# Package 'MF'

December 21, 2018

Type Package
Title Mitigated Fraction
Version 4.3.5
Date 21 December 2018
Author David Siev
Maintainer Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov></marie.c.vendettuoli@aphis.usda.gov>
<b>Description</b> Calculate MF (mitigated fraction) with clustering and bootstrap options. See http://goo.gl/pcXYVr for definition of MF. No endorsement, claim, or warranty is implied for this package. It is made available for investigational or pedagogical use only
License MIT + file LICENSE
LazyLoad yes
LazyData yes
<b>Depends</b> R (>= 3.4.4)
<b>Imports</b> methods, dplyr (>= 0.7.1), plyr, stringr, tidyr, forcats, purrr
Collate 'aaa.r' 'classes.r' 'generic_methods.r' 'MF-package.r'  'HLBoot.r' 'MFBoot.r' 'MFClus.r' 'MFClusBoot.r' 'MFClusHier.R'  'MFmp.r' 'MFnestBoot.r' 'MFr.r' 'MFSubj.r' 'MFHier-wrappers.r'
RoxygenNote 6.0.1
NeedsCompilation no
R topics documented:
•
MF-package       2         calflung       3         HLBoot       3         mf-class       5         MFBoot       6         mfboot-class       7         mfbootcluster-class       8         MFClus       9         MFClusBoot       10         MFClusBootHier       12         mfclusboothier-class       13         MFClusHier       14
2.22 - 2.20 - 2.2

2 MF-package

MF-p	ackage MF Package	
Index		31
	piglung	
	MFSubj	
	MFr	
	MFnestBoot	26
	MFnest	
	mfmp-class	
	mfhlboot-class	
	mfhierdata-class	
	MFhBoot	18
	MFh	
	mfcluster-class	
	mfclushier-class	

# Description

Includes functions related to mitigated fraction.

For internal use only at the USDA Center for Veterinary Biologics.

### **Details**

Package: MF-package Type: Package Version: 4.3.5 Date: 21-Dec-2018

MIT License: LazyLoad: yes

# Author(s)

David Siev <David.Siev@aphis.usda.gov>

# **Examples**

```
# Checking MF package
example(MFr)
#-----
# End examples
```

calflung 3

```
#-----invisible()
```

calflung calflung dataset

# Description

Post-mortem examination of the lungs of groups of calves.

### **Format**

```
a data frame with 50 observations of the following 2 variables, no NAs
```

**group** Treatment group. One of con = control or vac = vaccinate

lesion Fraction of lungs with gross lesions.

HLBoot

Bootstrap CI for MF, HL, and Qdif

### **Description**

Estimates bootstrap confidence intervals for MF, HL, and Qdif.

# Usage

```
HLBoot(formula, data, compare = c("con", "vac"), b = 100, B = 100,
alpha = 0.05, hpd = TRUE, bca = FALSE, return.boot = FALSE,
trace.it = FALSE, seed = sample(1:1e+05, 1))
```

# Arguments

formula	Formula of the form $y \sim x + cluster(w)$ , where y is a continuous response, x is a factor with two levels of treatment, and w is a factor indicating the clusters.
data	Data frame
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared
b	Number of bootstrap samples to take with each cycle
В	Number of cycles, giving the total number of samples = $B * b$
alpha	Complement of the confidence level
hpd	Boolean whether to estimate highest density intervals for MF and HL.
bca	Boolean whether to estimate BCa intervals for MF.
return.boot	Boolean whether to save the bootstrap samples of the statistics.
trace.it	Boolean whether to display verbose tracking of the cycles.
seed	to initialize random number generator for reproducibility. Passed to set.seed.

4 HLBoot

#### **Details**

Estimates bootstrap confidence intervals for the mitigated fraction (MF), Hodge-Lehmann estimator (HL), and the difference of medians and quartiles (Qdif). Equal tailed intervals are provided for all three, highest density intervals are optionally provided for MF and HL, and BCa intervals are optionally provided for MF. The Hodges-Lehmann estimator is the median difference; it assumes that the two distributions have the same shape and differ by a constant shift. Assumes data is single pool (no nesting).

#### Value

```
a mfhlboot-class data object
```

### Author(s)

David Siev <david.siev@aphis.usda.gov>

#### References

Hodges JL, Lehmann EL, (1963). Estimates of location based on rank tests. *Annals of Mathematical Statistics*. **34:598–611**.

Siev D, (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508**.

Efron B, Tibshirani RJ. An Introduction to the Bootstrap. Chapman and Hall, New York, 1993.

### See Also

```
mfhlboot-class
```

### Examples

```
HLBoot(lesion~group, calflung, seed = 12345)
# Bootstrapping
# 10000 bootstrap samples
# 95% confidence intervals
# Comparing vac to con
# Mitigated Fraction
# observed median lower upper
# Equal Tailed
              0.44 0.4496 0.152 0.7088
# Highest Density
                0.44 0.4496 0.152 0.7088
# Hodges-Lehmann
# observed
         median
                 lower
# Equal Tailed
             -0.07335 -0.07615 -0.17220 -0.01565000
```

mf-class 5

```
# Highest Density -0.07335 -0.07615 -0.15635 -0.00850065
#
# Quartile Differences (quartiles of vac - quartiles of con)
# observed
             median
                       lower
                                 upper
# 025 -0.041500 -0.041500 -0.10340 -0.000905
# Q50 -0.112525 -0.111175 -0.28115 0.019350
# Q75 -0.168000 -0.170425 -0.38890 0.005300
# Quartiles of con
# observed median
                    lower
# Q25 0.054000 0.054000 0.021005 0.11275
# Q50 0.139275 0.139275 0.061400 0.31000
# Q75 0.315000 0.315000 0.173000 0.44625
# Ouartiles of vac
# observed median
                    lower
                             upper
# Q25 0.01250 0.01250 0.00125 0.026000
# Q50 0.02675 0.02675 0.01665 0.144575
# Q75 0.14700 0.14700 0.02810 0.219250
```

mf-class

Class mf

# Description

Parent class for package MF data objects.

