Package 'evalITR'

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Title Evaluating Individualized Treatment Rules

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Depends stats, R (>= 3.5.0)

Description A collection of statistical methods for evaluating individualized treatment rules under randomized data. The provided metrics include PAV (Population Average Value), PAPE (Population Average Prescription Effect), and AUPEC (Area Under Prescription Effect Curve). It also provides the tools to analyze individualized treatment rules under budget constraints. Imai and Li (2019) <arXiv:1905.05389>.

License GPL (>= 2)

URL https://github.com/MichaelLLi/evalITR

BugReports https://github.com/MichaelLLi/evalITR/issues

RoxygenNote 7.0.2

Suggests testthat

NeedsCompilation no

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AUPEC

Estimation of the Area Under Prescription Evaluation Curve (AU-PEC) in Randomized Experiments

Description

This function estimates AUPEC. The details of the methods for this design are given in Imai and Li (2019).

Usage

AUPEC(Tr, tau, Y, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
tau	A vector of the unit-level continuous score for treatment assignment. We assume those that have tau<0 should not have treatment. Conditional Average Treatment Effect is one possible measure.
Y	A vector of the outcome variable of interest for each sample.
centered	If TRUE, the outcome variables would be centered before processing. This mini- mizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

aupec	The estimated Area Under Prescription Evaluation Curve
sd	The estimated standard deviation of AUPEC.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
tau = c(0,0.1,0.2,0.3,0.4,0.5,0.6,0.7)
Y = c(4,5,0,2,4,1,-4,3)
aupeclist <- AUPEC(Tr,tau,Y)
aupeclist$aupec
aupeclist$sd</pre>
```

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AUPECcv

Estimation of the Area Under Prescription Evaluation Curve (AU-PEC) in Randomized Experiments Under Cross Validation

Description

This function estimates AUPEC. The details of the methods for this design are given in Imai and Li (2019).

Usage

AUPECcv(Tr, tau, Y, ind, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
tau	A matrix where the ith column is the unit-level continuous score for treatment assignment generated in the ith fold.
Υ	The outcome variable of interest.
ind	A vector of integers (between 1 and number of folds inclusive) indicating which testing set does each sample belong to.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

aupec	The estimated AUPEC.
sd	The estimated standard deviation of AUPEC.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
tau = matrix(c(0,0.1,0.2,0.3,0.4,0.5,0.6,0.7,-0.5,-0.3,-0.1,0.1,0.3,0.5,0.7,0.9),nrow = 8, ncol = 2)
Y = c(4,5,0,2,4,1,-4,3)
ind = c(rep(1,4),rep(2,4))
aupeclist <- AUPECcv(Tr, tau, Y, ind)
aupeclist$aupec
aupeclist$aupec</pre>
```

PAPD

Estimation of the Population Average Prescription Difference in Randomized Experiments

Description

This function estimates the Population Average Prescription Difference with a budget constraint. The details of the methods for this design are given in Imai and Li (2019).

Usage

PAPD(Tr, Thatfp, Thatgp, Y, plim, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
Thatfp	A vector of the unit-level binary treatment that would have been assigned by the first individualized treatment rule. Please ensure that the percentage of treatment units of That is lower than the budget constraint.
Thatgp	A vector of the unit-level binary treatment that would have been assigned by the second individualized treatment rule. Please ensure that the percentage of treatment units of That is lower than the budget constraint.
Υ	A vector of the outcome variable of interest for each sample.
plim	The maximum percentage of population that can be treated under the budget constraint. Should be a decimal between 0 and 1.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

papd	The estimated Population Average Prescription Difference
sd	The estimated standard deviation of PAPD.

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PAPDcv

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
That = c(0,1,1,0,0,1,1,0)
That2 = c(1,0,0,1,1,0,0,1)
Y = c(4,5,0,2,4,1,-4,3)
papdlist <- PAPD(Tr,That,That2,Y,plim = 0.5)
papdlist$papd
papdlist$sd</pre>
```

PAPDcv	Estimation of the Population Average Prescription Difference in Ran-
	domized Experiments Under Cross Validation

Description

This function estimates the Population Average Prescription Difference The details of the methods for this design are given in Imai and Li (2019).

Usage

```
PAPDcv(Tr, Thatfp, Thatgp, Y, ind, plim, centered = TRUE)
```

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
Thatfp	A matrix where the ith column is the unit-level binary treatment that would have been assigned by the first individualized treatment rule generated in the ith fold. Please ensure that the percentage of treatment units of That is lower than the budget constraint.
Thatgp	A matrix where the ith column is the unit-level binary treatment that would have been assigned by the second individualized treatment rule generated in the ith fold. Please ensure that the percentage of treatment units of That is lower than the budget constraint.
Y	The outcome variable of interest.
ind	A vector of integers (between 1 and number of folds inclusive) indicating which testing set does each sample belong to.

Value

A list that contains the following items:

papd	The estimated Population Average Prescription Difference.
sd	The estimated standard deviation of PAPD.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
That = matrix(c(0,1,1,0,0,1,1,0,0,1,1,0,0,1), nrow = 8, ncol = 2)
That2 = matrix(c(0,0,1,1,0,0,1,1,1,0,0,1,1,0,0), nrow = 8, ncol = 2)
Y = c(4,5,0,2,4,1,-4,3)
ind = c(rep(1,4),rep(2,4))
papdlist <- PAPDcv(Tr, That, That2, Y, ind, plim = 0.5)
papdlist$papd
papdlist$sd</pre>
```

PAPE

Estimation of the Population Average Prescription Effect in Randomized Experiments

Description

This function estimates the Population Average Prescription Effect with and without a budget constraint. The details of the methods for this design are given in Imai and Li (2019).

