Package 'choplump'

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Title Permutation Test for Some Positive and Many Zero Responses

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Description Calculates permutation tests that can be powerful for comparing two groups with some positive but many zero responses (see Follmann, Fay, and Proschan < DOI:10.1111/j.1541-0420.2008.01131.x>).
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choplump-package

Permutation Test for Some Positive and Many Zero Responses

Description

This package has basically one important function, choplump for performing the choplump test, which is for comparing two groups with some positive response and many zero responses. These tests can often be more powerful than simpler permutation tests. Exact and approximation methods are available for calculating p-values.

Details

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License: GPL

See example below. There is also two vignettes. The vignette computation (see vignette("choplumpComputation")) gives computational details, and the vignette validation (see vignette("choplumpValidation")) details the way we have validated the function.

Author(s)

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References

Follmann, DA, Fay, MP, and Proschan, MA. (2009) "Chop-lump tests for Vaccine trials" Biometrics 65: 885-893. (see /doc/choplump.pdf)

Examples

```
set.seed(13921)
Ntotal<-200
Mtotal<-54
Z<-rep(0,Ntotal)
Z[sample(1:Ntotal,Ntotal/2,replace=FALSE)]<-1
test<-data.frame(W=c(rep(0,Ntotal-Mtotal),abs(rnorm(Mtotal))),Z=Z)
## defaults to asymptotic approximation if the number
## of calculations of the test statistic
## is >methodRuleParms=10^4
choplump(W~Z,data=test,use.ranks=TRUE,exact=FALSE)
```

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chooseMatrix

Create an (n choose m) by n matrix with unique rows.

Description

Create a choose(n,m) by n matrix. The matrix has unique rows with m ones in each row and the rest zeros.

Usage

```
chooseMatrix(n, m)
```

Arguments

m

an integer n an integer<=n

Value

A matrix with choose(n,m) rows n columns. The matrix has unique rows with m ones in each row and the rest zeros.

Note

Used for exact test method for choplump

Author(s)

M.P.Fay

Examples

```
chooseMatrix(5,2)
```

choplump

Choplump Test

Description

The choplump test is a two-sample permutation test, that is used when there are many responses that are zero with some positive.

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Usage

```
choplump(x, ...)
## Default S3 method:
choplump(x, y, alternative = c("two.sided", "less", "greater"),
            use.ranks=TRUE, exact = NULL, method=NULL,
            methodRule=methodRule1, methodRuleParms=c(10^4),
            nMC=10^4-1, seed=1234321, printNumCalcs=TRUE, ...)
## S3 method for class 'formula'
choplump(formula, data, subset, na.action, ...)
```

Arguments

a numeric vector of responses in first group, or a formula. Should have some

zeros and the rest positive.

numeric vector of responses in second group

alternative a character string specifying the alternative hypothesis, must be one of "two.sided"

(default), "greater" or "less".

use.ranks a logical indicating whether to use ranks for the responses

a logical indicating whether an exact p-value should be computed (see details) exact method

a character value, one of 'approx', 'exact', 'exactMC'. If NULL method chosen

by methodRule

methodRule a function used to choose the method (see details). Ignored if method is not

NULL

methodRuleParms

a vector of parameters passed to methodRule. Ignored if method is not NULL

nMC number of Monte Carlo replications, used if method='exactMC', ignored other-

wise

value used in set.seed if method='exactMC', ignored otherwise seed printNumCalcs logical, print number of calculations of test statistic for exact tests

formula a formula of the form lhs~rhs where lhs is a numeric variable giving the data

values and rhs a factor with two levels giving the corresponding groups.

data an optional matrix or data frame containing the variables in the formula

an optional vector specifying a subset of observations to be used. subset

na.action a function which indicates what should happen when the data contain NAs. De-

faults to getOption("na.action").

further arguments to be passed to or from methods.

Details

Consider a randomized trial where one wants to compare the responses in two groups, but there are many zeros in both groups. For example, in an HIV vaccine trial the response could be level of virus in the blood and very many in both groups will have zero values for the response. In order to gain choplump 5

power, the choplump test removes the same proportion of zeros from both groups, and compares the standardized means between the values left. The test can use ranks to obtain a Wilcoxon-like test. The choplump is a formal permutation test (in other words for each permutation, the chopping is redone) so the type I error is less than the nominal significance level either exactly (for exact methods) or approximately (for the approximate method).

There are a choice of 3 different methods to calculate the p-values: approx, an approximation method, see vignette("choplumpComputation"); exact, an exact method, see vignette("choplumpComputation"); exactMC, exact method using Monte Carlo resampling with nMC resamples.

