# Package 'madr' 

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Description Estimates average treatment effects using model average double robust (MA-DR) estimation. The MA-DR estimator is defined as weighted average of double robust estimators, where each double robust estimator corresponds to a specific choice of the outcome model and the propensity score model. The MA-DR estimator extend the desirable double robustness property by achieving consistency under the much weaker assumption that either the true propensity score model or the true outcome model be within a specified, possibly large, class of models.
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add.to.dictionary Worker function that fits propensity score models

## Description

This function fits propensity score models and saves necessary information

## Usage

add.to.dictionary(X, U, W, alpha)

## Arguments

| $X$ | vector of the treatment $(0 / 1)$ |
| :--- | :--- |
| $U$ | matrix of covariates to be considered for inclusion/exclusion |
| W | matrix of covariates that will be included in all models (optional) |
| alpha | vector of inclusion indicators (which columns of $U$ ) to included in the propensity <br> score model |

## Value

A list. The list contains the following named components:
out a list that contains the BIC and estimated propensity scores from propensity score models

```
add.to.dictionary.outcome
```

Worker function that fits outcome models

## Description

This function fits outcome models and saves necessary information

## Usage

add.to.dictionary.outcome(Y, X, U, W, alpha, binary = F)

## Arguments

| Y | vector of the outcome |
| :--- | :--- |
| X | vector of the treatment $(0 / 1)$ |
| U | matrix of covariates to be considered for inclusion/exclusion |
| W | matrix of covariates that will be included in all models (optional) |
| alpha | vector of inclusion indicators (which columns of U ) to included in the propensity <br> score model |
| binary | indicates if the outcome is binary |

## Value

A list. The list contains the following named components:
out a list that contains the BIC, predicted values, and estimated treatment effect from each outcome model

## Description

This function transforms BIC to model probabilities

## Usage

bic.to.prob(bic)

## Arguments

bic vector of BICs

## Value

A vector of model probabilities of the same dimension of bic

```
expit Expit (inverse logit) function
```


## Description

This function transforms the input using the expit function

## Usage <br> expit(x)

## Arguments

$x \quad$ vector of values to apply the expit function

## Value

A vector of the same dimension of $x$

```
madr
Calculate model averaged double robust estimate
```


## Description

This function estimates a model averaged double robust estimate.

## Usage

$\operatorname{madr}(\mathrm{Y}, \mathrm{X}, \mathrm{U}, \mathrm{W}=\mathrm{NULL}, \mathrm{M}=1000$, cut $=0.95$, enumerate $=\mathrm{F}$, tau $=$ NULL, two.stage $=$ NULL)

## Arguments

| Y | vector of the outcome |
| :--- | :--- |
| X | vector of the treatment (0/1) |
| U | matrix of covariates to be considered for inclusion/exclusion |
| W | matrix of covariates that will be included in all models (optional) <br> M |
| cut | cumulative probability of models to be retained for improved computational ef- <br> ficiency (1 retains all visited models) |
| enumerate | indicator if all possible models should be enumerated (default: FALSE) <br> tau |
| two.stage | scalar value for the prior model dependence (1 is an independent prior; defaults <br> to 0) <br> indicator if the two-stage procedure for calculating the model weights should be <br> used (defaults to TRUE) |

## Value

A list. The list contains the following named components:

| madr | the model averaged double robust estimate |
| :--- | :--- |
| weight.ps | a vector that contains the inclusion probability of each covariate in the propen- <br> sity score model |
| weight. om | a vector that contains the inclusion probability of each covariate in the outcome <br> model |

