Package 'TriMatch'

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TriMatch-package
as.data.frame.list
balance.plot
boxdiff.plot
covariateBalance
data.frame.to.list
distance.euclid
distances.plot
loess3.plot

2 TriMatch-package

naex		4
ndex		27
	unmatched	26
	tutoring	25
	trips	23
	trimatch.apply2	23
	trimatch	21
	summary.unmatched	
	summary.triangle.psa	20
	summary.triangle.matches	19
	summary.balance.plots	19
	star	18
	segment2	18
	segment1	18
	print.trimatch.summary	17
	print.triangle.plot	17
	print.balance.plots	16
	plot.triangle.psa	15
	plot.triangle.matches	14
	plot.balance.plots	
	perpPt	
	parallel.plot	
	OneToN	
	nmes	
	multibalance.plot	
	merge.triangle.psa	
	maximumTreat	

Description

TriMatch-package

This packages provides functions to estimate and visualize propensity score analyses including matching for non-binary treatments.

Propensity Score Analysis for Non-Binary Treatments

Author(s)

Jason Bryer < jason@bryer.org>

See Also

PSAgraphics multilevelPSA

as.data.frame.list 3

as.data.frame.list *Convert a list of vectors to a data frame.*

Description

This function will convert a list of vectors to a data frame. This function will handle three different types of lists of vectors. First, if all the elements in the list are named vectors, the resulting data frame will have have a number of columns equal to the number of unique names across all vectors. In cases where some vectors do not have names in other vectors, those values will be filled with NA.

Usage

```
## S3 method for class 'list'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

x a list to convert to a data frame.

row.names a vector equal to length(x) corresponding to the row names. If NULL, the row

names will be set to names (x).

optional not used.

... other parameters passed to data.frame.

Details

The second case is when all the vectors are of the same length. In this case, the resulting data frame is equivalent to applying rbind across all elements.

The third case handled is when there are varying vector lengths and not all the vectors are named. This condition should be avoided. However, the function will attempt to convert this list to a data frame. The resulting data frame will have a number of columns equal to the length of the longest vector. For vectors with length less than this will fill the row with NAs. Note that this function will print a warning if this condition occurs.

Value

a data frame.

Author(s)

Jason Bryer jason@bryer.org

References

http://stackoverflow.com/questions/4227223/r-list-to-data-frame

4 balance.plot

Examples

```
test1 <- list( c(a='a',b='b',c='c'), c(a='d',b='e',c='f'))
   as.data.frame(test1)
    test2 <- list( c('a', 'b', 'c'), c(a='d',b='e',c='f'))
    as.data.frame(test2)
    test3 <- list('Row1'=c(a='a',b='b',c='c'), 'Row2'=c(var1='d',var2='e',var3='f'))
    as.data.frame(test3)
    ## Not run:
    #This will print a warning.
    test4 <- list('Row1'=letters[1:5], 'Row2'=letters[1:7], 'Row3'=letters[8:14])</pre>
   as.data.frame(test4)
## End(Not run)
    test5 <- list(letters[1:10], letters[11:20])</pre>
   as.data.frame(test5)
    ## Not run:
    #This will throw an error.
    test6 <- list(list(letters), letters)</pre>
    as.data.frame(test6)
## End(Not run)
```

balance.plot

Balance plot for the given covariate.

Description

If the covariate is numeric, boxplots will be drawn with red points for the mean and green error bars for the standard error. For non-numeric covariates a barplot will be drawn.

Usage

```
balance.plot(x, covar, model, nstrata = attr(attr(tmatch, "triangle.psa"),
   "nstrata"), label = "Covariate", ylab = "", xlab = NULL, se.ratio = 2,
   print = TRUE, legend.position = "top", x.axis.labels,
   x.axis.angle = -45, ...)
```

Arguments

x results from trimatch.

covar vector of the covariate to check balance of.

model an integer between 1 and 3 indicating from which model the propensity scores

will be used.

boxdiff.plot 5

number of strata to use. nstrata label for the legend. label ylab label of the y-axis. label of the x-axis. xlab se.ratio a multiplier for how large standard error bars will be. print print the output if the Freidman Rank Sum Test and repeated measures ANOVA (for continuous variables). legend.position the position of the legend. See theme. x.axis.labels labels for the x-axis. x.axis.angle angle for x-axis labels.

Details

. . .

A Friedman rank sum test will be performed for all covariate types, printed, and stored as an attribute to the returned object named friedman. If a continuous covariate a repeated measures ANOVA will also be performed, printed, and returned as an attribute named rmanova.

parameters passed to plot.balance.plots.

Value

a ggplot2 figure or a list of ggplot2 figures if covar is a data frame.

boxdiff.plot Returns a ggplot2 box plot of the differences.

Description

A boxplot of differences between each pair of treatments.

Usage

