Package: SAMUR (via r-universe)

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Type Package
Title Stochastic Augmentation of Matched Data Using Restriction Methods
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Description Augmenting a matched data set by generating multiple stochastic, matched samples from the data using a multi-dimensional histogram constructed from dropping the input matched data into a multi-dimensional grid built on the full data set. The resulting stochastic, matched sets will likely provide a collectively higher coverage of the full data set compared to the single matched set. Each stochastic match is without duplication, thus allowing downstream validation techniques such as cross-validation to be applied to each set without concern for overfitting.
License GPL (>= 2)
Imports Matching
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samur	Stochastic	Augmentation	of	Matched	Datasets	Using	Restriction	
	Methods							

Description

This function generates multiple subsets of the data in which the distribution of covariates is balanced across treatment groups. It works by binning the output of a base matching algorithm into a multidimensional histogram, and drawing - without replacement - from the full data set according to the histogram. This leads to higher data coverage across multiple matched subsets without duplication of cases within each subset.

Usage

```
samur(
  formula, data
  , matched.subset = 1:nrow(data)
  , nsmp = 100
  , use.quantile = TRUE, breaks = 10
  , replace = length(unique(matched.subset)) < length(matched.subset)
  )
## S3 method for class 'samur'
print(x, ...)</pre>
```

Arguments

formula	Formula expression used to describe the treatment variable (lhs) and covariates used during matching (rhs).
data	Data frame containing the treatment variables and matched covariates as specified in the formula.
matched.subset	An integer vector representing the indexes of a subset of data that is the output of a base matching algorithm. It cannot contain duplicate values.
nsmp	Number of stochastically matched subsets to generate.
use.quantile	Should numeric covariates be binned using quantiles (TRUE) or not.
breaks	number of breaks to use in binning numeric covariates.
replace	Boolean flag indicating whether or not to perform sampling with replacement.
X	An object of class samur, typically the output of function samur.
	Arguments passed to/from other methods.

Value

An object of class samur, a matrix of size length(matched.subset) by nsmp, where each column is a matched subset without case duplication. It also has the following attributes:

call Copy of function call.

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formula Formula passed to the function.

mdg Multi-dimensional grid used for binning the matched data subsets.

mdh Multi-dimensional histogram resulting frm binning data[matched.subset,]

according to the grid specified in mdg.

data Copy of data frame passed to the function.

Author(s)

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See Also

```
summary.samur
```

Examples

```
## Not run:
library(SAMUR)
library(Matching)
data(lalonde)
myformula <- treat ~ age + educ</pre>
myglm <- glm(myformula, lalonde, family="binomial")</pre>
X <- myglm$fitted.values</pre>
# using M=1 and replace=F to ensure no duplication
bimatch <- Match(Tr = lalonde$treat, X = myglm$fitted.values</pre>
  , M = 1, replace = F, caliper = 0.25)
idx <- c(bimatch$index.control, bimatch$index.treated)</pre>
my.samur <- samur(formula = myformula, data = lalonde</pre>
  , matched.subset = idx, nsmp = 100
  , breaks = 10, use.quantile = TRUE)
summary(my.samur, nboots = 500)
## End(Not run)
```

summary.samur

Summarizing Output of SAMUR Augmentation Function

Description

summary method for class "samur".

Usage

```
## S3 method for class 'samur'
summary(object, ...)
## S3 method for class 'summary.samur'
print(x, ...)
```

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Arguments

object An object of class "samur", usually the result of a call to samur.

x An object of class "summary.samur", usually the result of a call to summary.samur.

Further arguments to be passed to/from other methods. Current implementation

of summary. samur passes arguments to MatchBalance function from Matching $\,$

package.

Value

A list with the following elements:

min.pval.new A vector of length equal to number of samples (nsmp) generated by samur, each

representing the minimum p-value from all univariate tests performed by the underlying function MatchBalance. It also has an attributed named min.pval.orig,

 $containing \ a \ similar \ number \ for \ the \ original \ matched \ subset, i.e. \ data \verb|[matched.subset||, i.e.]| \\$

].

min.pval.orig Same number as above, but for original matched subset.

coverage.new Percent of cases from full data set covered among all stochastic, matched sam-

ples.

coverage.orig Same as above, calculated for the original matched subset.

Note

All t-tests used for p-value calculations are "not" paired, since the philosophy of stochastic augmentation relaxes the notion of one-to-one matching.

Author(s)

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See Also

samur, MatchBalance

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