Package 'MatchIt'

January 9, 2018

Version 3.0.2

Date 2018-01-09

Title Nonparametric Preprocessing for Parametric Causal Inference

Description Selects matched samples of the original treated and control groups with similar covariate distributions -- can be used to match exactly on covariates, to match on propensity scores, or perform a variety of other matching procedures. The package also implements a series of recommendations offered in Ho, Imai, King, and Stuart (2007) <DOI:10.1093/pan/mpl013>.

Maintainer Kosuke Imai <kimai@Princeton.Edu>

Depends R (>= 2.6)

Imports MASS, Matching

Suggests cem, nnet, optmatch, rpart, mgcv, WhatIf, R.rsp, testthat, rgenoud

VignetteBuilder R.rsp

LazyLoad yes

LazyData yes

License GPL (>= 2)

URL http://gking.harvard.edu/matchit

RoxygenNote 6.0.1

NeedsCompilation no

Author Daniel Ho [aut], Kosuke Imai [aut, cre], Gary King [aut], Elizabeth Stuart [aut], Alex Whitworth [ctb]

Repository CRAN

Date/Publication 2018-01-09 17:41:26 UTC

R topics documented:

| et_matches | 2 |
|-------------|---|
| elp.matchit | 3 |
| matchit | |
| londe | 4 |
| natch.data | |
| natchit | 5 |
| ser.prompt | 8 |
| | • |
| | 9 |

Index

get_matches

Get matches from matchit object

Description

Get the resulting matches from a matchit model object. This function allows the user to extract the matches from the original dataset used in model building or from a new dataset that has a matching set of key column(s) (id_cols).

Usage

```
get_matches(object, model_frame, id_cols = NULL, newdata = NULL)
```

Arguments

| object | The 'matchit' class model object |
|-------------|---|
| model_frame | The 'data.frame' class object used in creation of object. |
| id_cols | A string indicating the ID for the datset used in the call to matchit. This can be used in combination with newdata to return the base dataset. Defaults to NULL. |
| newdata | A new data.frame object to extract matched observations from. Used in conjunction with id_cols. Defaults to NULL. |

Value

If newdata is NULL, a subset of model_frame containing the rows corresponding to the matched treatement and control observations with weights appended. If newdata is not NULL, an equivalent subset of newdata is returned.

help.matchit

Description

The help.matchit command launches html help for Matchit commands and supported methods. The full manual is available online at http://gking.harvard.edu/matchit.

Usage

```
help.matchit(object)
```

Arguments

object a character string representing a Matchit command or model. help.matchit("command") will take you to an index of Matchit commands and help.matchit("method") will take you to a list of matching methods. The following inputs are currently available: exact, nearest, subclass, full, optimal.

Author(s)

Daniel Ho <<daniel.ho@yale.edu>>; Kosuke Imai <<kimai@princeton.edu>>; Gary King <<king@harvard.edu>>; Elizabeth Stuart<<estuart@jhsph.edu>>

See Also

The complete document is available online at http://gking.harvard.edu/matchit.

is.matchit

Checks matchit Class

Description

Function that checks if the target object is a matchit object.

Usage

```
is.matchit(object)
```

Arguments

object any R object

Value

Returns TRUE if its argument has classs "matchit" among its classes and FALSE otherwise.

match.data

lalonde

Data from National Supported Work Demonstration and PSID, as analyzed by Dehejia and Wahba (1999).

Description

This is a subsample of the data from the treated group in the National Supported Work Demonstration (NSW) and the comparison sample from the Current Population Survey (CPS). This data was previously analyzed extensively by Lalonde (1986) and Dehejia and Wahba (1999).

Usage

data(lalonde)

Format

A data frame with 614 observations (185 treated, 429 control). There are 10 variables measured for each individual. "treat" is the treatment assignment (1=treated, 0=control). "age" is age in years. "educ" is education in number of years of schooling. "black" is an indicator for African-American (1=African-American, 0=not). "hispan" is an indicator for being of Hispanic origin (1=Hispanic, 0=not). "married" is an indicator for married (1=married, 0=not married). "nodegree" is an indicator for whether the individual has a high school degree (1=no degree, 0=degree). "re74" is income in 1974, in U.S. dollars. "re75" is income in 1975, in U.S. dollars. "re78" is income in 1978, in U.S. dollars.