# Usage

```
mf$new(nboot, alpha, seed, compare, rng)
```

### Fields

- nboot: numeric value specifying number of samples
- alpha: numeric value specifying complement of confidence interval
- seed: vector of integers specifying seed for pseudo-random number generator used
- compare: vector of character strings naming groups compared
- rng: character string naming type of random number generator

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

### See Also

Other mf: mfboot-class, mfbootcluster-class, mfhlboot-class

6 MFBoot

|--|

# Description

Estimates bootstrap confidence intervals for the mitigated fraction.

### Usage

```
MFBoot(formula, data, compare = c("con", "vac"), b = 100, B = 100,
alpha = 0.05, hpd = TRUE, bca = FALSE, return.boot = FALSE, trace.it = FALSE,
seed = sample(1:100000, 1))
```

#### **Arguments**

guments	
formula	Formula of the form $y \sim x$ , where y is a continuous response and x is a factor with two levels.
data	Data frame
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared
b	Number of bootstrap samples to take with each cycle
В	Number of cycles, giving the total number of samples = B * b
alpha	Complement of the confidence level
hpd	Estimate highest density intervals?
bca	Estimate BCa intervals?
return.boot	Save the bootstrap sample of the MF statistic?
trace.it	Verbose tracking of the cycles?
seed	to initialize random number generator for reproducibility. Passed to set.seed.

# **Details**

Resamples the data and produces bootstrap confidence intervals. Equal tailed intervals are estimated by the percentile method. Highest density intervals are estimated by selecting the shortest of all possible intervals. For BCa intervals, see Efron and Tibshirani section 14.3.

### Value

```
a mfboot-class data object
```

# Author(s)

David Siev <david.siev@aphis.usda.gov>

### References

Siev D. (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508** 

Efron B, Tibshirani RJ. An Introduction to the Bootstrap. Chapman and Hall, New York, 1993.

mfboot-class 7

### See Also

```
mfboot-class
```

### **Examples**

mfboot-class

Class mfboot

### **Description**

class for data objects produced by MFBoot, contains class mf with the two additional fields *stat* and *stuff*.

### Usage

```
mfboot$new(nboot, alpha, seed, compare, rng, sample, stat, stuff)
```

### **Fields**

- nboot: numeric value specifying number of samples
- alpha: numeric value specifying complement of confidence interval
- seed: vector of integers specifying seed for pseudo-random number generator used
- compare: vector of character strings naming groups compared
- rng: character string naming type of random number generator
- sample: what is this?
- stat: matrix of estimates

#### **Contains**

```
mf-class
```

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

# See Also

#### MFBoot

Other mf: mf-class, mfbootcluster-class, mfhlboot-class

8 mfbootcluster-class

mfbootcluster-class Class mfbootcluster

# **Description**

Class mfbootcluster is created from output of function MFClusBoot

### Usage

mfbootcluster\$new(nboot, alpha, seed, compare, rng, stat, what, excludedClusters,
call, sample)

### **Fields**

- nboot: numeric value specifying number of samples
- alpha: numeric value specifying complement of confidence interval
- seed: vector of integers specifying seed for pseudo-random number generator used
- compare: vector of character strings naming groups compared
- rng: character string naming type of random number generator
- stat: matrix matrix with columns observed, median, lower, upper for estimates
- what: character vector naming what was resampled: clusters, units, both
- excludedClusters: character vector naming clusters excluded because of missing treatment(s)
- call: the call to MFClusBoot
- sample: what is this?
- All: Field "All" from MFClus call.

#### **Contains**

mf-class

# Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

### See Also

### MFClusBoot

 $Other\ mf:\ mf-class,\ mfboot-class,\ mfhlboot-class$ 

MFClus 9

MFClus	Clustered mitigated fraction

### **Description**

Estimates mitigated fraction from clustered or stratified data.

# Usage

```
MFClus(formula, data, compare = c("con", "vac"), trace.it = FALSE)
```

# **Arguments**

formula	Formula of the form $y \sim x + cluster(w)$ , where y is a continuous response, x is a factor with two levels of treatment, and w is a factor indicating the clusters.
data	Data frame. See Note for handling of input data with more than two levels.
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared
trace.it	Verbose tracking of the cycles? Default FALSE.

### **Details**

Averages the U statistic over the clusters and computes MF from it. Clusters are excluded if they do not include both treatments.

### Value

```
a mfcluster-class data object
```

### Note

If input data contains more than two levels of treatment, rows associated with unused treatment levels will be removed.

Factor levels for treatments not present in the input data will be ignored.

Clusters with missing treatments will be excluded. See mfbootcluster-class or use trace.it to identify excluded clusters.

### Author(s)

```
David Siev <david.siev@aphis.usda.gov>
```

### References

Siev D. (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508** 

### See Also

```
mfcluster-class
```

10 MFClusBoot

### **Examples**

```
## Not run:
MFClus(lesion ~ group + cluster(litter), piglung)
# Comparing vac to con
#
# MF = 0.3533835
#
#
 By Cluster
#
                 r n1 n2
     w u
#
 U 25 10 0.4000000 5 5 -0.2000000
  K 12 2 0.2500000 4 2 -0.5000000
  Z 16 10 0.8333333 3 4 0.6666667
  D 3 2 1.0000000 1 2 1.0000000
  N 1
       0 0.0000000 1 3 -1.0000000
  T 8 5 0.8333333 2 3 0.6666667
  P 4 1 0.5000000 2 1 0.0000000
 L 3 2 0.6666667 1 3 0.3333333
 G 15 9 0.7500000 3 4 0.5000000
 J 15 9 1.0000000 3 3 1.0000000
 W 6 3 0.7500000 2 2 0.5000000
 A 9 3 0.3333333 3 3 -0.3333333
 X 12 6 1.0000000 3 2 1.0000000
 F 13 7 0.7777778 3 3 0.5555556
 S 21 11 0.9166667 4 3 0.8333333
 H 14 8 0.8888889 3 3 0.7777778
 Y 2 1 1.0000000 1 1 1.0000000
 E 2 1 1.0000000 1 1 1.0000000
#
#
  All
#
                   r n1 n2
        w u
                                 mf
  All 181 90 0.6766917 50 52 0.3533835
#
  Excluded Clusters
  [1] M, Q, R, B, O, V, I, C
## End(Not run)
```

MFClusBoot

Bootstrap MF CI from clustered data

### **Description**

Estimates bootstrap confidence intervals for the mitigated fraction from clustered or stratified data.