```
boxdiff.plot(tmatch, out, plot.mean = TRUE, ordering = attr(tmatch,
   "match.order"), ci.width = 0.5)
```

Arguments

tmatch	the results from trimatch.
out	a vector of the outcome measure of interest.
plot.mean	logical indicating whether the means should be plotted.
ordering	specify the order for doing the paired analysis, that is analysis will be conducted as: ordering[1] - ordering[2], ordering[1] - ordering[3], and ordering[2] - ordering[3].
ci.width	the width for the confidence intervals.

6 covariateBalance

Value

a ggplot2 boxplot of the differences.

covariateBalance Calculate covariate effect size differences before and after stratification.

Description

This function is modified from the cv.bal.psa function in the PSAgrpahics package.

Usage

```
covariateBalance(covariates, treatment, propensity, strata = NULL,
  int = NULL, tree = FALSE, minsize = 2, universal.psd = TRUE,
  trM = 0, absolute.es = TRUE, trt.value = NULL, use.trt.var = FALSE,
  verbose = FALSE, xlim = NULL, plot.strata = TRUE, ...)
```

Arguments

covariates	dataframe of interest
treatment	binary vector of 0s and 1s (necessarily? what if character, or 1, 2?)
propensity	PS scores from some method or other.
strata	either a vector of strata number for each row of covariate, or one number n in which case it is attempted to group rows by ps scores into n strata of size approximately 1/n. This does not seem to work well in the case of few specific propensity values, as from a tree.
int	either a number m used to divide [0,1] into m equal length subintervals, or a vector of cut points between 0 an 1 defining the subintervals (perhaps as suggested by loess.psa). In either case these subintervals define strata, so strata can be of any size.
tree	logical, if unique ps scores are few, as from a recursively partitioned tree, then TRUE will force each ps value to define a stratum.
minsize	smallest allowable stratum-treatment size. If violated, strata is removed.
universal.psd	If 'TRUE', forces standard deviations used to be unadjusted for stratification.
trM	trimming proportion for mean calculations.
absolute.es	logical, if 'TRUE' routine uses absolute values of all effect sizes.
trt.value	allows user to specify which value is active treatment, if desired.
use.trt.var	logical, if true then Rubin-Stuart method using only treatment variance with be used in effect size calculations.
verbose	logical, controls output that is visibly returned.
xlim	limits for the x-axis.
plot.strata	logical indicating whether to print strata.
	currently unused.

data.frame.to.list 7

Details

Note: effect sizes are calculated as treatment 1 - treatment 0, or treatment B - treatment A.

Author(s)

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data.frame.to.list

Convert a list of vectors to a data frame.

Description

This function will convert a list of vectors to a data frame. This function will handle three different types of lists of vectors. First, if all the elements in the list are named vectors, the resulting data frame will have have a number of columns equal to the number of unique names across all vectors. In cases where some vectors do not have names in other vectors, those values will be filled with NA.

Usage

```
data.frame.to.list(...)
```

Arguments

... other parameters passed to data.frame.

Details

The second case is when all the vectors are of the same length. In this case, the resulting data frame is equivalent to applying rbind across all elements.

The third case handled is when there are varying vector lengths and not all the vectors are named. This condition should be avoided. However, the function will attempt to convert this list to a data frame. The resulting data frame will have a number of columns equal to the length of the longest vector. For vectors with length less than this will fill the row with NAs. Note that this function will print a warning if this condition occurs.

Value

a data frame.

References

http://stackoverflow.com/questions/4227223/r-list-to-data-frame

8 distances.plot

|--|

Description

This method uses a simple Euclidean distance calculation for determining the distances between two matches. That is, lps1 - ps2l.

Usage

