Package ‘rbounds’

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Title Perform Rosenbaum bounds sensitivity tests for matched and unmatched data.

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Depends R (>= 2.8.1), Matching

Description Takes matched and unmatched data and calculates Rosenbaum bounds for the treatment effect. Calculates bounds for binary outcome data, Hodges-Lehmann point estimates, Wilcoxon signed-rank test for matched data and matched IV estimators, Wilcoxon sum rank test, and for data with multiple matched controls. Package is also designed to work with the Matching package and operate on Match() objects.

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AngristLavy

**Description**


**Usage**

AngristLavy

**Format**

A *data.frame* with 318 observations on the following 5 variables.
- enrollment: Enrolled number of students in each cohort
- pct_disadv: Percentage of students that are economically disadvantaged.
- classsize: Size of class or classes for each cohort.
- avgmath: Average math test score for each class.
- z: A recode of enrollment with 1 indicating a cohort with 41 or more students, i.e. two classes. This serves as the instrument which encourages smaller classes.

**Source**


**References**


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binarysens

**Description**

Function to calculate Rosenbaum bounds for binary data.

**Usage**

`binarysens(x, y, Gamma=6, GammaInc=1)`
Arguments

x  Count of the first set of discrepant pairs in a table of treated and control outcomes.

y  Count of the second set of discrepant pairs in a table of treated and control outcomes.

Gamma  Upper-bound on gamma parameter.

GammaInc  To set user specified increments for gamma parameter.

Author(s)

Luke Keele, Penn State University, <ljk20@psu.edu>

References


See Also

See also data.prep, psens, hlsens, Match, mcontrol

Examples

# Example From Rosenbaum Observational Studies Pg 112
# Success: Died From Lung Cancer
# 110 Discrepant Pairs
# 12 Discrepant Pairs

# Sensitivity Test
binarysens(12,110)

# Example Using Match()

# Load Matching Software and Data
library(Matching)
data(GerberGreenImai)

# Estimate Propensity Score
pscore.glm <- glm(PHN.C1 ~ PERSONS + VOTE96.1 + NEW + MAJORPTY + AGE + WARD + PERSONS:VOTE96.1 + PERSONS:NEW + AGE2, family = binomial(logit), data = GerberGreenImai)

# Save data objects
D <- GerberGreenImai$PHN.C1
Y <- GerberGreenImai$VOTED98
X <- fitted(pscore.glm)

# Match - without replacement
m.obj <- Match(Y = Y, Tr = D, X = X, M = 1, replace=FALSE)
summary(m.obj)

# One should check balance, but let's skip that step for now.

# Sensitivity Test
binarysens(m.obj, Gamma=2, GammaInc=.1)

data.prep

**Format Match() object for sensitivity test.**

**Description**

This function reshapes the output from Match() to create the necessary objects for mcontrol().

**Usage**

# Default Method
data.prep(obj, Y = NULL, group.size = 3)

**Arguments**

- **obj**
  An object from the Match() function.

- **Y**
  Internal argument to the Match object.

- **group.size**
  The size of the matched groups. Three for one treated unit and two control units.

**Details**

This functions takes a Match() object and formats it for use with the mcontrol() function. The output is a list with the three objects needed for the arguments of the mcontrol() function.

**Value**

- **Y**
  The matched outcomes

- **id**
  A vector which identifies the matched groups: 1, 1, 1 for matched group one; 2, 2, 2 for match group 2, etc.