### Usage

```
MFClusBoot(formula, data, compare = c("con", "vac"), boot.cluster = TRUE,
boot.unit = TRUE, b = 100, B = 100, alpha = 0.05, hpd = TRUE,
return.boot = FALSE, trace.it = FALSE, seed = sample(1:1e+05, 1))
```

MFClusBoot 11

### **Arguments**

formula of the form y ~ x + cluster(w), where y is a continuous response, x

is a factor with two levels of treatment, and w is a factor indicating the clusters.

data Data frame. See Note for handling of input data with more than two levels.

compare Text vector stating the factor levels - compare[1] is the control or reference

group to which compare[2] is compared

boot.cluster Boolean whether to resample the clusters.

boot.unit Boolean whether to resample the units within cluster.

b Number of bootstrap samples to take with each cycle

B Number of cycles, giving the total number of samples = B \* b

alpha Complement of the confidence level

hpd Boolean whether to estimate highest density intervals.

return.boot Boolean whether to save the bootstrap sample of the MF statistic.

trace.it Boolean whether to display verbose tracking of the cycles.

seed to initialize random number generator for reproducibility. Passed to set.seed.

### **Details**

Resamples the data and produces bootstrap confidence intervals. Equal tailed intervals are estimated by the percentile method. Highest density intervals are estimated by selecting the shortest of all possible intervals.

#### Value

a mfbootcluster-class data object

#### Note

If input data contains more than two levels of treatment, rows associated with unused treatment levels will be removed.

Factor levels for treatments not present in the input data will be ignored.

Clusters with missing treatments will be excluded. See mfbootcluster-class or use trace.it to identify excluded clusters.

### Author(s)

David Siev <david.siev@aphis.usda.gov>

#### References

Siev D. (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508** 

Efron B, Tibshirani RJ. An Introduction to the Bootstrap. Chapman and Hall, New York, 1993.

12 MFClusBootHier

### **Examples**

MFClusBootHier

**MFClusBootHier** 

### **Description**

Combines MFhBoot and MFnestBoot into a single function.

# Usage

```
MFClusBootHier(formula, data, compare = c("con", "vac"), nboot = 10000,
boot.unit = TRUE, boot.cluster = TRUE, which.factor = "All",
alpha = 0.05, seed = sample(1:1e+05, 1))
```

# Arguments

8	
formula	formula Formula of the form $y \sim x + a/b/c$ , where y is a continuous response, x is a factor with two levels of treatment, and $a/b/c$ are grouping variables corresponding to the clusters. Nesting is assumed to be in order, left to right, highest to lowest. So a single level of "a" will contain multiple levels of "b" and a single level of "b" will contain multiple levels of "c".
data	a data.frame or tibble with the variables specified in formula. Additional variables will be ignored.
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared.
nboot	number of bootstrapping events
boot.unit	Boolean whether to sample observations from within those of the same core.
boot.cluster	Boolean whether to sample which cores are present. If TRUE, some trees have all the cores while others only have a subset.
which.factor	Which variables to include in the mitigated fraction summation. Default is $\hat{a} \in TM$ All $\hat{a} \in TM$ , to sum over entire tree.

mfclusboothier-class 13

alpha	Passed to emp. hpd to calculate high tailed upper and high tailed lower of miti-
	gated fraction.
seed	Passed to MFhBoot to to initialize random number generator for reproducibility.

### Value

A list with the following elements:

- **MFhBoot** as output from MFhBoot.
- MFnestBoot as output from MFnestBoot.

#### Note

Core variable is the variable corresponding to the lowest nodes of the hierarchical tree. Nest variables are those above the core. All refers to a summary of the entire tree.

#### See Also

MFhBoot, MFnestBoot.

# **Examples**

mfclusboothier-class Class mfclusboothier

### **Description**

Class mfclusboothier is created from output of function MFClusBootHier.

### Usage

```
mfclusboothier$new(MFhBoot, MFnestBoot)
```

### **Fields**

- MFhBoot: output from MFhBoot. A list.
- MFnestBoot: output from MFnestBoot. A list.

14 MFClusHier

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

### See Also

MFhBoot, MFnestBoot

# Description

Calculate mitigated fraction directly from hierarchial nested data. Combines MFh and MFnest into a single function.

# Usage

```
MFClusHier(formula, data, compare = c("con", "vac"), which.factor = "All")
```

# Arguments

formula	Formula of the form $y \sim x + a/b/c$ , where y is a continuous response, x is a factor with two levels of treatment, and $a/b/c$ are grouping variables corresponding to the clusters. Nesting is assumed to be in order, left to right, highest to lowest. So a single level of "a" will contain multiple levels of "b" and a single level of "b" will contain multiple levels of "c".
data	a data.frame or tibble with the variables specified in formula. Additional variables will be ignored.
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared.
which.factor	one or more variable(s) of interest. This can be any of the core or nest variables from the data set. If none or NULL is specified, MF will be calculated for the whole tree.

# Value

A list with the following elements:

- **MFh** as output from MFh.
- **MFnest** as output from MFnest.

# Note

Core variable is the variable corresponding to the lowest nodes of the hierarchical tree. Nest variables are those above the core. All refers to a summary of the entire tree.