```
distance.euclid(x, grouping, id, groups, caliper, nmatch = Inf)
```

Arguments

X	vector of propensity scores.
grouping	vector or factor identifying group membership.
id	vector corresponding to unique identifer for each element in x and grouping.
groups	vector of length two indicating the unique groups to calculate the distance between. The first element will be the rows, the second columns.
caliper	a scaler indicating the caliper to use for matching within each step.
nmatch	number of smallest distances to retain.

Value

a list of length equal to x. Each element of the list is a named numeric vector where the values correspond to the distance and the name to the id.

distances.plot	Barplot for the sum of distances.

Description

Barplot for the sum of distances.

Usage

```
distances.plot(tmatch, caliper = 0.25, label = FALSE)
```

Arguments

tmatch	the results of trimatch.
caliper	a vector indicating where vertical lines should be drawn as a factor of the standard deviation. Rosenbaum and Rubin (1985) suggested one quarter of one standard deviation.
label	label the bars that exceed the minimum caliper.

loess3.plot

See Also

triangle.match

loess3.plot

Loess plot for matched triplets.

Description

This function will create a ggplot2 figure with propensity scores on the x-axis and the outcome on the y-axis. Three Loess regression lines will be plotted based upon the propensity scores from model. Since each model produces propensity scores for two of the three groups, the propensity score for the third group in each matched triplet will be the mean of the other two. If model is not specified, the default will be to use the model that estimates the propensity scores for the first two groups in the matching order.

Usage

```
loess3.plot(tmatch, outcome, model, ylab = "Outcome",
  plot.connections = FALSE, connections.color = "black",
  connections.alpha = 0.2, plot.points = geom_point, points.alpha = 0.1,
  points.palette = "Dark2", ...)
```

Arguments

tmatch the results of trimatch.

outcome a vector representing the outcomes.

model an integer between 1 and 3 indicating from which model the propensity scores

will be used.

ylab the label for the y-axis.

plot.connections

boolean indicating whether lines will be drawn connecting each matched triplet.

connections.color

the line color of connections.

connections.alpha

number between 0 and 1 representing the alpha levels for connection lines.

plot.points a ggplot2 function for plotting points. Usually geom_point or geom_jitter.

If NULL no points will be drawn.

points.alpha number between 0 and 1 representing the alpha level for the points.

points.palette the color palette to use. See scale_colour_brewer and http://colorbrewer2.

org/ for more information.

... other parameters passed to geom_smooth and stat_smooth.

Value

```
a ggplot2 figure.
```

maximumTreat	This method will return at least one treatment from groups one and two within the caliper.

Description

This method will attempt to return enough rows to use each treatment (the first two groups in the matching order) at least once. Assuming treat1 is the first group in the match order and treat2 the second, all duplicate treat1 rows are removed. Next, all treat2 units not in present in after removing duplicate treat1 units are identified. For each of those treat2 units, the matched triplet with the smallest overall distances where treat2 is one of the mathched units is retained.

Usage

```
maximumTreat(tmatch, ...)
```

Arguments

tmatch initial results from trimatch that contains all possible matches within the spec-

ified caliper.

... currently unused.

merge.triangle.matches

Merges outcomes with the matched set.

Description

The y parameter should be a subset of the original data used.

Usage

```
## S3 method for class 'triangle.matches' merge(x, y, ...)
```

Arguments

x the result of trimatch

y another data frame or vector to merge with.

... unused

Value

x with the additional column(s) added.

merge.triangle.psa 11

merge.triangle.psa

Merges covariate(s) with the results of trips.

Description

The y parameter should be a subset of the original data used.

Usage

```
## S3 method for class 'triangle.psa' merge(x, y, ...)
```

Arguments

x the result of trips

y another data frame or vector to merge with.

... unused

Value

x with the additional column(s) added.

multibalance.plot

Multiple covariate balance assessment plot.

Description

A graphic based upon cv.bal.psa function in the PSAgraphics package. This graphic plots the effect sizes for multiple covariates before and after propensity score adjustment.

Usage

```
multibalance.plot(tpsa, tmatch, grid = TRUE, cols)
```

Arguments

tpsa results of trips.
tmatch results of trimatch.

grid if TRUE, then a grid of three plots for each model will be displayed.

cols character vector of covariates (i.e. column names) from the original data to

include in the plot. By default all covariates used in the logistic regression model

are used.

Value

