

Package ‘DiPs’

July 21, 2025

Type Package

Title Directional Penalties for Optimal Matching in Observational Studies

Version 0.6.4

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Description Improves the balance of optimal matching with near-fine balance by giving penalties on the unbalanced covariates with the unbalanced directions. Many directional penalties can also be viewed as Lagrange multipliers, pushing a matched sample in the direction of satisfying a linear constraint that would not be satisfied without penalization. Yu and Rosenbaum (2019) <[doi:10.1111/biom.13098](https://doi.org/10.1111/biom.13098)>.

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Additional_repositories <https://errickson.net/rrelaxiv/>

Encoding UTF-8

LazyData true

Imports stats, plyr, mvnfast, methods, rlemon

Suggests optmatch, rrelaxiv

Note One minimum cost flow problem may have several or many solutions that are equivalent in providing the same minimum total or mean cost. Minor differences between computers or implementations may have the minor consequence of altering which equivalent solution is produced. The optmatch package, which is useful for running many of the provided functions, may be downloaded from Github at <<https://github.com/markmfredrickson/optmatch>> if not available on CRAN. The rrelaxiv package, which provides an alternative solver for the underlying network flow problems, carries an academic license and is not available on CRAN, but may be downloaded from Github at <<https://github.com/josherrickson/rrelaxiv/>>.

NeedsCompilation no

Repository CRAN

Date/Publication 2022-08-07 14:30:09 UTC

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addcaliper	<i>Add a caliper, that need not be symmetric, to a distance object.</i>
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Description

Imposes a caliper, that need not be symmetric, on p using a penalty function, adding the penalty to a distance matrix $dmat$ and returning a new distance matrix. The symmetric version of this function is discussed in Rosenbaum (2010).

Usage

```
addcaliper(dist, z, dx, rg, stdev = FALSE, penalty = 1000)
```

Arguments

<code>dist</code>	A distance object with three components: <code>d</code> , <code>start</code> , <code>end</code> , typically created by <code>maha_dense</code> or <code>maha_sparse</code> . <code>d[i]</code> gives the distance between the $(start[i])$ th treated and the $(end[i]-sum(z))$ th control.
<code>z</code>	A vector whose i th coordinate is 1 for a treated unit and is 0 for a control. Must have treated subjects ($z = 1$) before controls ($z = 0$).
<code>dx</code>	A vector of with $length(z)=length(dx)$ giving the variable used to define the caliper. For instance, <code>dx</code> might be the propensity score.
<code>rg</code>	A vector with $length(rg) = 2$ such that $rg[1] \leq 0 \leq rg[2]$. If the treated-minus-control difference in <code>dx</code> is $< rg[1]$ or $> rg[2]$, then penalty is added to the distance. If treated individuals have <code>dx</code> higher than controls, then you want to set $rg[2] < -rg[1]$, so that you tolerate smaller positive differences and larger negative differences.
<code>stdev</code>	If <code>stdev = TRUE</code> , <code>rg</code> is interpreted in units of an equally weighted pooled standard deviation; that is, <code>rg</code> is replaced by <code>rg*sp</code> where $sp = \sqrt{(\text{var}(dx[z==1]) + \text{var}(dx[z==0]))/2}$.
<code>penalty</code>	The number added to a distance when the caliper is violated. A large penalty, like the default value of <code>penalty = 1000</code> , will try to enforce the caliper to the extent that this is feasible. Small penalties can slightly tilt a match in a desired direction.

Value

Returns a new distance object whose distance component d is updated by the sum of d and the penalties for caliper violations.

References

Yu, R., & Rosenbaum, P. R. (2019). Directional penalties for optimal matching in observational studies. *Biometrics*, 75(4), 1380-1390.

Rosenbaum, P. R. (2010) *Design of Observational Studies*. New York: Springer.