### See Also

MFh, MFnest

mfclushier-class 15

### **Examples**

```
a <- data.frame(</pre>
room = paste('Room',rep(c('W','Z'),each=24)),
pen = paste('Pen',rep(LETTERS[1:6],each=8)),
litter = paste('Litter',rep(11:22,each=4)),
tx = rep(rep(c('vac', 'con'), each=2), 12),
stringsAsFactors = FALSE
set.seed(76153)
a$lung[a$tx=='vac'] <- rnorm(24,5,1.3)
a$lung[a$tx=='con'] <- rnorm(24,7,1.3)
thismf <- MFClusHier(lung ~ tx + room/pen/litter,a)</pre>
thismf$MFnest
aCore <- thismf$MFh
aCore
aCore$data
aCore$formula
aCore$compare
```

mfclushier-class

Class mfclushier

# Description

Class mfclushier is created from output of function MFClusHier

# Usage

```
mfclushier$new(MFh, MFnest)
```

### **Fields**

- MFh: output from MFh. A mfhierdata object.
- MFnest: output from MFnest. A tibble.

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

# See Also

```
MFh, MFnest
```

16 mfcomponents-class

mfcluster-class

Class mfcluster

### **Description**

Class mfcluster is created from output of function MFClus

# Usage

```
mfcluster$new(All, bycluster, excludedClusters, call, compare)
```

### **Fields**

- All: vector with elements:
  - w Wilcoxon statistic
  - u Mann-Whitney statistic
  - r mean ridit
  - n1 size of group 1
  - n2 size of group 2
  - mf mitigated fraction
- byCluster: As for All, by clusters
- excludedClusters: character vector naming clusters excluded because of missing treatment
- call: the call to MFClus
- compare: character vector naming groups compared

# Author(s)

 $Marie\ Vendettuoli\ \verb|\marie.c.vendettuoli@aphis.usda.gov>|$ 

### See Also

MFClus

mfcomponents-class

Class mfcomponents

# Description

Class mfcomponents is created from output of function MFSubj

# Usage

```
mfcomponents$new(mf, x, y, subj, compare)
```

MFh 17

#### **Fields**

- mf: numeric estimator for mitigated fraction
- x: numeric vector containing responses of group 1
- y: numeric vector containing responses of group 2
- subj: matrix where mf. j are the subject components
- compare: character vector naming groups being compared

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

### See Also

MFSubj

MFh

*Identify ranks for use when evaluating MF for nested hierarchy.* 

### Usage

```
MFh(formula, data, compare = c("con", "vac"))
```

### **Arguments**

formula Formula of the form  $y \sim x + a/b/c$ , where y is a continuous response, x is a factor

with two levels of treatment, and a/b/c are vgrouping variables corresponding to the clusters. Nesting is assumed to be in order, left to right, highest to lowest. So a single level of "a" will contain multiple levels of "b" and a single level of

"b" will contain multiple levels of "c".

data a data.frame or tibble with the variables specified in formula. Additional vari-

ables will be ignored.

compare Text vector stating the factor levels - compare[1] is the control or reference group

to which compare[2] is compared.

# Value

A mfhierdata object, which is a list of three items.

**coreTbl** A tibble with one row for each unique core level showing values for:

- con\_n & vac\_n counts of observations for each treatment level in the core level.
- *con\_*medResp & *vac\_*medResp median of the y continuous response for each treatment level.
- n1n2 product of the counts, con\_n \* vac\_n.
- w Wilcoxon statistic
- u Mann-Whitney statistic

data A tibble of the restructured input data used for calculations.

**compare** The compare variables as input by user.

formula The formula as input by user.

18 MFhBoot

#### Note

Core variable is the variable corresponding to the lowest nodes of the hierarchial tree. Nest variables are those above the core.

### See Also

MFnest for calculation of MF for nest, core and all variables. mfhierdata for returned object. MFClusHier for a wrapper.

### **Examples**

```
a <- data.frame(</pre>
room = paste('Room',rep(c('W','Z'),each=24)),
pen = paste('Pen',rep(LETTERS[1:6],each=8)),
 litter = paste('Litter',rep(11:22,each=4)),
 tx = rep(rep(c('vac', 'con'), each=2), 12),
 stringsAsFactors = FALSE
set.seed(76153)
a$lung[a$tx=='vac'] <- rnorm(24,5,1.3)
a$lung[a$tx=='con'] <- rnorm(24,7,1.3)
aCore <- MFh(lung ~ tx + room/pen/litter,a)
aCore
# A tibble: 12 x 10
     room pen litter con_medResp con_n
                                            w vac_medResp vac_n n1n2
     <chr> <chr> <chr>
                        <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
  1 Room W Pen A Litter 11
                              8.24 2 7
                                                   5.13
                                                           2
  2 Room W Pen A Litter 12
                              4.91 2 5
                                                   3.81
                                                            2
                                                                  4
                                                                       2
                              8.10 2 7
                                                                       4
  3 Room W Pen B Litter 13
                                                   5.23
                                                            2
                                                                 4
                             8.11 2 7
                                                                       4
  4 Room W Pen B Litter 14
                                                   5.59
                                                            2
                             8.09 2 7
                                                                       4
  5 Room W Pen C Litter 15
                                                   5.26
                                                            2
                                                                 4
                             6.77
                                                           2
                                    2
  6 Room W Pen C Litter 16
                                            7
                                                                       4
                                                   4.50
                                                                 4
                              5.58
   7 Room Z Pen D Litter 17
                                      2
                                            7
                                                    4.26
                                                            2
                                      2
                                                   6.33
   8 Room Z Pen D Litter 18
                               7.44
                                            6
                                                            2
                                                                       3
                              7.98
   9 Room Z Pen E Litter 19
                                      2
                                                    4.58
                                                            2
                                                                       4
# 10 Room Z Pen E Litter 20
                               6.78
                                       2
                                            7
                                                    4.86
                                                            2
                                                                       4
# 11 Room Z Pen F Litter 21
                               6.82
                                       2
                                            7
                                                    5.36
                                                            2
                                                                  4
                                                                       4
                                       2 7
# 12 Room Z Pen F Litter 22
                              7.27
                                                    5.13
                                                            2
                                                                  4
```

MFhBoot MFhBoot

### **Description**

Calculate rank tables for MF using bootstrapping.