```
a ggplot2 figure.
```

12 OneToN

nmes

Results from the 1987 National Medical Expenditure Study

Description

This file was originally prepared by Anders Corr (corr@fas.harvard.edu) who reports on December 8, 2007 that the resulting numbers closely match with those reported in the published article. It was later modified by Jason Bryer (jason@bryer.org) to an R data object to be included in this package. See http://imai.princeton.edu/research/pscore.html for more information

Format

a data frame with 9,708 observations of 12 variables.

Author(s)

United States Department of Health and Human Services. Agency for Health Care Policy and Research

Source

http://imai.princeton.edu/research/pscore.html

References

National Center For Health Services Research, 1987. National Medical Expenditure Survey. Methods II. Questionnaires and data collection methods for the household survey and the Survey of American Indians and Alaska Natives. National Center for Health Services Research and Health Technology Assessment.

Imai, K., & van Dyk, D.A. (2004). Causal Inference With General Treatment Regimes: Generalizing the Propensity Score, Journal of the American Statistical Association, 99(467), pp. 854-866.

Elizabeth Johnson, E., Dominici, F., Griswold, M., & Zeger, S.L. (2003). Disease cases and their medical costs attributable to smoking: An analysis of the national medical expenditure survey. Journal of Econometrics, 112.

OneToN

This method will use a M1-to-M2-to-1 matching.

Description

In this method, M2 corresponds to the number of times a treat1 unit can be matched with a treat2 unit. The M1 parameter corresponds to the number of times a treat1 unit can be used in total.

Usage

```
OneToN(tmatch, M1 = 2, M2 = 1, ...)
```

parallel.plot 13

Arguments

tmatch	initial results from trimatch that contains all possible matches within the specified caliper.
M1	a scaler indicating the number of unique subjects in group one to retain. This applies only to the first group in the matching order.
M2	a scaler indicating the number of unique matches to retain. This applies to the first two groups in the matching order.
	currently unused

... currently unused.

parallel.plot

Parallel coordinate plot for the three groups and dependent variable.

Description

Creates a ggplot2 figure of a parallel coordinate plot.

Usage

```
parallel.plot(tmatch, outcome)
```

Arguments

tmatch results from trimatch.
outcome vector of the outcome

perpPt

Internal method for plotting. Finds a point d distance from x, y

Description

Internal method for plotting. Finds a point d distance from x, y

Usage

```
perpPt(x, y, d = 0.05)
```

Arguments

Χ	x coordinate
у	y coordinate
d	the distance

14 plot.triangle.matches

plot.balance.plots

Prints a grid of balance plots.

Description

Prints a grid of balance plots.

Usage

```
## $3 method for class 'balance.plots'
plot(x, rows, cols, byrow = TRUE,
    plot.sequence = seq_along(bplots), ...)
```

Arguments

x the results of balance.plot when a data frame is specified.

rows if covar is a data frame of covariates, the number of rows in the grid of figures.

cols if covar is a data frame of covariates, the number of columns in the grid of

figures.

byrow if TRUE (default), plots will be drawn by rows, otherwise by columns.

plot. sequence (or subset) of plots to draw.

... currently unused.

plot.triangle.matches Triangle plot drawing matched triplets.

Description

This plot function adds a layer to plot.triangle.psa drawing matched triplets. If p is supplied, this function will simply draw on top of the pre-existing plot, otherwise plot.triangle.psa will be called first.

Usage

```
## S3 method for class 'triangle.matches'
plot(x, sample = 0.05,
  rows = sample(nrow(tmatch), nrow(tmatch) * sample), line.color = "black",
  line.alpha = 0.5, point.color = "black", point.size = 3, p, ...)
```

plot.triangle.psa 15

Arguments

X	matched triplets from link{triangle.match}.
sample	an number between 0 and 1 representing the percentage of matched triplets to draw.
rows	an integer vector corresponding to the rows in tmatch to draw.
line.color	the line color.
line.alpha	the alpha for the lines.
point.color	color of matched triplet points.
point.size	point size for matched triplets.
p	a ggplot to add the match lines. If NULL, then plot.triangle.psa.
	other parameters passed to plot.triangle.psa.

Details

If this function calls plot.triangle.psa, it will only draw line segments and points for those data rows that were used in the matching procedure. That is, data elements not matched will be excluded from the figure. To plot all segments and points regardless if used in matching, set p = plot(tpsa).

Value

```
a ggplot2 graphic.
```

See Also

```
plot.triangle.psa
triangle.match
```

Description

Triangle plot showing the fitted values (propensity scores) for three different models.

Usage