Examples

```
data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
      family = binomial)$fitted.values
d<-cbind(nh0506Homocysteine, p)
detach(nh0506Homocysteine)
dist0<-maha_dense(d$z, X)
#symmetric caliper
dist1<-addcaliper(dist0, d$z, d$p, c(-.3,.3), stdev = TRUE,
                 penalty = 1000)
head(dist1$d)
#asymmetric caliper
dist2<-addcaliper(dist0, d$z, d$p, c(-.5,.1), stdev = TRUE,
                 penalty = 1000)
head(dist2$d)
```

addDirectPenalty	<i>Add a directional penalty to a distance object</i>
------------------	---

Description

Add a directional penalty to a distance object.

Usage

```
addDirectPenalty(dist, z, dx, positive = TRUE, penalty = 1)
```

Arguments

dist	A distance object with three components: d , $start$, end , typically created by <code>maha_dense</code> or <code>maha_sparse</code> . $d[i]$ gives the distance between the $(start[i])$ th treated and the $(end[i]-sum(z))$ th control.
z	A vector whose i th coordinate is 1 for a treated unit and is 0 for a control. Must have treated subjects ($z = 1$) before controls ($z = 0$).

dx	A vector of with $\text{length}(z) = \text{length}(dx)$ giving the variable used to define the caliper. For instance, dx might be the propensity score.
positive	If <code>positive = TRUE</code> , a treated-minus-control difference in dx that is positive is increased by penalty, but if <code>positive = FALSE</code> a a treated-minus-control difference in dx that is negative is increased by penalty. Zero differences are never penalized.
penalty	The number added to a distance when the desired direction is violated.

Value

Returns a new distance matrix that is the sum of dmat and the penalties for direction violations.

References

Yu, R., & Rosenbaum, P. R. (2019). Directional penalties for optimal matching in observational studies. *Biometrics*, 75(4), 1380-1390.

Examples

```
data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
       family=binomial)$fitted.values
d<-cbind(nh0506Homocysteine, p)
detach(nh0506Homocysteine)
dist<-maha_dense(d$z, X)
head(dist$d)
dist<-addDirectPenalty(dist, d$z, d$p, positive=TRUE, penalty = 1)
head(dist$d)
```

addMagnitudePenalty *Add a directional magnitude penalty to a distance matrix*

Description

Adds a penalty to the distance component d of the distance object dist depending upon value of dx. The distance object dist has three components: d, start, end. $d[i]$ gives the distance between the tth treated and the cth control, with $t = \text{start}[i]$ and $c = \text{end}[i] - \text{sum}(z)$. The value of dx for treated unit t, say dxt, is $dx[z==1][t]$ and the value of dx for control c, say dxc, is $dx[z==0][c]$. Then, $d[i]$ is adjusted using $dxt - dxc$. If `positive=TRUE`, the adjustment changes $d[i]$ to $d[i] + \text{multiplier} * (\max(0, (dxt - dxc) - \text{hstick}))$. That is, a penalty is imposed if dxt exceeds dxc by more than hstick. If `positive = FALSE`, the adjustment changes $d[i]$ to $d[i] + \text{multiplier} * (\max(0, (dxc - dxt) - \text{hstick}))$.

Usage

```
addMagnitudePenalty(dist, z, dx, positive = TRUE, hstick = 0,
                    multiplier = 2)
```

Arguments

<code>dist</code>	A distance object with three components: <code>d</code> , <code>start</code> , <code>end</code> , typically created by <code>maha_dense</code> or <code>maha_sparse</code> . <code>d[i]</code> gives the distance between the <code>(start[i])</code> th treated and the <code>(end[i]-sum(z))</code> th control.
<code>z</code>	A vector whose <code>i</code> th coordinate is 1 for a treated unit and is 0 for a control. Must have treated subjects (<code>z=1</code>) before controls (<code>z=0</code>).
<code>dx</code>	A vector of with <code>length(z)=length(dx)</code> giving the variable used to define the caliper. For instance, <code>dx</code> might be the propensity score.
<code>positive</code>	If <code>positive = TRUE</code> , a treated-minus-control difference <code>di</code> in <code>dx</code> that is positive is increased by a multiple of <code>ldil</code> , but if <code>positive = FALSE</code> a a treated-minus-control difference in <code>dx</code> that is negative is increased by a multiple of <code>ldil</code> .
<code>hstick</code>	Hockey-stick value, which is a nonnegative number. See the description.
<code>multiplier</code>	The magnitude added is <code>multiplier*ldil/s</code> where <code>s</code> is an equally weighted, pooled within group standard deviation of <code>dx</code> .