- **treat**
  A vector with 1’s for treated units and 0’s for control units

**Author(s)**

Luke Keele, Penn State University, <ljk20@psu.edu>

**References**

FisherSens

See Also

See also binarysens, psens, hlsens, Match, mcontrol

Examples

```r
# Load Matching Software and Data
library(Matching)
data(lalonde)

# Estimate Propensity Score
dWglm <- glm(treat~age + I(age^2) + educ + I(educ^2) + black + hisp +
married + nodegr + re74 + I(re74^2) + re75 + I(re75^2) +
u74 + u75, family=binomial, data=lalonde)

# Save data objects
Y <- lalonde$re78  # the outcome of interest
Tr <- lalonde$treat  # the treatment of interest

# Match
mDW <- Match(Y = Y, Tr = Tr, X = dWglm$fitted, M = 2)

# One should check balance, but let's skip that step for now.

# Create Data Object
tmp <- data.prep(mDW, group.size=3)

# Sensitivity Test
mcontrol(tmp$Y, tmp$id, tmp$treat, group.size = 3)
```

FisherSens

**Rosenbaum Sensitivity Analysis for Fisher’s Exact Test**

Description

Calculates sensitivity to hidden bias for Fisher’s exact test for a two-by-two contingency table, following the method described in Rosenbaum (2002, sec. 4.4).

Usage

FisherSens(totalN, treatedN, totalSuccesses, treatedSuccesses, Gammas)

Arguments

- `totalN`: total number of observations
- `treatedN`: number of treated observations
- `totalSuccesses`: total number of “successes”
### hlsens

**Rosenbaum Bounds for Hodges-Lehmann Point Estimate**

**Description**

Function to calculate Rosenbaum bounds for continuous or ordinal outcomes based on Hodges-Lehmann point estimate.

**Usage**

```r
# Default Method
hlsens(x, y = NULL, pr = 0.1, Gamma = 6, GammaInc = 1)
```

- **treatedSuccesses**
  - number of successes in treatment group
- **Gammas**
  - vector of Gammas (bounds on the differential odds of treatment) at which to test the significance of the results

**Value**

Returns a matrix with three columns and number of rows equal to the length of "Gammas". Each row indicates the upper and lower bounds for the (one-sided) p-value for a given value of Gamma.

**Author(s)**

Devin Caughey, MIT, <caughey@mit.edu>

**References**


**See Also**

See also `data.prep`, `binarysens`, `hlsens`, `Match`, `mcontrol`

**Examples**

```r
## Fisher's Lady Tasting Tea: milk first or tea first?
LadyTastingTea <- matrix(c(4, 0, 0, 4), nrow = 2,
                        dimnames = list(Guess = c("Milk", "Tea"),
                                        Truth = c("Milk", "Tea")))
## Define "Milk" as "treated"/"success"
FisherSens(totalN = sum(LadyTastingTea),
           treatedN = sum(LadyTastingTea["Milk", ]),
           totalSuccesses = sum(LadyTastingTea["Milk"], "Milk"],
           treatedSuccesses = sum(LadyTastingTea["Milk", "Milk"],
                                   Gammas = seq(1, 2, .2))
## Interpretation: Rejection of the null hypothesis
## (that the lady cannot discriminate between milk-first and tea-first)
## is insensitive to bias as large as Gamma = 2.
```
**Arguments**

- **x**: Treatment group outcomes in same order as treatment group outcomes or an objects from Match().
- **y**: Control group outcomes in same order as treatment group outcomes unnecessary when using Match() object.
- **pr**: Search precision parameter.
- **Gamma**: Upper-bound on gamma parameter.
- **GammaInc**: To set user specified increments for gamma parameter.

**Details**

For large data sets this function can be quite slow if pr is set to low. If the data set is larger, it is best to set pr to .5 before trying values such as .01. Generally, the results from the function are insensitive to the value for pr.