## Examples

```
set.seed(122)
## generate data
n = 100 # number of observations
k = 4 # number of covariates
U = matrix(rnorm(n*k),n,k)
colnames(U) = paste0("U",1:k)
A = rbinom(n,1, expit(-1+.5*rowSums(U)))
Y = rnorm(n,1+A+.25*rowSums(U))
## A is confounded -- true effect is 1
lm(Y~A)
## fit ma-dr -- can enumerate models if k isnt too big
res = madr (Y=Y,X=A,U=U, enumerate=TRUE,tau=1,two.stage=FALSE) # independent prior
res
res = madr (Y=Y, X=A,U=U, enumerate=TRUE, tau=0, two.stage=TRUE) # tau=0 and using two-stage weights
res
## no need to refit madr each time when enumerating -- use summarize and specify different taus
summary(res,tau=1, two.stage=FALSE) # independent prior
summary(res, tau=0, two.stage=FALSE)
summary(res,tau=0,two.stage=TRUE) # two-stage procedure for calculating weights
## use mcmc instead of enumerating (the default)
madr (Y=Y, X=A,U=U,M=1000, cut=1) #should approximate tau=0 and two.stage=TRUE
```

```
madr.enumerate Model averaged double robust estimate with enumeration of all possi-
    ble models (linear terms only)
```


## Description

This function enumerates all possible models and estimates a model averaged double robust estimate

## Usage

madr.enumerate $(\mathrm{Y}, \mathrm{X}, \mathrm{U}, \mathrm{W}=\mathrm{NULL}$, tau $=1$, two. stage $=\mathrm{F})$

## Arguments

| Y | vector of the outcome |
| :--- | :--- |
| X | vector of the treatment indicator $(0 / 1)$ |
| U | matrix of covariates to be considered for inclusion/exclusion |
| W | matrix of covariates that will be included in all models (optional) |
| tau | scalar value for the prior model dependence (1 is an independent prior) <br> two.stage |
| indicator if the two-stage procedure for calculating the model weights should be <br> used |  |

## Value

A object of class madr.enumerate. The object contains the following named components:
out a matrix that contains the BIC and estimated treatment from each outcome model
ps a matrix that contains the BIC from each propensity score model
dr a matrix that contains the model-specific double robust estimates
U. names the column names of $U$

| madr.mcmc | Calculate model averaged double robust estimate using a pseudo-MC3 <br> algorithm |
| :--- | :--- |

## Description

This function uses a pseudo-MC3 algorithm to search the model space, then estimate a model averaged double robust estimate using the two-stage procedure for estimating model weights with tau $=0$.

## Usage

madr.mcmc $(\mathrm{Y}, \mathrm{X}, \mathrm{U}, \mathrm{W}=\mathrm{NULL}, \mathrm{M}=1000$, cut $=0.95$ )

## Arguments

Y
$X \quad$ vector of the treatment ( $0 / 1$ )
U matrix of covariates to be considered for inclusion/exclusion
W matrix of covariates that will be included in all models (optional)
M the number of MCMC iteration
cut cumulative probability of models to be retained for improved computational efficiency (1 retains all visited models)

## Value

A list. The list contains the following named components:
madr the model averaged double robust estimate
weight.ps a vector that contains the inclusion probability of each covariate in the propensity score model
weight.om a vector that contains the inclusion probability of each covariate in the outcome model

```
OM.MA
```

Calculate model probabilities for the outcome models using a pseudoMC3 algorithm

## Description

This function uses a pseudo-MC3 algorithm to search the outcome model space.

## Usage

OM. MA (Y, X, U, $W=$ NULL, $M=1000$, alpha $=$ NULL, binary = F)

## Arguments

Y
X
U
W matrix of covariates that will be included in all models (optional)
M
alpha
binary
vector of the outcome
vector of the treatment $(0 / 1)$
matrix of covariates to be considered for inclusion/exclusion
the number of MCMC iteration
vector of inclusion indicators (which columns of U ) to start MCMC algorithm (optional)
indicator if the outcome is binary (optional)

## Value

A list. The list contains the following named components:
dict a list that contains the BIC, predicted values, and estimated treatment effect from each outcome model
alpha the last model visited by the algorithm
out. table a matrix that contains the BIC and estimated treatment effect from each outcome model

## Description

This function enumerates and fits all possible outcome models

## Usage

OM.MA.enumerate(Y, X, U, W = NULL)