Value

Returns a new distance object whose distance component `d` is updated by the sum of `dmat` and the penalties.

References

Yu, R., & Rosenbaum, P. R. (2019). Directional penalties for optimal matching in observational studies. *Biometrics*, 75(4), 1380-1390.

Examples

```
## Not run:
library(MASS)
data("nh0506Homocysteine")
attach(nh0506Homocysteine)
# Select covariates
X<-cbind(female, age, black, education, povertyr, bmi)
#Propensity score
p<-glm(z ~ female + age + black + education + povertyr + bmi,
      family=binomial)$fitted.values
d<-cbind(nh0506Homocysteine, p)
detach(nh0506Homocysteine)
dist<-maha_dense(d$z, X)
head(dist$d)
# Impose a penalty when a treated individual has a higher propensity
# score than a control
dist<-addMagnitudePenalty(dist, d$z, d$p, positive=TRUE, multiplier = 20)
head(dist$d)

## End(Not run)
```

check	<i>Check standardized mean differences (SMDs) of the matched data set.</i>
-------	--

Description

The function is used to create a table of mean and SMDs to check the balance before and after matching.

Usage

```
check(fdata, mdata, fz, mz)
```

Arguments

<code>fdata</code>	A full data frame with <code>length(fz)</code> rows and columns being variables that need to check SMDs. <code>fdata</code> and <code>mdata</code> must have the same variables with same column names in the same order.
<code>mdata</code>	A matched data frame with <code>length(mz)</code> rows and columns being variables that need to check SMDs. <code>fdata</code> and <code>mdata</code> must have the same variables with same column names in the same order.
<code>fz</code>	A vector whose <i>i</i> th coordinate is 1 for a treated unit and is 0 for a control for subjects in the full data set.
<code>mz</code>	A vector whose <i>i</i> th coordinate is 1 for a treated unit and is 0 for a control for subjects in the matched data set.

Value

A matrix with one row for each variable and five columns being the mean of treated group, mean of matched control group, mean of full control group, SMD of matched control group and SMD of full control group.

References

Rosenbaum, P. R. (2010) Design of Observational Studies. New York: Springer.

Examples

```
# To run this example, you must load the optmatch package.
# The optmatch is available on CRAN and Github.

data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
      family = binomial)$fitted.values
d<-cbind(nh0506Homocysteine, p)
detach(nh0506Homocysteine)
```

```

dist<-maha_dense(d$z, X)
o<-match(d$z, dist, d)
matcheddata<-o$data
Xm<-subset(matcheddata, select=c('female', 'age', 'black', 'education',
                                'poverty', 'bmi', 'p'))
check(cbind(X, p), Xm, d$z, matcheddata$z)

```

edgenum

Computes the number of edges in the reduced bipartite graph.

Description

Computes the number of edges in the reduced bipartite graph after applying the caliper and number of nearest neighbors (constant). Equivalently, this is the number of candidate pairs for matching in the observational study.

This function can provide users some idea of the required computation time. Smaller caliper and constant removes more edges, hence accelerates computation, but risks infeasibility.

Usage

```
edgenum(z, p, caliper, constant=NULL, exact=NULL, ties.all=TRUE)
```

Arguments

<code>z</code>	A vector whose <i>i</i> th coordinate is 1 for a treated unit and is 0 for a control.
<code>p</code>	A vector of length(<code>z</code>)=length(<code>p</code>) giving the variable used to define the caliper. Typically, <code>p</code> is the propensity score or its rank.
<code>caliper</code>	If two individuals differ on <code>p</code> by more than <code>caliper</code> , we will not calculate the distance for this pair.
<code>constant</code>	If the number of pairs within a caliper is greater than <code>constant</code> , we will select the constant closest ones.
<code>exact</code>	If not <code>NULL</code> , then a vector of length(<code>z</code>)=length(<code>p</code>) giving variable that need to be exactly matched.
<code>ties.all</code>	If <code>ties.all</code> is <code>True</code> , include all ties while choosing nearest neighbors. In this case, some treated may have more than <code>constant</code> controls. Otherwise, randomly select one or several controls to make sure there are not more than <code>constant</code> controls for each treated.