**Author(s)**

Luke Keele, Penn State University, <ljk20@psu.edu>

**References**


**See Also**

See also `data.prep`, `binarysens`, `psens`, `Match`, `mcontrol`

**Examples**

```r
# Replication of Rosenbaum Sensitivity Tests From Chapter 4 of
# Observational Studies

# Data: Matched Data of Lead Blood Levels in Children
trt <- c(38, 23, 41, 18, 37, 36, 23, 62, 31, 34, 24, 14, 21, 17, 16, 20, 15, 10, 45, 39, 22, 35, 49, 48, 44, 35, 43, 39, 34, 13, 73, 25, 27)
ctrl <- c(16, 18, 18, 24, 19, 11, 10, 15, 16, 18, 18, 13, 19, 10, 16, 16, 24, 13, 9, 14, 21, 19, 7, 18, 19, 12, 11, 22, 25, 16, 13, 11, 13)
hlsens(trt, ctrl)

# Load Matching Software and Data
library(Matching)
data(lalonde)

# Estimate Propensity Score
DWglm <- glm(treat~age + I(age^2) + educ + I(educ^2) + black + hisp +
    married + nodegr + re74 + I(re74^2) + re75 + I(re75^2) +
    u74 + u75, family=binomial, data=lalonde)
```
# Save data objects
Y <- lalonde$re78  # the outcome of interest
Tr <- lalonde$treat  # the treatment of interest

# Match - without replacement
mDW <- Match(Y=Y, Tr=Tr, X=DWglm$fitted, replace=FALSE)

# One should check balance, but let's skip that step for now.
# Sensitivity Test:
hlsens(mDW, pr=.1, Gamma=1.5, GammaInc=.25)

---

**iv_sens**  
*Function to calculate Rosenbaum bounds for IV Estimator based on Wilcoxon sign rank test.*

**Description**

`iv_sens` performs a non-parametric, instrumental variable sensitivity analysis on matched pairs following the logic of the Neyman-Rubin framework for causal inference. The function supports a variable-valued instrument.

**Usage**

`iv_sens(Rt, Rc, Dt, Dc, Gamma = 6, GammaInc = 1)`

**Arguments**

- **Rt, Rc**  
  Vectors of observed response outcomes for matched treatment and control observations, respectively.

- **Dt, Dc**  
  Vectors of observed doses for matched observations, respectively. This is level of dose encouraged by the instrument.

- **Gamma**  
  Upper-bound on gamma parameter.

- **GammaInc**  
  To set user specified increments for gamma parameter.

**Details**

Given matched pairs of observations on an instrument Z, which encourages dose D, this function performs a Rosenbaum’s bounds sensitivity analysis. Note that matching is done on levels of the instrument. See example below.

**Value**

Returns an object of class `rbounds`. 
mcontrol

Author(s)

Luke Keele, Penn State University, <ljk20@psu.edu>
Jason W. Morgan, Ohio State University, <morgan.746@osu.edu>

References


See Also

See also data.prep, binarysens, hlsens, Match, mcontrol

Examples

```r
## Example from Rosenbaum (2010, ch. 5).

data(AngristLavy)

#Match on Economic Status Across Levels of the Instrument
rr <- Match(Y=AngristLavy$avgmath, Tr=AngristLavy$z, X=AngristLavy$pct_disadv, estimand ="ATC", M=2, replace=FALSE)

#Extract Matched Outcome Data
ctrl <- AngristLavy$avgmath[rr$index.control]
trt <- AngristLavy$avgmath[rr$index.treated]

#Extract Matched Doses
#Doses Encouraged By Instrument - Here Class Size
csize.trt <- AngristLavy$classize[rr$index.treated]
csize.ctrl <- AngristLavy$classize[rr$index.control]

#Run Sensitivity Analysys
iv_sens(trt, ctrl, csizetrt, csizectl, Gamma=2, GammaInc=.1)
```

Description

Function to calculate Rosenbaum bounds for continuous or ordinal outcomes based on Wilcoxon sign rank test p-value when there are multiple matched control units.
Usage

# Default Method
mcontrol(y, grp.id, treat.id, group.size = 3, Gamma = 4, GammaInc = 1)

Arguments

y Vector of grouped matched outcomes.
treat.id A vector indicating the treated unit in each matched group.
grp.id A vector indicating matched groups.
group.size The size of the matched groups. Three for one treated unit and two control units.
Gamma Upper-bound on gamma parameter.
GammaInc To set increments for gamma parameter.