```
## S3 method for class 'triangle.psa'
plot(x, point.alpha = 0.3, point.size = 1.5,
  legend.title = "Treatment", text.size = 4, draw.edges = FALSE,
  draw.segments = TRUE, edge.alpha = 0.2, edge.color = "grey",
  edge.labels = c("Model 1", "Model 2", "Model 3"), sample = c(1), ...)
```

16 print.balance.plots

Arguments

x the results from trips.
point.alpha alpha level for points.

point.size point size.

legend.title title for the legend.

text.size text size.

draw.edges draw edges of the triangle.

draw.segments draw segments connecting points across two models.

edge.alpha alpha level for edges if drawn.
edge.color the color for edges if drawn.

edge.labels the labels to use for each edge of the triangle.

sample a vector of length 1 or 3 representing the sample of points to plot. The position of

each element corresponds to the groups as returned by attr(tpsa, 'groups'). If equal to one, all points will be plotted. Values less than one will plot a percentage of points. Values greater than one exactly that number of points will be

plotted.

... currently unused.

Value

ggplot2 figure

See Also

triangle.psa

print.balance.plots

Print the results of balance.plot *for a data frame of covariates.*

Description

Print the results of balance.plot for a data frame of covariates.

Usage

```
## S3 method for class 'balance.plots'
print(x, ...)
```

Arguments

x the results of balance.plot when a data frame is specified.

... parameters passed to plot.balance.plots and summary.balance.plots.

print.triangle.plot 17

```
print.triangle.plot Print method for plot.triangle.psa. The primary purpose is to suppress the "Removed n rows containing missing values" warning printed by ggplot2.
```

Description

Print method for plot.triangle.psa. The primary purpose is to suppress the "Removed n rows containing missing values" warning printed by ggplot2.

Usage

```
## S3 method for class 'triangle.plot'
print(x, ...)
```

Arguments

```
x a plot from plot.triangle.psa.... other parameters passed to ggplot2.
```

```
print.trimatch.summary
```

Prints the results of summary.triangle.matches.

Description

This is an S3 generic function to print the results of summary.triangle.matches.

Usage

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

Arguments

```
x results of summary.triangle.matches.
```

```
... multiple results of summary.triangle.matches. These must be named. For example, "Method 1" = summary(tmath, outcome).
```

18 star

segment1

Internal method for plotting. Position along the left side segment

Description

Internal method for plotting. Position along the left side segment

Usage

```
segment1(d)
```

Arguments

d

the distance

segment2

Internal method for plotting. Position along the right side segment

Description

Internal method for plotting. Position along the right side segment

Usage

```
segment2(d)
```

Arguments

d

the distance

star

Returns significance level.

Description

Returns the significance level as stars, or NA if a non-numeric value is passed in.

Usage

star(x)

Arguments

Х

p-value.

summary.balance.plots 19

summary.balance.plots *Prints a summary table of the test statistics of each balance plot.*

Description

The balance.plot function will create a grid of balance plots if a data frame is provided. The returned object is a list of ggplot2 figures with the statistical tests (i.e. Friedmen Rank Sum tests and if a continuous variable, repeated measures ANOVA as well) saved as attributes. This function will return a data frame combining all of those results.

Usage

```
## S3 method for class 'balance.plots'
summary(object, ...)
```

Arguments

```
object the results of balance.plot when a data frame is specified.
... currently unused.
```

Value

a data frame

```
summary.triangle.matches
```

Provides a summary of the matched triplets including analysis of outcome measure if provided.

Description

If an outcome measure is provided this function will perform a Freidman Rank Sum Test and repeated measures ANOVA. If either test has a statistically significant difference (as determined by the value of the p parameter), a Pairwise Wilcoxon Rank Sum Test will also be provided.

Usage

```
## S3 method for class 'triangle.matches'
summary(object, outcome, p = 0.05,
   ordering = attr(object, "match.order"), ...)
```

20 summary.triangle.psa

Arguments

object result of trimatch.

outcome vector representing the outcome measure.

p threshold of the p value to perform a

ordering specify the order for doing the paired analysis, that is analysis will be conducted as: ordering[1] - ordering[2], ordering[1] - ordering[3], and ordering[2] - ordering[3].

... parameters passed to other statistical tests.

Value

a trimatch.summary object.

See Also