Details

A given choice of caliper and number of nearest neighbors (`constant`) removes candidate pairs, so there exists a corresponding reduced bipartite graph.

Smaller caliper and `constant` removes more edges from the original dense graph, hence the computation is faster. However, this risks infeasibility. A smallest caliper that permits a feasible match and its corresponding smallest number of nearest neighbors can be computed by functions `optcal()` and `optconstant()`.

Value

Number of edges in the reduced bipartite graph with the constraints on caliper and number of nearest neighbors (constant).

References

Yu, R., Silber, J. H., & Rosenbaum, P. R. (2020). Matching methods for observational studies derived from large administrative databases (with Discussion). *Statistical Science*, 35(3), 338-355.

Examples

```
data(nh0506Homocysteine)
attach(nh0506Homocysteine)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
       family = binomial)$fitted.values
edgenum(z, p, 0.2)
edgenum(z, p, 0.2, 10, exact=female)
detach(nh0506Homocysteine)
```

maha_dense	<i>Creates a robust Mahalanobis distance for matching based on a dense network.</i>
------------	---

Description

Computes a robust Mahalanobis distance list for use in dense matching. In this case, we compute the distance for all possible pairs of treated and control.

This function and its use are discussed in Rosenbaum (2010). The robust Mahalanobis distance in described in Chapter 8 of Rosenbaum (2010).

Usage

```
maha_dense(z, X, exact=NULL, nearexact=NULL, penalty=100)
```

Arguments

z	A vector whose ith coordinate is 1 for a treated unit and is 0 for a control.
X	A matrix with length(z) rows giving the covariates. X should be of full column rank.
exact	If not NULL, then a vector of length(z) = length(p) giving variable that need to be exactly matched.
nearexact	If not NULL, then a vector of length length(z) giving variable that need to be exactly matched.
penalty	The penalty for a mismatch on nearexact.

Details

The usual Mahalanobis distance works well for multivariate Normal covariates, but can exhibit odd behavior with typical covariates. Long tails or an outlier in a covariate can yield a large estimated variance, so the usual Mahalanobis distance pays little attention to large differences in this covariate. Rare binary covariates have a small variance, so a mismatch on a rare binary covariate is viewed by the usual Mahalanobis distance as extremely important. If you were matching for binary covariates indicating US state of residence, the usual Mahalanobis distance would regard a mismatch for Wyoming as much worse than a mismatch for California.

The robust Mahalanobis distance uses ranks of covariates rather than the covariates themselves, but the variances of the ranks are not adjusted for ties, so ties do not make a variable more important. Binary covariates are, of course, heavily tied.

Value

d	A distance object for each pair of treated and control.
start	The treated subject for each distance.
end	The control subject for each distance.

References

Rosenbaum, P. R. (2010) Design of Observational Studies. New York: Springer.

Examples

```
data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
dist<-maha_dense(z, X)
head(dist$d)
detach(nh0506Homocysteine)
```

maha_sparse	<i>Creates a robust Mahalanobis distance for matching based on a sparse network.</i>
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Description

Computes a robust Mahalanobis distance list for use in sparse matching. In this case, we will only calculate the distance for pairs within the caliper on p. If the caliper is too small, the matching may be infeasible. For the smallest caliper that keeps feasibility, refer to `optcal()` in package 'bigmatch'.

This function and its use are discussed in Rosenbaum (2010). It is preferred when the dataset is large. The robust Mahalanobis distance is described in Chapter 8 of Rosenbaum (2010).