Usage

PAPE(Tr, That, Y, plim = NA, centered = TRUE)

PAPEcv

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
That	A vector of the unit-level binary treatment that would have been assigned by the individualized treatment rule. If plim is specified, please ensure that the percentage of treatment units of That is lower than the budget constraint.
Υ	A vector of the outcome variable of interest for each sample.
plim	The maximum percentage of population that can be treated under the budget constraint. Should be a decimal between 0 and 1. Default is NA which assumes no budget constraint.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

pape	The estimated Population Average Prescription Effect.
sd	The estimated standard deviation of PAPE.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
That = c(0,1,1,0,0,1,1,0)
Y = c(4,5,0,2,4,1,-4,3)
papelist <- PAPE(Tr,That,Y)
papelist$pape
papelist$sd</pre>
```

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PA	PE	CV

Estimation of the Population Average Prescription Effect in Randomized Experiments Under Cross Validation

Description

This function estimates the Population Average Prescription Effect with and without a budget constraint. The details of the methods for this design are given in Imai and Li (2019).

Usage

PAPEcv(Tr, That, Y, ind, plim = NA, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
That	A matrix where the ith column is the unit-level binary treatment that would have been assigned by the individualized treatment rule generated in the ith fold. If plim is specified, please ensure that the percentage of treatment units of That is lower than the budget constraint.
Υ	The outcome variable of interest.
ind	A vector of integers (between 1 and number of folds inclusive) indicating which testing set does each sample belong to.
plim	The maximum percentage of population that can be treated under the budget constraint. Should be a decimal between 0 and 1. Default is NA which assumes no budget constraint.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

pape	The estimated Population Average Prescription Effect.
sd	The estimated standard deviation of PAPE.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
That = matrix(c(0,1,1,0,0,1,1,0,0,1,1,0,0,1), nrow = 8, ncol = 2)
Y = c(4,5,0,2,4,1,-4,3)
ind = c(rep(1,4),rep(2,4))
papelist <- PAPEcv(Tr, That, Y, ind)
papelist$pape
papelist$pape
papelist$sd</pre>
```

Estimation of the Population Average Value in Randomized Experiments

Description

This function estimates the Population Average Value. The details of the methods for this design are given in Imai and Li (2019).

Usage

PAV(Tr, That, Y, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
That	A vector of the unit-level binary treatment that would have been assigned by the individualized treatment rule. If plim is specified, please ensure that the percentage of treatment units of That is lower than the budget constraint.
Y	A vector of the outcome variable of interest for each sample.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

pav	The estimated Population Average Value.
sd	The estimated standard deviation of PAV.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

Examples

Tr = c(1,0,1,0,1,0,1,0)
That = c(0,1,1,0,0,1,1,0)
Y = c(4,5,0,2,4,1,-4,3)
pavlist <- PAV(Tr,That,Y)
pavlist\$pav
pavlist\$pav</pre>

PAV

PAVcv

Description

This function estimates the Population Average Value. The details of the methods for this design are given in Imai and Li (2019).

Usage

PAVcv(Tr, That, Y, ind, centered = TRUE)

Arguments

Tr	A vector of the unit-level binary treatment receipt variable for each sample.
That	A matrix where the ith column is the unit-level binary treatment that would have been assigned by the individualized treatment rule generated in the ith fold. If plim is specified, please ensure that the percentage of treatment units of That is lower than the budget constraint.
Υ	The outcome variable of interest.
ind	A vector of integers (between 1 and number of folds inclusive) indicating which testing set does each sample belong to.
centered	If TRUE, the outcome variables would be centered before processing. This minimizes the variance of the estimator. Default is TRUE.

Value

A list that contains the following items:

pav	The estimated Population Average Value.
sd	The estimated standard deviation of PAV.

Author(s)

Michael Lingzhi Li, Operations Research Center, Massachusetts Institute of Technology <mlli@mit.edu>, http://mlli.mit.edu;

References

Imai and Li (2019). "Experimental Evaluation of Individualized Treatment Rules",

PAVcv

Examples

```
Tr = c(1,0,1,0,1,0,1,0)
That = matrix(c(0,1,1,0,0,1,1,0,0,1,1,0,0,1), nrow = 8, ncol = 2)
Y = c(4,5,0,2,4,1,-4,3)
ind = c(rep(1,4),rep(2,4))
pavlist <- PAVcv(Tr, That, Y, ind)
pavlist$pav
pavlist$pav</pre>
```

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