# Usage

```
MFhBoot(formula, data, compare = c("con", "vac"), nboot = 10000,
boot.unit = TRUE, boot.cluster = TRUE, seed = sample(1:1e+05, 1))
```

**MFhBoot** 19

# **Arguments**

formula	Formula of the form $y \sim x + a/b/c$ , where y is a continuous response, x is a factor with two levels of treatment, and a/b/c are grouping variables corresponding to the clusters. Nesting is assumed to be in order, left to right, highest to lowest. So a single level of "a" will contain multiple levels of "b" and a single level of "b" will contain multiple levels of "c".
data	a data.frame or tibble with the variables specified in formula. Additional variables will be ignored.
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared.
nboot	number of bootstrapping events
boot.unit	Boolean whether to sample observations from within those of the same core.
boot.cluster	Boolean whether to sample which cores are present. If TRUE, some trees have all the cores while others only have a subset.
seed	to initialize random number generator for reproducibility. Passed to set. seed.

### Value

A list with the following elements:

bootmfh Rank table for the bootstrapped values as output from MFh. Includes a new bootID variable to distinguish each bootstrapped incidence.

clusters Table of unique nodes with an ID.

compare Compare vector as specified by user.

mfh MFh run on original data input.

### See Also

MFClusBootHier, MFnestBoot

# **Examples**

```
set.seed(76153)
a <- data.frame(room = paste('Room', rep(c('W', 'Z'), each = 24)),
                    pen = paste('Pen', rep(LETTERS[1:6], each = 8)),
                    litter = paste('Litter', rep(11:22, each = 4)),
tx = rep(rep(c('vac', 'con'), each = 2), 12),
                    stringsAsFactors = FALSE)
a[a$tx == 'vac', 'lung'] <- rnorm(24, 5, 1.3)
a[a$tx == 'con', 'lung'] <- rnorm(24, 7, 1.3)
formula <- lung \sim tx + room/pen/litter
nboot <- 10000
boot.cluster <- TRUE</pre>
boot.unit <- TRUE</pre>
which.factors <- c('All', 'room', 'pen', 'litter')</pre>
system.time(test1 <- MFhBoot(formula, a,</pre>
                                   nboot = 10000,
```

20 mfhlboot-class

boot.cluster = TRUE, boot.unit = TRUE, seed = 12345))

test1\$bootmfh

mfhierdata-class

Class mfhierdata

### **Description**

Class mfhierdata is created from output of function MFh

# Usage

```
mfhierdata$new(coreTbl, data)
```

#### **Fields**

- coreTb1: data.frame with one row for each unique core level showing values for nx, ny, N, w, u, and median observed response.
- data: data.frame is the restructured input data used for calculations in MFh and MFnest.
- compare: character vector naming groups being compared.
- formula: formula that was called by user.

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

# See Also

MFh

mfhlboot-class

Class mfhlboot

# Description

class for data objects produced by HLBoot, contains class mf with additional fields *MFstat*, *HLstat*, *QDIFstat*, *QXstat*, *QYstat* 

# Usage

```
mfhlboot$new(nboot, alpha, seed, compare, rng, sample, MFstat, HLstat,
QDIFstat, QXstat, QYstat)
```

MFmp 21

### **Fields**

- nboot: Numeric value specifying number of samples.
- alpha: Numeric value specifying complement of confidence interval.
- seed: Vector of integers specifying seed for pseudo-random number generator used.
- compare: Vector of character strings naming groups compared.
- rng: Character string naming type of random number generator.
- sample: The bootstrapped values.
- MFstatMatrix with columns *observed, median, lower, upper* for Equal Tailed and Highest Density estimates of mitigated fraction (MF).
- HLstatMatrix with columns *observed*, *median*, *lower*, *upper* for Equal Tailed and Highest Density estimates of Hodge-Lehmann estimator (HL).
- QDIFstatMatrix with columns *observed, median, lower, upper* for estimates of Quartile Differences.
- QXstatMatrix with columns *observed, median, lower, upper* for quartiles of treatments, equal tailed.
- QYstatMatrix with columns observed, median, lower, upper for quartiles of response, equal tailed.

#### **Contains**

mf-class

# Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

### See Also

**HLBoot** 

Other mf: mf-class, mfboot-class, mfbootcluster-class

MFmp

Mitigated fraction from matched pairs

# **Description**

Estimates mitigated fraction from matched pairs.

### Usage

```
MFmp(formula=NULL, data=NULL, compare = c("con", "vac"), x=NULL, alpha=0.05, df=NULL, tdist=T)
```

22 MFmp

# **Arguments**

formula	Formula of the form $y \sim x + \text{cluster}(w)$ , where y is a continuous response, x is a factor with two levels of treatment, and w is a factor indicating the clusters.	
data	Data frame	
compare	Text vector stating the factor levels - compare[1] is the control or reference group to which compare[2] is compared	
x	Trinomial vector $\{\Sigma I(x < y), \Sigma I(x = y), \Sigma I(x > y)\}$	
alpha	Complement of the confidence level.	
df	Degrees of freedom. Default N-2	
tdist	Use quantiles of t or Gaussian distribution for confidence interval? Default t distribution.	

### **Details**

Estimates MF from matched pairs by the difference of multinomial fractions  $(\Sigma I(x < y) - \Sigma I(x > y))/N$ . The trinomial vector is  $\{\Sigma I(x < y), \Sigma I(x = y), \Sigma I(x > y)\}$ 

### Value

```
a mfmp-class data object
```

### Note

upper confidence interval is truncated to 1; lower confidence interval is truncated to -1. Point estimate of 1.0 indicates complete separation.