Usage

```
maha_sparse(z, X, p=rep(1,length(z)), caliper=1, stdev=FALSE,
            constant=NULL, ncontrol=1, exact=NULL, nearexact=NULL,
            penalty=100, subX=NULL, ties.all=TRUE)
```

Arguments

<code>z</code>	A vector whose <i>i</i> th coordinate is 1 for a treated unit and is 0 for a control.
<code>X</code>	A matrix with <code>length(z)</code> rows giving the covariates. <code>X</code> should be of full column rank.
<code>p</code>	A vector of <code>length(z) = length(p)</code> giving the variable used to define the caliper. Typically, <code>p</code> is the propensity score or its rank.
<code>caliper</code>	If two individuals differ on <code>p</code> by more than <code>caliper</code> , we will not calculate the distance for this pair. If <code>caliper</code> is a positive number, then a symmetric caliper is applied. If <code>caliper</code> is a vector of a negative number and a positive number, then an asymmetric caliper is applied.
<code>stdev</code>	If <code>stdev = TRUE</code> , <code>caliper</code> is interpreted in units of an equally weighted pooled standard deviation; that is, <code>caliper</code> is replaced by <code>caliper*sp</code> where <code>sp = sqrt((var(dx[z==1])+var(dx[z==0])))</code>
<code>constant</code>	If the number of pairs within a caliper is greater than <code>constant</code> , we will select the constant closest ones.
<code>ncontrol</code>	A positive integer giving the number of controls to be matched to each treated subject. If <code>ncontrol</code> is too large, the match will be infeasible.
<code>exact</code>	If not <code>NULL</code> , then a vector of <code>length(z) = length(p)</code> giving variable that need to be exactly matched.
<code>nearexact</code>	If not <code>NULL</code> , then a vector of length <code>length(z)</code> giving variable that need to be exactly matched.
<code>penalty</code>	The penalty for a mismatch on <code>nearexact</code> .
<code>subX</code>	If a subset matching is required, the variable that the subset matching is based on. That is, for each level of <code>subX</code> , extra treated will be discarded in order to have the number of matched treated subjects being the minimum size of treated and control groups. If exact matching on a variable <code>x</code> is desired and discarding extra treated is fine if there are more treated than controls for a certain level <code>k</code> , set <code>exact = x</code> , <code>subX = x</code> .
<code>ties.all</code>	If <code>ties.all</code> is <code>True</code> , include all ties while choosing nearest neighbors. In this case, some treated may have more than constant controls. Otherwise, randomly select one or several controls to make sure there are not more than constant controls for each treated.

Details

The usual Mahalanobis distance works well for multivariate Normal covariates, but can exhibit odd behavior with typical covariates. Long tails or an outlier in a covariate can yield a large estimated variance, so the usual Mahalanobis distance pays little attention to large differences in this covariate. Rare binary covariates have a small variance, so a mismatch on a rare binary covariate is viewed by the usual Mahalanobis distance as extremely important. If you were matching for binary covariates indicating US state of residence, the usual Mahalanobis distance would regard a mismatch for Wyoming as much worse than a mismatch for California.

The robust Mahalanobis distance uses ranks of covariates rather than the covariates themselves, but the variances of the ranks are not adjusted for ties, so ties do not make a variable more important. Binary covariates are, of course, heavily tied.

Value

d	A distance list for each pair within the caliper distance and constant constraint.
start	The treated subject for each distance.
end	The control subject for each distance.

References

Yu, R., Silber, J. H., & Rosenbaum, P. R. (2020). Matching methods for observational studies derived from large administrative databases (with Discussion). *Statistical Science*, 35(3), 338-355.

Examples

```
data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
      family=binomial)$fitted.values
d<-cbind(nh0506Homocysteine,p)
detach(nh0506Homocysteine)

#apply symmetric caliper 0.15 on propensity score
dist1<-maha_sparse(d$z, X, p, 0.15)
length(dist1$d)

#apply asymmetric caliper c(-0.2,0.1) on propensity score
dist2<-maha_sparse(d$z, X, p, c(-0.2,0.1))
length(dist2$d)
```

match	<i>Minimum-distance near-fine matching.</i>
-------	---

Description

The program finds an optimal near-fine match with a given caliper on p, plus directional penalties on dx to offset the distribution imbalances. That is, it finds a near-fine match that minimizes the penalized Mahalanobis distance. In order to avoid the distortion of the original distribution by large penalties, it has the option of apply asymmetric calipers on those covariates.