```
friedman.test, ezANOVA, pairwise.wilcox.test
```

Description

The trips function estimates three separate logistic regression models for each pair of groups. This function will print a combined table of the three summaries.

Usage

```
## S3 method for class 'triangle.psa'
summary(object, ...)
```

Arguments

object the results of trips.... currently unused.

summary.unmatched 21

summary.unmatched

Provides a summary of unmatched subjects.

Description

Will return as a list and print the percentage of total unmatched rows and percent by treatment.

Usage

```
## S3 method for class 'unmatched'
summary(object, digits = 3, ...)
```

Arguments

object results of unmatched digits number of digits to print. . . . currently unused.

Value

a list of summary results.

trimatch

Creates matched triplets.

Description

Create matched triplets by minimizing the total distance between matched triplets within a specified caliper.

Usage

```
trimatch(tpsa, caliper = 0.25, nmatch = c(15), match.order, exact, method = maximumTreat, ...)
```

Arguments

tpsa the results from trips

caliper a vector of length one or three indicating the caliper to use for matching within

each step. This is expressed in standardized units such that .25 means that matches must be within .25 of one standard deviation to be kept, otherwise the

match is dropped.

22 trimatch

nmatch number of closest matches to retain before moving to next edge. This can be Inf in which case all matches within the caliper will be retained through to the next step. For large datasets, evaluating all possible matches within the caliper could be time consuming. match.order character vector of length three indicating the order in which the matching algorithm will processes. The default is to use start with the group the middle number of subjects, followed by the smallest, and then the largest. a vector or data frame of representing covariates for exact matching. That is, exact matched triplets will first be matched exactly on these covariates before evaluating distances. method This is a function that specifies which matched triplets will be retained. If NULL, all matched triplets within the specified caliper will be returned (equivalent to caliper matching in two group matching). The default is maximumTreat that

attempts include each treatment at least once. Another option is OneToN which mimicks the one-to-n matching where treatments are matched to multiple control units.

other parameters passed to method. . . .

Details

The trips function will estimate the propensity scores for three models. This method will then find the best matched triplets based upon minimizing the summed differences between propensity scores across the three models. That is, the algorithm works as follows:

- The first subject from model 1 is selected.
- The nmatch[1] smallest distances are selected using propensity scores from model 1.
- For each of the matches identified, the subjects propensity score from model 2 is retrieved.
- The nmatch[2] smallest distances are selected using propensity score from model 3.
- For each of those matches identified, the subjects propensity score from model 2 is retrieved.
- The distances is calculated from the first and last subjects propensity scores from model 2.
- The three distances are summed.
- The triplet with the smallest overall distance is selected and returned.

Examples

```
## Not run:
data(turoing)
formu <- ~ Gender + Ethnicity + Military + ESL + EdMother + EdFather + Age +
     Employment + Income + Transfer + GPA
tpsa <- trips(tutoring, tutoring$treat, formu)</pre>
tmatch <- trimatch(tpsa, status=FALSE)</pre>
## End(Not run)
```

trimatch.apply2 23

trimatch.apply2	Recursive function to find possible matched triplets using the apply functions.

Description

Internal method. This version does not use the exact matching. Instead, this function should be called separately for each grouping.

Usage

```
trimatch.apply2(tpsa, caliper, nmatch, match.order, sd1, sd2, sd3)
```

Arguments

tpsa	the results from trips
caliper	a vector of length one or three indicating the caliper to use for matching within each step. This is expressed in standardized units such that .25 means that matches must be within .25 of one standard deviation to be kept, otherwise the match is dropped.
nmatch	number of closest matches to retain before moving to next edge. This can be Inf in which case all matches within the caliper will be retained through to the next step. For large datasets, evaluating all possible matches within the caliper could be time consuming.
match.order	character vector of length three indicating the order in which the matching algorithm will processes. The default is to use start with the group the middle number of subjects, followed by the smallest, and then the largest.
sd1	standard deviation for propensity scores from model 1.
sd2	standard deviation for propensity scores from model 2.
sd3	standard deviation for propensity scores from model 3.

trips	Estimates propensity scores for three groups

Description

The propensity score is

$$e(X) = P(W = 1|X)$$

This function will estimate the propensity scores for each pair of groups (e.g. two treatments and one control).

24 trips

Usage