References

Lalonde, R. (1986). Evaluating the econometric evaluations of training programs with experimental data. American Economic Review 76: 604-620.

Dehejia, R.H. and Wahba, S. (1999). Causal Effects in Nonexperimental Studies: Re-Evaluating the Evaluation of Training Programs. Journal of the American Statistical Association 94: 1053-1062.

match.data

Output Matched Data Sets

Description

match.data outputs matched data sets from matchit().

Usage

```
match.data(object, group="all", distance = "distance",
weights = "weights", subclass = "subclass")
```

matchit

Arguments

| object | The output object from matchit. This is a required input. |
|----------|--|
| group | This argument specifies for which matched group the user wants to extract the data. Available options are "all" (all matched units), "treat" (matched units in the treatment group), and "control" (matched units in the control group). The default is "all". |
| distance | This argument specifies the variable name used to store the distance measure. The default is "distance". |
| weights | This argument specifies the variable name used to store the resulting weights from matching. The default is "weights". |
| subclass | This argument specifies the variable name used to store the subclass indicator. The default is "subclass". |

Value

Returns a subset of the original data set sent to matchit(), with just the matched units. The data set also contains the additional variables distance, weights, and subclass. The variable distance gives the estimated distance measure, and weights gives the weights for each unit, generated in the matching procedure. The variable subclass gives the subclass index for each unit (if applicable). See the http://gking.harvard.edu/matchit/ for the complete documentation and type demo(match.data) at the R prompt to see a demonstration of the code.

Author(s)

Daniel Ho <daniel.ho@yale.edu>; Kosuke Imai <kimai@princeton.edu>; Gary King <king@harvard.edu>; Elizabeth Stuart <estuart@jhsph.edu>

See Also

Please use help.matchit to access the matchit reference manual. The complete document is available online at http://gking.harvard.edu/matchit.

matchit

MatchIt: Matching Software for Causal Inference

Description

matchit is the main command of the package *MatchIt*, which enables parametric models for causal inference to work better by selecting well-matched subsets of the original treated and control groups. MatchIt implements the suggestions of Ho, Imai, King, and Stuart (2004) for improving parametric statistical models by preprocessing data with nonparametric matching methods. MatchIt implements a wide range of sophisticated matching methods, making it possible to greatly reduce the dependence of causal inferences on hard-to-justify, but commonly made, statistical modeling assumptions. The software also easily fits into existing research practices since, after preprocessing with MatchIt, researchers can use whatever parametric model they would have used without MatchIt, but produce inferences with substantially more robustness and less sensitivity

to modeling assumptions. Matched data sets created by MatchIt can be entered easily in Zelig (http://gking.harvard.edu/zelig) for subsequent parametric analyses. Full documentation is available online at http://gking.harvard.edu/matchit, and help for specific commands is available through help.matchit.

Usage

Arguments

| formula | This argument takes the usual syntax of R formula, treat ~ $x1 + x2$, where treat is a binary treatment indicator and x1 and x2 are the pre-treatment co-variates. Both the treatment indicator and pre-treatment covariates must be contained in the same data frame, which is specified as data (see below). All of the usual R syntax for formula works. For example, x1:x2 represents the first order interaction term between x1 and x2, and I(x1^2) represents the square term of x1. See help(formula) for details. |
|-----------------|--|
| data | This argument specifies the data frame containing the variables called in formula. |
| method | This argument specifies a matching method. Currently, "exact" (exact match- ing), "full" (full matching), "genetic" (genetic matching), "nearest" (near- est neighbor matching), "optimal" (optimal matching), and "subclass" (sub- classification) are available. The default is "nearest". Note that within each of these matching methods, <i>MatchIt</i> offers a variety of options. |
| distance | This argument specifies the method used to estimate the distance measure. The default is logistic regression, "logit". A variety of other methods are available. |
| distance.optior | IS |
| | This optional argument specifies the optional arguments that are passed to the model for estimating the distance measure. The input to this argument should be a list. |
| discard | This argument specifies whether to discard units that fall outside some measure of support of the distance score before matching, and not allow them to be used at all in the matching procedure. Note that discarding units may change the quantity of interest being estimated. The options are: "none" (default), which discards no units before matching, "both", which discards all units (treated and control) that are outside the support of the distance measure, "control", which discards only control units outside the support of the distance measure of the treated units, and "treat", which discards only treated units outside the support of the distance measure of the control units. |
| reestimate | This argument specifies whether the model for distance measure should be re- estimated after units are discarded. The input must be a logical value. The default is FALSE. |
| | Additional arguments to be passed to a variety of matching methods. |