Usage

```
match(z, dist, dat, p=rep(1,length(z)), exact=NULL,
      fine=rep(1,length(z)), ncontrol=1,
      penalty=round(max(dist$d)*1000), s.cost=100, subX=NULL)
```

Arguments

<code>z</code>	A vector whose i th coordinate is 1 for a treated unit and is 0 for a control.
<code>dist</code>	A distance object with three components: <code>d</code> , <code>start</code> , <code>end</code> , typically created by <code>maha_dense</code> or <code>maha_sparse</code> . <code>d[i]</code> gives the distance between the $(start[i])$ th treated and the $(end[i]-sum(z))$ th control.
<code>dat</code>	A data frame with <code>length(z)</code> rows. If the match is feasible, the matched portion of <code>dat</code> is returned with additional columns that define the match.
<code>p</code>	A vector of <code>length(z)=length(p)</code> giving the variable used to define the caliper. Typically, <code>p</code> is the propensity score or its rank. If the dense match is performed, use the default <code>p=rep(1,length(z))</code> .
<code>exact</code>	If not <code>NULL</code> , then a vector of <code>length(z) = length(p)</code> giving variable that need to be exactly matched.
<code>fine</code>	A vector of with <code>length(z) = length(fine)</code> giving the nominal levels that are to be nearly-finely balanced.
<code>ncontrol</code>	A positive integer giving the number of controls to be matched to each treated subject. If <code>ncontrol</code> is too large, the match will be infeasible.
<code>penalty</code>	A numeric penalty imposed for each violation of fine balance.
<code>s.cost</code>	The scaling factor for cost of the each pair of treated and control while rounding the cost.
<code>subX</code>	If a subset matching is required, the variable that the subset matching is based on. That is, for each level of <code>subX</code> , extra treated will be discarded in order to have the number of matched treated subjects being the minimum size of treated and control groups. If exact matching on a variable <code>x</code> is desired and discarding extra treated is fine if there are more treated than controls for a certain level <code>k</code> , set <code>exact = x</code> , <code>subX = x</code> .

Details

The match minimizes the total distance between treated subjects and their matched controls subject to a near-fine balance constraint imposed as a penalty on imbalances. Another set of directional penalties on `dx` can be imposed in order to offset the distribution imbalances. In order to avoid the case of matching far pairs to get close means, the user can the option of apply asymmetric calipers on covariates in `dx`. We add a larger penalty for pairs outside the asymmetric caliper to avoid infeasibility issue. But a match may be infeasible if the caliper on `p` is too small. In this case, increase the caliper, or find the smallest caliper that gives a feasible matching by using `optcal()` in package 'bigmatch'.

For discussion of networks for fine-balance, see Rosenbaum (1989, Section 3) and Rosenbaum (2010). For near-fine balance, see Yang et al. (2012).

You MUST install and load the `optmatch` package to use `match()`.

Value

If the match is infeasible, a warning is issued. Otherwise, a list of results is returned.

A match may be infeasible if the caliper or constant on `p` is too small, or `ncontrol` is too large, or if exact matching for `exact` is impossible.

feasible	Indicator of whether matching is feasible or not.
data	The matched sample, selected rows of dat.
x	A vector of indicators of whether each treated-control pair is included in the matched sample.

References

- Bertsekas, D. P. and Tseng, P. (1988) The relax codes for linear minimum cost network flow problems. *Annals of Operations Research*, 13, 125-190. Fortran and C code: <http://www.mit.edu/~dimitrib/home.html>. Available in R via the optmatch package.
- Rosenbaum, P.R. (1989) Optimal matching in observational studies. *Journal of the American Statistical Association*, 84, 1024-1032.
- Rosenbaum, P. R. (2010) *Design of Observational Studies*. New York: Springer.
- Yang, D., Small, D. S., Silber, J. H., and Rosenbaum, P. R. (2012) Optimal matching with minimal deviation from fine balance in a study of obesity and surgical outcomes. *Biometrics*, 68, 628-636.
- Yu, R., & Rosenbaum, P. R. (2019). Directional penalties for optimal matching in observational studies. *Biometrics*, 75(4), 1380-1390.

