_sample, n_treat = 2, treat_sizes = c(6,6), control = FALSE)
Z_crd_iter[i,] = fc$Treat
# Test for the sharp null H0: Y_i(1) = Y_i(0) for all i.
# Alternative: not H0 (two-sided test).
perm = perm_test(Y_obs = Y_obs, alloc_obs = Z_crd_obs, alloc = Z_crd_iter,
contrast = c(1,-1), tau = 0, method = "marginal mean", alternative = 'not equal')
# Obtain the p-value.
perm$p_value
```

som

Selection Order Matrix (SOM)

### **Description**

Generates a Selection Order Matrix (SOM) in a deterministic/random manner.

## Usage

```
som(
  data_frame = NULL,
  n_treat,
  treat_sizes,
```

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```
include_discard = FALSE,
method = "SCOMARS",
control = FALSE,
marginal_treat = NULL
)
```

### **Arguments**

data\_frame A (optional) data frame corresponding to the full sample of units. Required if

include\_discard = TRUE.

n\_treat Number of treatment groups.

treat\_sizes A vector of treatment group sizes. If control = TRUE, the first element of

treat\_sizes should be the control group size.

include\_discard

TRUE if a discard group is considered.

method Specifies the selection strategy used among 'global percentage', 'randomized

chunk', 'SCOMARS'. 'SCOMARS' is applicable only if n\_treat = 2.

control If TRUE, treatments are labeled as 0,1,...,g-1 (0 representing the control group).

If FALSE, they are labeled as 1,2,...,g.

marginal\_treat A vector of marginal probabilities, the jth element being the probability that

treatment group (or treatment group 2 in case control = FALSE) gets to choose at the jth stage given the total number of choices made by treatment group upto

the (j-1)th stage. Only applicable when method = 'SCOMARS'.

### Value

A data frame containing the selection order of treatments, i.e. the labels of treatment groups at each stage of selection. If method = 'SCOMARS', the data frame contains an additional column of the conditional selection probabilities.

### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

Morris, C. (1983), "Sequentially controlled Markovian random sampling (SCOMARS)", Institute of Mathematical Statistics Bulletin, 12(5), 237.

```
# Generate an SOM with N = 12, n1 = n2 = 6.

som_sample = som(data_frame = NULL, n_treat = 2, treat_sizes = c(6,6), include_discard = FALSE,

method = 'SCOMARS', control = FALSE, marginal_treat = rep(6/12,12))
```

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tasmd\_rand

Target Absolute Standardized Mean Differences (TASMD)

### Description

Computes the mean and standard deviation of Target Absolute Standardized Mean Differences (TASMD) of multiple covariates (or transformations thereof) in a treatment group relative to a target population or a target individual for a set of assignments under one or two designs.

### Usage

```
tasmd_rand(
  data_frame,
  index_col = FALSE,
  alloc1,
  alloc2,
  treat_lab = 1,
  legend = c("CRD", "FSM"),
  mean_tar = NULL,
  sd_tar = NULL,
  denom = "target",
  roundoff = 3
)
```

### **Arguments**

data_frame	Data frame containing a column of unit indices (optional) and covariates (or transformations thereof).
index_col	if TRUE, data_frame contains a column of unit indices.
alloc1	A matrix or vector of treatment assignments. If alloc1 is a matrix, then each row should correspond to an assignment vector.
alloc2	A (optional) matrix or vector of treatment assignment. If alloc2 is a matrix, then each row should correspond to an assignment vector.
treat_lab	Label of the treatment group in which the TASMD is computed.
legend	Legend of the two designs under consideration.
mean_tar	A (optional) vector of target profile of the covariates under consideration, e.g., mean of the covariates in the target population. Applicable only when imbalance = 'TASMD'. If mean_tar = NULL, the full-sample average of the covariates is considered as the target profile.
sd_tar	A optional vector of the standard deviation of the covariates in the target population. Applicable only when imbalance = 'TASMD'.
denom	Specifies the denominator for the computation of TASMD. If denom = 'target', the standard deviations of the covariates in the target population are used. If denom = 'group', the standard deviations of the covariates in the treatment group given by treat_lab are used. Applicable only when imbalance = 'TASMD'.

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roundoff

A number indicating the number of decimal places to be used for rounding off the TASMDs.

### Value

A list containing the following items (if alloc1 and alloc2 are matrices)

tasmd\_table: A matrix containing the means (standard deviations in parenthesis) of the TASMDs for the designs under consideration. If alloc1 or alloc2 is a vector, the TASMD of the corresponding assignment is returned.

tasmd\_mean: A matrix containing the means of the TASMDs for the designs under consideration.

tasmd\_sd: A matrix containing the standard deviations of the TASMDs for the designs under consideration.

If alloc1 and alloc2 are vectors, tasmd\_rand produces a data frame of the corresponding TAS-MDs.

#### Author(s)

Ambarish Chattopadhyay, Carl N. Morris and Jose R. Zubizarreta.

#### References

Chattopadhyay, A., Morris, C. N., and Zubizarreta, J. R. (2020), "Randomized and Balanced Allocation of Units into Treatment Groups Using the Finite Selection Model for R".

```
# Consider the Lalonde dataset.
# Get the full sample size.
N = nrow(Lalonde)
# Get the treatment group sizes.
n1 = floor(N/2)
n2 = N-n1
# Generate an SOM.
som_obs = som(n_treat = 2, treat_sizes = c(n1,n2),include_discard = FALSE,
method = 'SCOMARS', marginal_treat = rep((n2/N),N), control = FALSE)
# Generate a treatment assignment given som_obs.
f = fsm(data_frame = Lalonde, SOM = som_obs, s_function = 'Dopt', eps = 0.0001,
ties = 'random', intercept = TRUE, standardize = TRUE, units_print = FALSE)
# Get assignment vector under the FSM.
Z_fsm_obs = f$data_frame_allocated$Treat
# Draw a random CRD.
Z_crd_obs = crd(data_frame = Lalonde, n_treat = 2, treat_sizes = c(n1, n2),
control = FALSE)$Treat
# Calculate the TASMD.
TASMD = tasmd_rand(data_frame = Lalonde, index_col = TRUE, alloc1 = Z_crd_obs,
alloc2 = Z_fsm_obs, treat_lab = 1, mean_tar = NULL, sd_tar = NULL,
denom = 'target', legend = c('CRD', 'FSM'), roundoff = 3)
```

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