```
trips(thedata, treat, formu = ~., groups = unique(treat), nstrata = 5,
  method = "logistic", ...)
```

Arguments

thedata	the data frame.
treat	vector or factor indicating the treatment/control assignment for the data. Length must be equal to $nrow(the data)$.
formu	the logistic regression formula. Note that the dependent variable should not be specified and will be modified.
groups	a vector of exactly length three corresponding the values in treat for each control/treatment.
nstrata	the number of strata marks to plot on the edge.
method	the method to use to estimate the propensity scores. Current options are logistic or randomForest.
	other parameters passed to glm.

Details

$$PS_1 = e(X_{T_1C}) = Pr(z = 1|X_{T_1C})$$

$$PS_2 = e(X_{T_2C}) = Pr(z = 1|X_{T_2C})$$

$$PS_3 = e(X_{T_2T_1}) = Pr(z = 1|X_{T_2T_1})$$

Examples

tutoring 25

tutoring	Results from a study examining the effects of tutoring services on course grades.

Description

- treat Treatment indicator.
- Course The course id the student was enrolled in.
- Grade The course grade the student earned (4=A, 3=B, 2=C, 1=D, 0=F or W).
- Gender Gender of the student.
- Ethnicity Ethnicity of the student, either White, Black, or Other.
- Military Is the student an active military student.
- ESL English second language student.
- EdMother Education level of the mother (1 = did not finish high school; 2 = high school grad; 3 = some college; 4 = earned associate degree; 5 = earned baccalaureate degree; 6 = Earned Master's degree; 7 = earned doctorate).
- EdFather Education level of the father (levels same as EdMother).
- Age Age at the start of the course.
- Employment Employment level at college enrollment (1 = No; 2 = part-time; 3 = full-time).
- Income Household income level at college enrollment (1 = <25K; 2 = <35K; 3 = <45K; 4 = <55K; 5 = <70K; 6 = <85K; 7 = <100K; 8 = <120K; 9 = >120K).
- Transfer Number of transfer credits at the start of the course.
- GPA GPA as of the start of the course.
- GradeCode Letter grade.
- Level Level of the course, either Lower or Upper.
- ID Randomly assigned student ID.

Format

a data frame with 17 variables.

26 unmatched

unmatched

Returns rows from trips that were not matched by trimatch.

Description

This function returns a subset of trips that were not matched by trimatch. All data frame methods work with the returned object but special summary function will provided relevant information.

Usage

unmatched(tmatch)

Arguments

tmatch

the results of trimatch.

Value

a data frame of unmatched rows.

Index

* analysis	multibalance.plot, 11
TriMatch-package, 2	
* datasets	nmes, 12
nmes, 12	
tutoring, 25	OneToN, 12, 22
* matching	pairwise wilesy test 20
TriMatch-package, 2	pairwise.wilcox.test, 20
* propensity	parallel.plot, 13
TriMatch-package, 2	perpPt, 13
* psa	plot.balance.plots, 5, 14, 16
TriMatch-package, 2	plot.triangle.matches, 14
* score	plot.triangle.psa, 14, 15, 15, 17
TriMatch-package, 2	print.balance.plots, 16
	print.triangle.plot, 17
as.data.frame.list,3	print.trimatch.summary, 17
balance.plot, 4, 14, 16, 19	scale_colour_brewer,9
boxdiff.plot, 5	segment1, 18
boxairi.piot, 3	segment2, 18
covariateBalance, 6	star, 18
cv.bal.psa, 6, 11	stat_smooth, 9
CV. ba1. p3a, 0, 11	summary.balance.plots, 16, 19
data.frame, 3, 7	summary.triangle.matches, 17, 19
data.frame.to.list,7	summary.triangle.psa, 20
distance.euclid, 8	summary.unmatched, 21
distances.plot, 8	
410 tall 1000 F 100 t, 6	theme, 5
ezANOVA, 20	TriMatch (TriMatch-package), 2
,	trimatch, 4, 5, 8–11, 13, 20, 21, 26
friedman.test, 20	TriMatch-package, 2
	trimatch.apply2,23
<pre>geom_jitter, 9</pre>	trips, <i>11</i> , <i>16</i> , <i>20–23</i> , 23, 26
<pre>geom_point, 9</pre>	tutoring, 25
$geom_smooth, 9$	
glm, 24	unmatched, 21, 26
loess3.plot,9	
maximumTreat, 10, 22 merge.triangle.matches, 10 merge.triangle.psa, 11	