Examples

```
# To run this example, you must install and load the optmatch package.
# The optmatch is available on CRAN and Github.

data("nh0506Homocysteine")
attach(nh0506Homocysteine)
X<-cbind(female, age, black, education, povertyr, bmi)
p<-glm(z ~ female + age + black + education + povertyr + bmi,
       family=binomial)$fitted.values
d<-cbind(nh0506Homocysteine,p)
detach(nh0506Homocysteine)
dist<-maha_dense(d$z,X)
dist$d<-dist$d+1000*as.numeric(dist$d>7)
dist<-addcaliper(dist, d$z, d$p, c(-.5,.15), stdev=TRUE, penalty=1000)
dist<-addMagnitudePenalty(dist, d$z, d$p, positive=TRUE, multiplier=20)
dist<-addDirectPenalty(dist, d$z, d$p, positive=TRUE, penalty=1)
dist<-addDirectPenalty(dist, d$z, d$black, positive=FALSE, penalty=2)
dist<-addDirectPenalty(dist, d$z, d$bmi, positive=FALSE, penalty=2)
dist<-addDirectPenalty(dist, d$z, d$female, positive=FALSE, penalty=4)
o<-match(d$z, dist, d, fine=d$education, ncontrol=2)
md<-o$data
head(md)
```

net	<i>Optimal near-fine match from a distance matrix.</i>
-----	--

Description

The function creates the network for optimal near-fine matching to be passed via `callrelax` to the Fortran code for Bertsekas and Tseng's (1988) Relax IV.

Of limited interest to most users; function `netfine()` would typically be called by some other functions.

Usage

```
net(z, dist, ncontrol=1, fine=rep(1,length(z)),
    penalty=round(max(dist$d)*100), s.cost=100, subX=NULL)
```

Arguments

<code>z</code>	A vector whose <i>i</i> th coordinate is 1 for a treated unit and is 0 for a control.
<code>dist</code>	A distance list with the starting node (treated subject), ending node (control), the extra distance between them based on directional penalty.
<code>ncontrol</code>	A positive integer giving the number of controls to be matched to each treated subject.
<code>fine</code>	A vector of with <code>length(z) = length(fine)</code> giving the nominal levels that are to be nearly-finely balanced.
<code>penalty</code>	A numeric penalty imposed for each violation of fine balance.
<code>s.cost</code>	The scaling factor for cost of the each pair of treated and control while rounding the cost.
<code>subX</code>	If a subset matching is required, the variable that the subset matching is based on. That is, for each level of <code>subX</code> , extra treated will be discarded in order to have the number of matched treated subjects being the minimum size of treated and control groups. If exact matching on a variable <code>x</code> is desired and discarding extra treated is fine if there are more treated than controls for a certain level <code>k</code> , set <code>exact = x</code> , <code>subX = x</code> .

Details

The network contains a bipartite graph for treated and control subjects plus additional nodes for fine balance categories, plus additional nodes accept needed deviations from fine balance yielding near-fine balance.

For discussion of fine-balance, see Rosenbaum (1989, Section 3) and Rosenbaum (2010). For near-fine balance balance, see Yang et al. (2012).

Value

A network for optimal near-fine matching.

References

- Bertsekas, D. P. and Tseng, P. (1988) The relax codes for linear minimum cost network flow problems. *Annals of Operations Research*, 13, 125-190. Fortran and C code: <http://www.mit.edu/~dimitrib/home.html>. Available in R via the optmatch package.
- Rosenbaum, P.R. (1989) Optimal matching in observational studies. *Journal of the American Statistical Association*, 84, 1024-1032.
- Rosenbaum, P. R. (2010) *Design of Observational Studies*. New York: Springer.
- Yang, D., Small, D. S., Silber, J. H., and Rosenbaum, P. R. (2012) Optimal matching with minimal deviation from fine balance in a study of obesity and surgical outcomes. *Biometrics*, 68, 628-636.

nh0506Homocysteine	<i>Homocysteine and Smoking</i>
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Description

NHANES 2005-2006 data on smoking and homocysteine levels in adults.

