# Package 'PropScrRand' 

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Type Package
Title Propensity Score Methods for Assigning Treatment in Randomized
Trials
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Description Contains functions to run propensity-biased allocation to balance covariate distribu-
tions in sequential trials and propensity-constrained randomization to balance covariate distribu-
tions in trials with known baseline covariates at time of randomization. Currently only sup-
ports trials comparing two groups.
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## $R$ topics documented:

PropScrRand-package ..... 2
genPerms ..... 2
getVar ..... 3
pba ..... 4
pcr ..... 5
piFunction ..... 7
plotpi ..... 8
Index ..... 9

## Description

Provides propensity score-based methods for allocating units to treatment experiments with two treatment levels (e.g., treatment and control).

## Details

| Package: | PropScrRand |
| :--- | :--- |
| Type: | Package |
| Version: | 1.1 |
| Date: | $2013-11-14$ |
| License: | GPL-3 |

For sequential allocation, the functions the user will interact with directly are pba and pbaAgain. Both of these functions perform propensity-biased allocation, producing a treatment assignment for the current unit, among other information. The function plotpi can be used to investigate the strength of balance forced by various values of the tuning parameter $k$, with curves for new values of $k$ added to the plot via addpi. For randomization when all baseline covariates are known, use pcr, which will conduct propensity-constrained randomization. The remaining functions are called from these internally.

## Author(s)

Travis M. Loux
Maintainer: Travis M. Loux [loux@slu.edu](mailto:loux@slu.edu)
genPerms Generate Treatment Permutations

## Description

Used within calls to pcr to generate a set of unique treatment permutations for randomization.

## Usage

genPerms(n, n1, nPerms)

## Arguments

n
Total number of units to be randomized.
$\mathrm{n} 1 \quad$ Number of units to receive treatment.
nPerms

Number of permutations to generate.

## Details

This function randomly samples $n P e r m s$ of the choose $(n, n 1)$ possible treatment permutations. If nPerms > choose ( $n, n 1$ ), then all choose ( $n, n 1$ ) permutations are generated systematically. Also, in the case of 1-to-1 allocation, the complement treatment vectors are also produced, so the returned matrix has $2 *$ nPerms permutations. Uniqueness is checked throughout and duplicate permutations disgarded.

## Value

The result is an $n 1 \times n$ Perms (or $n 1 \times$ choose $(n, n 1)$ or $n 1 \times 2 * n P e r m s)$ matrix. Each column represents one treatment permutation, with the values in the column giving the index of the treated units.

## Author(s)

Travis M. Loux

## Examples

genPerms ( $\mathrm{n}=50$, $\mathrm{n} 1=25$, $\mathrm{nPerms=500)}$
genPerms ( $n=50, n 1=35$, nPerms=500)

```
getVar Compute Propensity Score Variance
```


## Description

Given a data set and vector of indices for treated units, computes the variance of the propensity score fitted via logistic regression.

## Usage

getVar(covs, tIndex)

## Arguments

covs A data frame of baseline covariates.
tIndex A vector indicating which units are to receive treatment.

## Value

Returns the variance of the fitted propensity scores.

## Author(s)

Travis M. Loux

## Description

Performs propensity-biased allocation for assigning a new unit to treatment in a sequential design with two treatment levels (i.e., treatment and control).

## Usage

pba(x, tr, newx, $k=1$, global = 0.5)
pbaAgain(previous, newx, $k=N A$ )

## Arguments

x
tr A vector of treatment assignments (0 or 1) for previously assigned units.
newx Data frame of covariate values of the new unit.
$k \quad$ Balancing parameter.
global Global target proportion to be treated.
previous The output of a previous call to pba or pbaAgain

## Details

The function pba generates a treatment assignment for a new unit. The steps of the process include regressesing tr on $x$ by logistic regression, computeing the fitted value of the new unit using covarate values in newx, and transforming the fitted propensity score into the probability of treatment by a call to piFunction using k and global as parameters. The balancing parameter k must be one of 0 , Inf, or the ratio of two positive odd integers. Small values of $k$ result in less restrictive randomization while larger values of $k$ result in more forced balance. In particular, $k=0$ is equivalent to pure randomization and $k=I n f$ results in deterministic allocation. Finally, a treatment assignment for the new unit is generated via a Bernoulli trial with probability from piFunction.
The function pbaAgain takes as input the output from a previous call to pba or pbaAgain and runs pba for the new unit using the values of newx. If $k=$ NA (the default), the value of $k$ from previous is used; otherwise, the provided value of $k$ is used. The parameter global is assumed to stay the same throughout the trial. The output of pbaAgain contains the same information as pba.