### Author(s)

David Siev <david.siev@aphis.usda.gov>

### References

Siev D. (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508** 

### See Also

```
mfmp-class
```

# **Examples**

```
MFmp(les ~ tx + cluster(cage), mlesions, compare = c('con', 'vac')) MFmp(x = c(12, 12, 2))
```

mfmp-class 23

mfmp-class

Class mfmp

### **Description**

Class mfmp is created from output of function MFmp

### Usage

```
mfmp$new(ci, x, what, alpha, tdist, df)
```

### **Fields**

- ci: numeric vector of point and interval estimates
- x: numeric vector of length three holding data
- what: text string describing interval type
- alpha: numeric value specifying complement of confidence interval
- tdist: Logical indicating if t distribution(TRUE) or gaussian (FALSE)
- df: numeric value indicating degrees freedom

### Author(s)

Marie Vendettuoli <marie.c.vendettuoli@aphis.usda.gov>

# See Also

MFmp

MFnest

Summations to calculate the MF for nested data from a rank table.

### Usage

```
MFnest(Y, which.factor = "All")
```

# **Arguments**

Y rank table (tibble or data.frame), structured as \$coreTbl output from MFh or returned object from MFh().

which.factor

one or more grouping variable(s) of interest. This can be any of the core or nest variables from the data set. If none or All is specified, a summary MF will be calculated for the whole tree.

24 MFnest

#### Value

A tibble with each unique level of a variable as a row. Other values include:

MF Mitigated fraction for the particular level of the variable in this row.

- N1N2 Sum of the n1n2 variable in \$coreTb1 field of mfhierdata object output by MFh for this particular variable-level combination.
- U Sum of u variable in \$coreTbl field of mfhierdata object output by MFh for this particular variable-level combination.
- \_N Sum of the \_n variable in \$coreTbl field of mfhierdata object output by MFh for this particular variable-level combination.
- \_medResp Median of observed responses for each comparison group for this particular variable-level combination.

#### Note

Core variable is the variable corresponding to the lowest nodes of the hierarchial tree. Nest variables are those above the core. All refers to a summary of the entire tree.

### See Also

MFh

### **Examples**

```
a <- data.frame(
 room = paste('Room',rep(c('W','Z'),each=24)),
 pen = paste('Pen',rep(LETTERS[1:6],each=8)),
 litter = paste('Litter',rep(11:22,each=4)),
 tx = rep(rep(c('vac', 'con'), each=2), 12),
 stringsAsFactors = FALSE
set.seed(76153)
a\log[a\tx=='vac'] <- rnorm(24,5,1.3)
a = con' < -rnorm(24,7,1.3)
aCore <- MFh(lung \sim tx + room/pen/litter,a)
MFnest(aCore)
# # A tibble: 1 x 9
   variable level
                      MF N1N2
                                   U con_N vac_N con_medResp vac_medResp
            <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
                                                       <dbl>
    <fct>
                                                                   <dbl>
# 1 All
             All 0.875
                            48
                                  45
                                        24
                                                        7.24
                                                                    4.91
MFnest(aCore$coreTbl)
# Skipping median summary, no response data provided.
# # A tibble: 1 x 7
                      MF N1N2
   variable level
                                   U con_N vac_N
            <chr> <dbl> <dbl> <dbl> <dbl> <dbl>
    <fct>
# 1 All
             All 0.875
                            48
                                  45
                                        24
MFnest(aCore, 'room')
# # A tibble: 2 x 9
                       MF N1N2
                                    U con_N vac_N con_medResp vac_medResp
             <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
                                                        <dbl>
                                                                     <dbl>
```

MFnest 25

```
Room W 0.833
                          24
                                22
                                     12
                                           12
                                                    7.79
                                                               4.85
# 1 room
# 2 room
           Room Z 0.917
                          24
                                23
                                     12
                                           12
                                                    6.71
                                                               4.98
MFnest(aCore, 'pen')
# Complete separation observed for variable(s): pen
# # A tibble: 6 x 9
# variable level
                   MF N1N2
                                U con_N vac_N con_medResp vac_medResp
           <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
                                                  <dbl>
# 1 pen
           Pen A 0.5
                        8
                             6
                                   4
                                                  6.79
                                                              4.24
# 2 pen
           Pen B 1
                          8
                                8
                                                  8.11
                                                              5.59
# 3 pen
           Pen C 1
                                8
                                                   7.69
                                                              4.85
                             7
                                    4
# 4 pen
           Pen D 0.75
                          8
                                           4
                                                   6.10
                                                              4.98
# 5 pen
           Pen E 1
                          8
                               8
                                   4
                                           4
                                                   6.86
                                                              4.86
           Pen F 1
                          8
                                8
# 6 pen
                                     4
                                                   6.88
                                                              5.13
MFnest(aCore, c('All', 'litter'))
# Complete separation observed for variable(s): litter
# # A tibble: 13 x 9
    variable level
                       MF N1N2
                                    U con_N vac_N con_medResp vac_medResp
    <fct>
            <chr>
                      <dbl> <dbl> <dbl> <dbl> <dbl> <
                                                   <dbl>
                                                                  <dbl>
# 1 All
            A11
                      0.875
                            48
                                   45
                                        24
                                              24
                                                       7.24
                                                                   4.91
# 2 litter
            Litter 11 1
                              4
                                    4
                                          2
                                               2
                                                       8.24
                                                                   5.13
            Litter 12 0
# 3 litter
                               4
                                    2
                                          2
                                               2
                                                        4.91
                                                                   3.81
# 4 litter Litter 13 1
                              4
                                    4
                                          2
                                               2
                                                        8.10
                                                                   5.23
                                   4
# 5 litter Litter 14 1
                                                        8.11
                              4
                                          2
                                               2
                                                                   5.59
                                 4
# 6 litter Litter 15 1
                                                       8.09
                                                                   5.26
                               4
                                          2
                                               2
                               4 4
# 7 litter Litter 16 1
                                                       6.77
                                          2
                                               2
                                                                   4.50
                                        2
# 8 litter Litter 17 1
                                 4
                                               2
                                                        5.58
                                                                   4 26
                               4
                               4 3 2
# 9 litter
            Litter 18 0.5
                                               2
                                                       7.44
                                                                   6.33
# 10 litter Litter 19 1
                                          2
                                               2
                                                       7.98
                                                                   4.58
# 11 litter Litter 20 1
                                          2
                                               2
                                                       6.78
                                                                   4.86
# 12 litter Litter 21 1
                                          2
                                                                   5.36
                                               2
                                                        6.82
# 13 litter Litter 22 1
                               4
                                          2
                                                2
                                                        7.27
                                                                   5.13
MFnest(aCore, 'litter')
# Complete separation observed for variable(s): litter
# # A tibble: 12 x 9
                        MF N1N2
#
    variable level
                                    U con_N vac_N con_medResp vac_medResp
#
    <fct>
            <chr>
                      <dbl> <dbl> <dbl> <dbl> <dbl> <
                                                       <dbl>
                                                                  <fdh>>
  1 litter
            Litter 11 1
                                  4
                                        2
                                               2
                                                        8.24
                                                                   5.13
                             4
            Litter 12
  2 litter
                       0
                               4
                                    2
                                          2
                                               2
                                                        4.91
                                                                   3.81
# 3 litter
            Litter 13
                                          2
                                               2
                                                        8.10
                                                                   5.23
                       1
                               4
                                    4
# 4 litter
            Litter 14
                                    4
                                          2
                                               2
                                                        8.11
                                                                   5.59
                               4
            Litter 15
# 5 litter
                       1
                               4
                                    4
                                          2
                                                2
                                                        8.09
                                                                   5.26
# 6 litter
            Litter 16
                       1
                               4
                                    4
                                          2
                                               2
                                                        6.77
                                                                   4.50
# 7 litter
            Litter 17
                       1
                               4
                                    4
                                          2
                                               2
                                                        5.58
                                                                   4.26
                                    3
# 8 litter Litter 18
                       0.5
                               4
                                          2
                                               2
                                                        7.44
                                                                   6.33
# 9 litter Litter 19
                                    4
                                          2
                                               2
                                                       7.98
                                                                   4.58
                       1
                               4
# 10 litter
            Litter 20
                                    4
                                          2
                                               2
                                                        6.78
                                                                   4 86
                       1
# 11 litter
            Litter 21
                                         2
                                               2
                                                        6.82
                                                                   5.36
                       1
# 12 litter Litter 22
                       1
                                          2
                                               2
                                                        7.27
                                                                   5.13
MFnest(aCore, c('room', 'pen', 'litter'))
# # A tibble: 20 x 9
                        MF N1N2
#
    variable level
                                    U con_N vac_N con_medResp vac_medResp
    <fct> <chr>
                      <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
                                                                  < db1>
            Room W
# 1 room
                      0.833 24
                                                       7.79
                                                                   4.85
                                   22 12 12
```