Usage

```
data("nh0506Homocysteine")
```

Format

A data frame with 2759 observations on the following 11 variables.

x Row number, 1 to 2759

SEQN NHANES identification number

z Smoking status, 1 = daily smoker, 0 = never smoker

female 1 = female, 0 = male

age Age in years, ≥ 20 , capped at 85

black 1=black race, 0=other

education Level of education

povertyr Ratio of family income to the poverty level, capped at 5 times poverty

bmi BMI or body-mass-index

cigspersday30 Cigarettes smoked per day, 0 for never smokers

cotinine Blood cotinine level, a biomarker of recent exposure to tobacco

homocysteine Level of homocysteine

Details

The following code constructed the data as used here. Attention is confined to adults, excluding children. Also, people who have smoked in the past, but do not now smoke daily, are excluded. A moderate number of individuals with missing povertyr, cotinine or homocysteine were excluded.

```
library(foreign)
DEMO<-read.xport("DEMO_D.XPT")
HCY<-read.xport("HCY_D.XPT")
SMQ<-read.xport("SMQ_D.XPT")
BMX<-read.xport("BMX_D.XPT")
COT<-read.xport("COT_D.XPT")
d<-merge(DEMO, HCY, by="SEQN", all.x=TRUE)
d<-merge(d, SMQ, by="SEQN", all.x=TRUE)
d<-merge(d, COT, by="SEQN", all.x=TRUE)
d<-merge(d, BMX, by="SEQN", all.x=TRUE)
rm(DEMO, HCY, SMQ, COT, BMX)
SEQN<-d$SEQN
age<-d$RIDAGEYR
race<-d$RIDRETH1
black<-1*(race==4)
hispanic<-1*((race==1)|(race==2))
female<-1*(d$RIAGENDR==2)
education<-d$DMDEDUC2
education[education>6]<-NA
povertyr<-d$INDFMPIR
homocysteine<-d$LBXHCY
bmi<-d$BMXBMI
cotinine<-d$LBXCOT
cigs100life<-d$SMQ020
cigs100life[cigs100life>3]<-NA
cigs100life<-(cigs100life==1)*1
smokenow<-1*(d$SMQ040<2.5)
smokenow[cigs100life==0]<-0
cigsdays30<-d$SMD641
cigsdays30[cigsdays30>32]<-NA
cigsdays30[smokenow==0]<-0
cigsperday30<-d$SMD650
cigsperday30[cigsperday30>100]<-NA
```



```

cigsperday30[smokenow==0]<-0
dailysmoker<-1*((cigs100life==1)&(cigsdays30==30)&(smokenow==1)&(cigsperday30>=10))
neversmoker<-1*((cigs100life==0)&(smokenow==0))
z<-dailysmoker
z[(neversmoker==0)&(dailysmoker==0)]<-(-999)
ds<-data.frame(SEQN, female, age, black, education, povertyr, bmi, homocysteine, cotinine, cigs100life,
smokenow, cigsdays30, cigsperday30, dailysmoker, neversmoker, z)
use<-age>=20
ds1<-ds[use,]
use<-complete.cases(ds1)
ds1$z[ds1$z==-999]<-NA
ds2<-ds1[use&!is.na(ds1$z),]
rm(SEQN, female, age, black, hispanic, education, povertyr, homocysteine, cotinine, cigs100life,
smokenow, cigsdays30, cigsperday30, dailysmoker, neversmoker, z, race, use, bmi)
ds2<-ds2[order(1-ds2$z),]
attach(ds2)
nh0505Homocysteine<-data.frame(SEQN, z, female, age, black, education, povertyr, bmi, cigsper-
day30, cotinine, homocysteine)
write.csv(nh0506Homocysteine, "nh0506Homocysteine.csv")

```

Source

From the NHANES web page, for NHANES 2005-2006.

References

US National Health and Nutrition Examination Survey, 2005-2006. From the US Center for Health Statistics.

Examples

```

data(nh0506Homocysteine)
summary(nh0506Homocysteine)

```

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