matchit

Details

The matching is done using the matchit(treat ~ X, ...) command, where treat is the vector of treatment assignments and X are the covariates to be used in the matching. There are a number of matching options, detailed below. The full syntax is matchit(formula, data=NULL, discard=0, exact=FALSE, replace A summary of the results can be seen graphically using plot(matchitobject), or numerically using summary(matchitobject). print(matchitobject) also prints out the output.

Value

| call | The original matchit call. |
|--------------|---|
| formula | The formula used to specify the model for estimating the distance measure. |
| model | The output of the model used to estimate the distance measure. summary(m.out\$model) will give the summary of the model where m.out is the output object from matchit. |
| match.matrix | An n_1 by ratio matrix where the row names, which can be obtained through row.names(match.matrix), represent the names of the treatment units, which come from the data frame specified in data. Each column stores the name(s) of the control unit(s) matched to the treatment unit of that row. For example, when the ratio input for nearest neighbor or optimal matching is specified as 3, the three columns of match.matrix represent the three control units matched to one treatment unit). NA indicates that the treatment unit was not matched. |
| discarded | A vector of length $n\$ that displays whether the units were ineligible for matching due to common support restrictions. It equals TRUE if unit <i>i</i> was discarded, and it is set to FALSE otherwise. |
| distance | A vector of length n with the estimated distance measure for each unit. |
| weights | A vector of length n that provides the weights assigned to each unit in the match- ing process. Unmatched units have weights equal to 0 . Matched treated units have weight 1. Each matched control unit has weight proportional to the number of treatment units to which it was matched, and the sum of the control weights is equal to the number of uniquely matched control units. |
| subclass | The subclass index in an ordinal scale from 1 to the total number of subclasses as specified in subclass (or the total number of subclasses from full or exact matching). Unmatched units have NA. |
| q.cut | The subclass cut-points that classify the distance measure. |
| treat | The treatment indicator from data (the left-hand side of formula). |
| х | The covariates used for estimating the distance measure (the right-hand side of formula). |
| nn | A basic summary table of matched data (e.g., the number of matched units) |

Author(s)

Daniel Ho <daniel.ho@yale.edu>; Kosuke Imai <kimai@princeton.edu>; Gary King <king@harvard.edu>; Elizabeth Stuart<estuart@jhsph.edu>

References

Daniel Ho, Kosuke Imai, Gary King, and Elizabeth Stuart (2007). Matching as Nonparametric Preprocessing for Reducing Model Dependence in Parametric Causal Inference. Political Analysis 15(3): 199-236. http://gking.harvard.edu/files/abs/matchp-abs.shtml

See Also

Please use help.matchit to access the matchit reference manual. The complete document is available online at http://gking.harvard.edu/matchit.

user.prompt

Pause in demo files

Description

Use user.prompt while writing demo files to force users to hit return before continuing.

Usage

user.prompt()

Author(s)

Olivia Lau <olau@fas.harvard.edu>

See Also

readline

Examples

Not run: user.prompt()

End(Not run)

Index

*Topic datasets
 lalonde, 4
*Topic documentation
 help.matchit, 3
*Topic environment
 matchit, 5
*Topic file
 user.prompt, 8
*Topic methods
 match.data, 4
get_matches, 2

help.matchit, 3

is.matchit, 3

lalonde, 4

match.data,4
MatchIt(matchit),5
Matchit(matchit),5
matchit,2,5

user.prompt, 8