Details

The matched data needs to be in a very particular form for this function to work. The data must be
sorted by matched groups with indicators for each matched group and for treated and control units.
The simplest way to prepare the data is to use the Match() function and use the data.prep() function
to format the data.
Currently this function only takes matched data with 2 or 3 controls units matched to each treated
unit.
This function does cannot handle data where the number of control units is not the same for every
controlled unit.

Author(s)

Luke Keele, Penn State University, <ljk20@psu.edu>

References


See Also

See also data.prep, binarysens, psens, Match, hlsens

Examples

# Load Matching Software and Data
library(Matching)
data(lalonde)

# Estimate Propensity Score
dWglm <- glm(treat~age + I(age^2) + educ + I(educ^2) + black + hisp +
mixed + nodegr + re74 + I(re74^2) + re75 + I(re75^2) +
u74 + u75, family=binomial, data=lalonde)
# Save data objects
Y <- lalonde$re78  # the outcome of interest
Tr <- lalonde$treat  # the treatment of interest

# Match - without replacement
mDW <- Match(Y=Y, Tr=Tr, X=DWglm$fitted, M=2, replace=FALSE)

# One should check balance, but let's skip that step for now.

# Create Data Object:
tmp <- data.prep(mDW, group.size=3)

# Sensitivity Test
mcontrol(tmp$Y, tmp$id, tmp$treat, group.size=3)

---

**print.rbounds**

**Summarize output from rbounds object**

---

**Description**

Summary method for rbounds object.

**Usage**

```r
## S3 method for class 'rbounds'
print(x, ...)
```

**Arguments**

- `x` An object of class rbounds as produced by `binarysens`, `hlsens`, or `psens`.
- `...` Any additional arguments.

**Author(s)**

Jason W. Morgan, Ohio State University, <morgan.746@osu.edu>

**See Also**

Also see `binarysens`, `psens`, and `hlsens`.
psens

Rosenbaum Bounds for Sign Rank

Description
Function to calculate Rosenbaum bounds for continuous or ordinal outcomes based on Wilcoxon
sign rank test.

Usage
# Default Method
psens(x, y = NULL, Gamma = 6, GammaInc = 1)

Arguments
x Treatment group outcomes in same order as treatment group outcomes or an
objects from a Match.
y Control group outcomes in same order as treatment group outcomes unnecessary
when using a Match object.
Gamma Upper-bound on gamma parameter.
GammaInc To set user-specified increments for gamma parameter.

Author(s)
Luke Keele, Penn State University, <ljk20@psu.edu>

References

See Also
See also data.prep, binarysens, hlsens, Match, mcontrol

Examples

# Replication of Rosenbaum Sensitivity Tests From Chapter 4 of
# Observational Studies

# Data: Matched Data of Lead Blood Levels in Children
trt <- c(38, 23, 41, 18, 37, 36, 23, 62, 31, 34, 24, 14, 21, 17, 16, 20,
15, 10, 45, 39, 22, 35, 49, 48, 44, 35, 43, 39, 34, 13, 73, 25,
27)
ctrl <- c(16, 18, 18, 24, 19, 11, 10, 15, 16, 18, 18, 13, 19, 10, 16,
16, 24, 13, 9, 14, 21, 19, 7, 18, 19, 12, 11, 22, 25, 16, 13,
11, 13)
```r
psens(trt, ctrl)

# Example With Match()

# Load Matching Software and Data
library(Matching)
data(lalonde)

# Estimate Propensity Score
DWglm <- glm(treat ~ age + I(age^2) + educ + I(educ^2) + black + hisp + 
            married + nodegr + re74 + I(re74^2) + re75 + I(re75^2) + 
            u74 + u75, family=binomial, data=lalonde)

# Save data objects
Y <- lalonde$re78 #the outcome of interest
Tr <- lalonde$treat #the treatment of interest

# Match - without replacement
mDW <- Match(Y=Y, Tr=Tr, X=DWglm$fitted, replace=FALSE)

# One should check balance, but let's skip that step for now.

# Sensitivity Test
psens(mDW, Gamma = 2, GammaInc = 0.1)
```

---

### SumTestSens

**Rosenbaum Sensitivity Analysis for Unmatched Groups**

**Description**

Calculates sensitivity to hidden bias for tests based on sum statistics (e.g., Wilcoxon’s rank sum test), following the method described by Rosenbaum (2002, sec. 4.6). It is meant for unmatched/unstratified data with ordinal or continuous responses.