26 MFnestBoot

#	2	room	Room Z	0.917	24	23	12	12	6.71	4.98
#	3	pen	Pen A	0.5	8	6	4	4	6.79	4.24
#	4	pen	Pen B	1	8	8	4	4	8.11	5.59
#	5	pen	Pen C	1	8	8	4	4	7.69	4.85
#	6	pen	Pen D	0.75	8	7	4	4	6.10	4.98
#	7	pen	Pen E	1	8	8	4	4	6.86	4.86
#	8	pen	Pen F	1	8	8	4	4	6.88	5.13
#	9	litter	Litter 11	1	4	4	2	2	8.24	5.13
#	10	litter	Litter 12	0	4	2	2	2	4.91	3.81
#	11	litter	Litter 13	1	4	4	2	2	8.10	5.23
#	12	litter	Litter 14	1	4	4	2	2	8.11	5.59
#	13	litter	Litter 15	1	4	4	2	2	8.09	5.26
#	14	litter	Litter 16	1	4	4	2	2	6.77	4.50
#	15	litter	Litter 17	1	4	4	2	2	5.58	4.26
#	16	litter	Litter 18	0.5	4	3	2	2	7.44	6.33
#	17	litter	Litter 19	1	4	4	2	2	7.98	4.58
#	18	litter	Litter 20	1	4	4	2	2	6.78	4.86
#	19	litter	Litter 21	1	4	4	2	2	6.82	5.36
#	20	litter	Litter 22	1	4	4	2	2	7.27	5.13

MFnestBoot MFn	nestBoot
----------------	----------

# Description

MFnest using bootstrapping

# Usage

```
MFnestBoot(x, which.factor = "All", alpha = 0.05)
```

# **Arguments**

Х	output from MFhBoot
which.factor	one or more grouping variable(s) of interest. This can be any of the core or nest variables from the data set. A MF value will be calculated for each level of the variable(s) specified. Default is $\hat{a} \in TM$ All $\hat{a} \in TM$ , to sum over entire tree.
alpha	Passed to emp. hpd to calculate eq tailed upper and high lower of mitigated fraction

### Value

A list with the following elements:

**mfnest\_details** The MF and summary statistics as calculated for each bootstrap event. Variables as in MFnest output.

**mfnest\_summary** Statistical summary of bootstrapped MF with each unique level of a core or nest variable passed to which.factor as a row. Other variables include:

- median Median of MFs from all of the bootstrap events.
- etlower Lower value of equal tailed range.

MFr 27

- etupper Upper value of equal tailed range.
- hdlower Lower value of the highest posterior density range.
- hdupper Upper value of the highest posterior density range.
- mf.obs MF calculated from data using MFh.