## Value

results $\quad$ A list of results from the PBA procedure.
phat The fitted propensity score for the new unit.
ptreat The probability of assignment to the treatment group for the new unit.
newtr $\quad$ Result of random assignment using ptreat.
input A list of inputs to PBA procdure. Used in future calls to pbaAgain.

| $x$ | Input $x$. |
| :--- | :--- |
| $t r$ | Input tr. |
| newx | Input newx. |
| $k$ | Input $k$. |
| global | Input global. |

## Author(s)

Travis Loux

## References

Loux, T.M. (2013) A simple, flexible, and effective covariate-adaptive treatment allocation procedure. Statistics in Medicine 32(22), 3775-3787. DOI: 10.1002/sim. 5837

## Examples

```
    x0 = data.frame(matrix(rnorm(60), ncol=3))
    t0 = rbinom(nrow(x0), size=1, prob=0.5)
    x1 = data.frame(matrix(rnorm(3), ncol=3))
    trial1 = pba(x=x0, tr=t0, newx=x1, k=Inf)
    x2 = data.frame(matrix(rnorm(3), ncol=3))
    trial2 = pbaAgain(previous=trial1, newx=x2)
    x3 = data.frame(matrix(rnorm(3), ncol=3))
    trial3 = pbaAgain(previous=trial2, newx=x3, k=5/3)
```

    pcr
    
## Description

Performs propensity-contstrained randomization on a given data set with measured covariates on all units.

## Usage

pcr (x, nTreat, M, m)

## Arguments

x
nTreat
M
m

Data frame of covariates.
Number of units to be treated.
Number of candidate permutations to create.
Number of permutations to keep.

## Details

Given the parameters, pcr generates $M$ unique permutations by calling genPerms. For each permutation, the empirical propensity scores are computed and the variance returned by getVar. Finally, the $m$ permutations with the smallest propensity score variance are found. The $m$ permutations returned in best can then be used to perform randomization and randomization inference. We suggest $M>=10000$ and $\mathrm{m} / \mathrm{M}<=0.10$.

## Value

treatments
variance
cutoff
best

The (approximately) M permutations generated by genPerms.
A vector of the propensity score variances for all $M$ permutations in treatments. The calculated $\mathrm{m} / \mathrm{M}$ quantile of propensity score variances.
The column indices of the permutations in treatments with propensity score variance below cutoff.

## Author(s)

Travis Loux

## References

Loux, T.M. (2015) Randomization, matching, and propensity scores in the design and analysis of experimental studies with known covariates. Statistics in Medicine. 34(4), 558-570. DOI: 10.1002/sim. 6361

## Examples

```
# 1:1 allocation, M small
dat1 = data.frame(x1=rnorm(50),
    x2=rnorm(50),
    x3=sample(c('a','b','c'), size=50, replace=TRUE))
trial1 = pcr(x=dat1, nTreat=25, M=500, m=50)
# 1:1 allocation, M large
dat2 = data.frame(x1=rnorm(10),
    x2=rnorm(10),
    x3=sample(c('a','b','c'), size=10, replace=TRUE))
trial2 = pcr(x=dat2, nTreat=5, M=200, m=20)
# non-1:1 allocation, M small
dat3 = data.frame(x1=rnorm(50),
    x2=rnorm(50),
    x3=sample(c('a','b','c'), size=50, replace=TRUE))
trial3 = pcr(x=dat3, nTreat=35, M=200, m=20)
# non-1:1 allocation, M large
dat4 = data.frame(x1=rnorm(10),
    x2=rnorm(10),
    x3=sample(c('a','b','c'), size=10, replace=TRUE))
trial4 = pcr(x=dat4, nTreat=6, M=300, m=30)
```

piFunction Get PBA Treatment Probability

## Description

Used within calls to pba and pbaAgain to obtain the probability a unit is assigned treatment given its fitted propensity score.

## Usage

piFunction(fit, kparam, qparam)

## Arguments

fit Fitted propensity score.
kparam Balancing parameter.
qparam Global target for proportion of units treated.

## Details

The input kparam must be one of 0 , Inf, or the ratio of two positive odd integers. Both fit and qparam must be between 0 and 1 .

## Value

A numeric object. In the conext of PBA, the probability of assignment to treatment for the current unit.

## Author(s)

Travis M. Loux

## Examples

```
piFunction(fit=0.6, kparam=1, qparam=0.5)
piFunction(fit=0.6, kparam=5, qparam=0.5)
piFunction(fit=0.6, kparam=1/5, qparam=0.5)
piFunction(fit=0.6, kparam=1, qparam=2/3)
piFunction(fit=0.6, kparam=5, qparam=2/3)
piFunction(fit=0.6, kparam=1/5, qparam=2/3)
```

```
    plotpi Plots of piFunction
```


## Description

Can be used to investigate the strength of balance forced by various values of the tuning parameter k.

## Usage

plotpi(k, global = 0.5)
addpi(k, global = 0.5, ...)

## Arguments

| k | Balancing parameter. |
| :--- | :--- |
| global | Global target for proportion of units treated. |
| $\ldots$. | Parameters in addpi passed to lines function. |

## Details

The function plotpi creates a plot of treatment probability against fitted propensity score for values of $k$ and global. The function addpi adds a curve for a new combination of $k$ and global to an existing plot.

## Author(s)

Travis M. Loux

## Examples

```
plotpi(k=3, global=0.5)
addpi(k=5/3, lty=2, col='red')
plotpi(k=1/3, global=2/3)
```


## Index

```
* design
    genPerms, 2
    pba,4
    pcr, 5
    PropScrRand-package, 2
* multivariate
    pba,4
    pcr, 5
    PropScrRand-package, 2
addpi (plotpi), 8
genPerms, 2, }
getVar, 3,6
pba,4
pbaAgain (pba),4
pcr, 5
piFunction, 4,7
plotpi,8
PropScrRand (PropScrRand-package), 2
PropScrRand-package, 2
```