**Usage**

```
SumTestSens(T, q, n, m, Gamma)
```

**Arguments**

- **T**: observed value of the test statistic (e.g., the sum of the ranks of the responses of the treated units; note that a higher rank corresponds to a higher response)
- **q**: vector of functions of the responses (e.g., their ranks), sorted in decreasing order
- **n**: total number of observations
- **m**: number treated units
- **Gamma**: scalar indicating upper limit on the ratio of the a priori odds of treatment assignment between the treated and control groups
Value

This function prints the upper bound of the normal approximation one-sided p-value for the test at
the given value of Gamma. It also invisibly returns a list of intermediate statistics.

Warning

Since ‘SumTestSens’ calculates through enumeration the exact expectation and variance of the test
under the null, it is very computationally intensive and may be unworkable for even medium-sized
datasets.

Author(s)

Devin Caughey, MIT, <caughey@mit.edu>

References


See Also

See also data.prep, binarysens, hlsens, Match, mcontrol

Examples

```r
## Example from Rosenbaum (2002, p. 146)
mercury <- data.frame(matrix(c(1, 0, 2.7, 5.3,
                               2, 0, 0.5, 15.0,
                               3, 0, 0.0, 11.0,
                               4, 0, 0.0, 5.8,
                               5, 0, 5.0, 17.0,
                               6, 0, 0.0, 7.0,
                               7, 0, 0.0, 8.5,
                               8, 0, 1.3, 9.4,
                               9, 0, 0.0, 7.8,
                               10, 0, 1.8, 12.0,
                               11, 0, 0.0, 8.7,
                               12, 0, 0.0, 4.0,
                               13, 0, 1.0, 3.0,
                               14, 0, 1.8, 12.2,
                               15, 0, 0.0, 6.1,
                               16, 0, 3.1, 10.2,
                               17, 1, 0.7, 100.0,
                               18, 1, 4.6, 70.0,
                               19, 1, 0.0, 196.0,
                               20, 1, 1.7, 69.0,
                               21, 1, 5.2, 370.0,
                               22, 1, 0.0, 270.0,
                               23, 1, 5.0, 150.0,
                               24, 1, 9.5, 60.0,
                               25, 1, 2.0, 330.0,
                               26, 1, 3.0, 1100.0),
```

27, 1, 1.0, 40.0,
28, 1, 3.5, 100.0,
29, 1, 2.0, 70.0,
30, 1, 5.0, 150.0,
31, 1, 5.5, 200.0,
32, 1, 2.0, 304.0,
33, 1, 3.0, 236.0,
34, 1, 4.0, 178.0,
35, 1, 0.0, 41.0,
36, 1, 2.0, 120.0,
37, 1, 2.2, 330.0,
38, 1, 0.0, 62.0,
39, 1, 2.0, 12.8),
ncol = 4, byrow = TRUE))
colnames(mercury) <- c("ID", "Tr", "Pct.cu.cells", "Hg.in.blood")

T_test <- rank(mercury$Hg.in.blood) %>% mercury$Tr)
q_test <- sort(rank(mercury$Hg.in.blood), decreasing = TRUE))
n_test <- nrow(mercury))
m_test <- sum(mercury$Tr))

## Note: since this function uses exact rather than approximate
## formulas for the mean and variance of T, the p-values it
## calculates do not precisely match those in Rosenbaum (2002).

# A single Gamma value - example not run
# testOut2 <- SumTestSens(T = T_test,
# q = q_test,
# n = n_test,
# m = m_test,
# Gamma = 35)

## Apply to vector of Gamma values
sapply(c(1, 5, 35), SumTestSens,
T = T_test, q = q_test, n = n_test, m = m_test)
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