### See Also

MFClusBootHier, MFhBoot

### **Examples**

```
set.seed(76153)
a <- data_frame(room = paste('Room', rep(c('W','Z'), each = 24)),</pre>
                 pen = paste('Pen', rep(LETTERS[1:6], each = 8)),
                 litter = paste('Litter', rep(11:22, each = 4)),
                 tx = rep(rep(c('vac', 'con'), each = 2), 12)) %>%
  mutate(lung = ifelse(tx == 'vac', rnorm(24, 5, 1.3), rnorm(24, 7, 1.3)))
formula <- lung ~ tx + room/pen/litter</pre>
nboot <- 10000
boot.cluster <- TRUE</pre>
boot.unit <- TRUE</pre>
which.factors <- c('All', 'room', 'pen', 'litter')</pre>
##################
test1 <- MFhBoot(formula, a,</pre>
                  nboot = 10000,
                  boot.cluster = TRUE, boot.unit = TRUE, seed = 12345)
MFnestBoot(test1, c('All', 'litter'))
## Not run:
system.time(test2 <- MFnestBoot(test1, which.factors))</pre>
system.time(test3 <- MFnestBoot(test1, which.factors[1]))</pre>
system.time(test4 <- MFnestBoot(test1, which.factors[2]))</pre>
system.time(test5 <- MFnestBoot(test1, which.factors[2:3]))</pre>
system.time(test6 <- MFnestBoot(test1, which.factors[2:4]))</pre>
test6
## End(Not run)
```

MFr

Mitigated fraction

### **Description**

Mitigated fraction comparing treatment to control.

28 MFSubj

#### Usage

```
MFr(formula, data, compare = c("con", "vac"))
```

#### **Arguments**

formula Formula of the form  $y \sim x$ , where y is a continuous response and x is a factor

with two levels

data Data frame

compare Text vector stating the factor levels - compare[1] is the control or reference

group to which compare[2] is compared

### **Details**

The mitigated fraction is an estimator that quantifies an intervention's effect on reducing the severity of a condition. Since its units are on the probability scale, it is often a good idea to accompany it with an estimator on the original scale of measurement.

#### Value

The estimated mitigated fraction.

### Author(s)

David Siev

#### References

Siev D, 2005. An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. 4:500-508

# **Examples**

```
MFr(lesion~group,calflung)
# [1] 0.44
```

MFSubj

Subject components of mitigated fraction

## **Description**

Estimates the subject components of the mitigated fraction.

# Usage

```
MFSubj(formula, data, compare = c("con", "vac"))
```

### **Arguments**

formula Formula of the form  $y \sim x$ , where y is a continuous response and x is a factor

with two levels

data Data frame

compare Text vector stating the factor levels - compare[1] is the control or reference

group to which compare[2] is compared

mlesions 29

#### **Details**

The mitigated fraction is an estimator that quantifies an intervention's effect on reducing the severity of a condition. Since its units are on the probability scale, it is often a good idea to accompany it with an estimator on the original scale of measurement.

The subject components are the individual contributions of the treated subjects to MF, which is the average of the subject components.

### Value

```
a mfcomponents-class data object
```

### Author(s)

David Siev <david.siev@aphis.usda.gov>

#### References

Siev D. (2005). An estimator of intervention effect on disease severity. *Journal of Modern Applied Statistical Methods*. **4:500–508** 

### **Examples**

```
x <- MFSubj(lesion ~ group, calflung)</pre>
#
  MF = 0.44 comparing vac to con
#
# MF Subject Components
#
    mf.j freq
                 min.y max.y
            6 0.000030 0.00970
    1.00
    0.84
            1 0.012500 0.01250
#
    0.76
            3 0.016650 0.02030
#
    0.68
            6 0.023250 0.03190
#
    0.04
            1 0.132100 0.13210
   -0.04
            3 0.144575 0.16325
   -0.20
            2 0.210000 0.21925
   -0.36
            1 0.292000 0.29200
   -0.52
            1 0.356500 0.35650
   -0.84
            1 0.461500 0.46150
mean(x$subj[,'mf.j'])
# [1] 0.44
```

mlesions

mlesions dataset

### **Description**

Post-mortem examination of the lungs of dogs housed in cages by pairs.

30 piglung

### **Format**

A data frame with 52 observations of the following 3 variables, no NAs.

```
cage Cage ID. 1 - 26.tx Treatment. One of 'con' or 'vac'.les Percent gross lung lesions.
```

piglung

piglung dataset

# Description

Post-mortem examination of the lungs of pigs in litters.

# **Format**

A data frame with 102 observations of the following 3 variables, no NAs.

lesion Percent gross lung lesions.

group Treatment group. One of 'con' or 'vac'.

litter Litter ID.

# Index

*Topic datasets calflung, 3 mlesions, 29 piglung, 30 *Topic documentation mf-class, 5 mfboot-class, 7 mfbootcluster-class, 8 mfclusboothier-class, 13 mfclushier-class, 15 mfcluster-class, 16 mfcomponents-class, 16 mfhierdata-class, 20 mfhlboot-class, 20 mfmp-class, 23	mfcomponents-class, 16 MFh, 14, 15, 17, 19, 20, 23, 24, 27 MFhBoot, 12-14, 18, 26, 27 mfhierdata, 15, 17, 18, 24 mfhierdata (mfhierdata-class), 20 mfhierdata-class, 20 mfhlboot (mfhlboot-class), 20 mfhlboot-class, 20 MFmp, 21, 23 mfmp (mfmp-class), 23 mfmp-class, 23 MFnest, 14, 15, 18, 23, 26 MFnestBoot, 12-14, 19, 26 MFr, 27 MFSubj, 17, 28 mlesions, 29
<pre>calflung, 3 calflung-data(calflung), 3</pre>	mlesions-data (mlesions), 29
emp.hpd, 13, 26	piglung, 30 piglung-data(piglung), 30
HLBoot, 3, 21	tibble, <i>17</i>
MF (MF-package), 2 mf (mf-class), 5 mf-class, 5 MF-package, 2 MFBoot, 6, 7 mfboot (mfboot-class), 7 mfboot-class, 7 mfbootcluster (mfbootcluster-class), 8 mfbootcluster-class, 8 MFClus, 9, 16 MFClusBoot, 8, 10 MFClusBootHier, 12, 19, 27 mfclusboothier (mfclusboothier-class),  13 mfclusboothier-class, 13 MFClusHier, 14, 18 mfclushier (mfclushier-class), 15 mfclushier (mfclushier-class), 16 mfcluster (mfcluster-class), 16 mfcluster-class, 16 mfcomponents (mfcomponents-class), 16	