## Arguments

$Y$ vector of the outcome
$X \quad$ vector of the treatment indicator (0/1)
U matrix of covariates to be considered for inclusion/exclusion
W matrix of covariates that will be included in all models (optional)

## Value

A list. The listcontains the following named components:
dict a list that contains the BIC, predicted values, and estimated treatment effect from each outcome model
out. table a matrix that contains the BIC and estimated treatment effect from each outcome model
print.madr.enumerate Print function for madr.enumerate class

## Description

This function prints results from madr.enumerate class

## Usage

\#\# S3 method for class 'madr.enumerate'
print(x, ...)

## Arguments

$\begin{array}{ll}x & \text { madr.enumerate object } \\ \ldots & \text { ignored }\end{array}$

```
print.madr.mcmc Print function for madr.memc class
```


## Description

This function prints results from madr.memc class

## Usage

\#\# S3 method for class 'madr.mcmc'
print(x, ...)

## Arguments

| $x$ | madr.mcmc object |
| :--- | :--- |
| $\ldots$ | ignored |

```
    print.summary.madr.enumerate
```

            Print function for summary.madr.enumerate class
    
## Description

This function prints results from summary.madr.enumerate class

## Usage

\#\# S3 method for class 'summary.madr.enumerate'
print(x, ...)

## Arguments

x
...
summary.madr.enumerate object
ignored

Calculate model probabilities for the propensity score model using a pseudo-MC3 algorithm

## Description

This function uses a pseudo-MC3 algorithm to search the propensity score model space.

## Usage

```
PS.MA(X, U, W = NULL, M = 1000, alpha = NULL, master.index = NULL,
    master. dict \(=\) list())
```


## Arguments

$X \quad$ vector of the treatment $(0 / 1)$
U matrix of covariates to be considered for inclusion/exclusion
W matrix of covariates that will be included in all models (optional)
M the number of MCMC iteration
alpha vector of inclusion indicators (which columns of U ) to start MCMC algorithm (optional)
master.index indexes which columns of $U$ should be considered for inclusion in the propensity score model (optional)
master.dict list containing information from previous propensity score model fits (optional)

## Value

A list. The list contains the following named components:
dict a list that contains the BIC and estimated propensity scores from propensity score models
alpha the last model visited by the algorithm
out.table a matrix that contains the BIC from each propensity score model

## Description

This function enumerates and fits all possible propensity score models

## Usage

PS.MA.enumerate(X, U, W = NULL)

## Arguments

$X \quad$ vector of the treatment indicator $(0 / 1)$
U matrix of covariates to be considered for inclusion/exclusion
W matrix of covariates that will be included in all models (optional)

## Value

A list. The list contains the following named components:
dict a list that contains the BIC and estimated propensity scores from propensity score models
out.table a matrix that contains the BIC from each propensity score model
summary.madr.enumerate
Provides model averaged double robust estimate for different values of tau

## Description

This function estimates model averaged double robust estimate for different values of tau using a madr.enumerate object

## Usage

\#\# S3 method for class 'madr.enumerate'
summary (object, tau $=$ NULL, two.stage $=$ NULL,.. )

## Arguments

| object | madr.enumerate object |
| :--- | :--- |
| tau | scalar value for the prior model dependence (1 is an independent prior; defaults <br> to value used in madr.enumerate) |
| two.stage | indicator if the two-stage procedure for calculating the model weights should be <br> used (defaults to value used in madr.enumerate) |
| $\ldots$ | ignored |

## Value

A list. The list contains the following named components:

| madr | the model averaged double robust estimate |
| :--- | :--- |
| weight.ps | a vector that contains the inclusion probability of each covariate in the propen- <br> sity score model <br> a vector that contains the inclusion probability of each covariate in the outcome <br> model |
| weight.om | value of tau used in estimation |
| tau | indicator if the two-stage procedure for calculating the model weights was used |
| two.stage |  |

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