The associated functions for the above methods (choplumpApprox, choplumpExact, choplumpExactMC), are internal and not to be called directly.

A methodRule function has 4 input values: W (a vector of all responses), Z (a vector of 0 or 1 denoting group membership), exact (a logical value, same as exact in the choplump call), and parms (the vector of parameters, same as methodRuleParms in the choplump call). The methodRule function returns a character vector with one of the allowed methods. The default method rule is methodRule1. It gives a result of 'approx' if either exact=FALSE or exact=NULL and there are more than parms calculations of the test statistic. It gives a result of 'exact' if there are less than methodRuleParms calculations of the test statistic, and it gives a result of 'exactMC' if exact=TRUE and there are more than methodRuleParms calculations of the test statistic.

Value

A htest object, a list with elements

p.value p value associated with alternative alternative description of alternative hypothesis

p. values a vector giving lower, upper, and two-sided p-values

METHOD a character vector describing the test

data.name a character vector describing the two groups

Author(s)

M.P. Fay

References

Follmann, DA, Fay, MP, and Proschan, MA. (2009) "Chop-lump tests for Vaccine trials" Biometrics 65: 885-893. (see /doc/choplump.pdf)

Examples

```
set.seed(1)
Ntotal<-200
Mtotal<-12
Z<-rep(0,Ntotal)
Z[sample(1:Ntotal,Ntotal/2,replace=FALSE)]<-1
test<-data.frame(W=c(rep(0,Ntotal-Mtotal),abs(rnorm(Mtotal))),Z=Z)
## defaults to asymptotic approximation if
## the number of calculations of the test</pre>
```

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```
## statistic is greater than parms
## see help for methodRule1
choplump(W~Z,data=test,use.ranks=TRUE)
## alternate form
cout<-choplump(test$W[test$Z==0],test$W[test$Z==1],use.ranks=TRUE,exact=TRUE)
cout
cout$p.values</pre>
```

choplumpGeneral

General choplump test

Description

This function does a general choplump test. For simple difference in standardized means (on the responses or on the ranks), use the much faster choplump function.

Usage

```
choplumpGeneral(W, Z, testfunc=testfunc.wilcox.ties.general)
```

Arguments

W numeric vector of responses, some should be zero

Z numeric vector of group membership, values either 0 or 1

test function, inputs a data frame with two columns labeled W and Z, outputs

test statistic

Value

Returns a p-value vector of length 3, with 3 named values: p.lower, p.upper, p.2sided.

Examples

```
### compare speed and results using two different functions
W < -c(0,0,0,0,0,0,0,0,2,4,6)
Z < -c(0,0,0,0,1,1,1,1,0,1,1)
Testfunc<-function(d){</pre>
     W<-d$W
     Z<-d$Z
     N<-length(Z)
     sqrt(N-1)*(sum(W*(1-Z)) - N*mean(W)*mean(1-Z))/
       sqrt(var(W)*var(1-Z))
}
time0<-proc.time()</pre>
choplumpGeneral(W,Z,Testfunc)
time1<-proc.time()</pre>
choplump(W~Z,use.ranks=FALSE)$p.values
time2<-proc.time()</pre>
time1-time0
time2-time1
```

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Description

This is the default function which determines which method to use in choplump.

Usage

```
methodRule1(W,Z, exact, parms)
```

Arguments

W	numeric vector of response scores, usually many zeros and the rest positive
Z	group membership vector, values all 0 (control) or 1 (treated)
exact	logical, TRUE=exact method, FALSE=approximate method, NULL=see below
parms	numeric value of maximum number of calculations of test statistic, if number of calculations greater than parms then use Monte Carlo for exact method

Details

This function determines which of several methods will be used in choplump; see that help for description of methods.

When exact=FALSE then returns 'approx'. When exact=TRUE then returns either 'exact' if the number of calculations of the test statistic is less than or equal to parms or 'exactMC' otherwise. When exact=NULL then returns either 'exact' if the number of calculations of the test statistic is less than or equal to parms or or 'approx' otherwise.

Value

```
a character vector with one of the following values: "approx", "exact", "exactMC"
```

See Also

choplump

wilcox.manyzeros.exact

Wilcoxon Rank Sum Test

Description

This function gives exact p-values for the Wilcoxon rank sum. This algorithm is designed for the case when the responses are either positive or zero, and there are many zero responses. Its purpose is mostly for the validation of the choplump function (see vignette("choplumpValidation")).

Usage

```
wilcox.manyzeros.exact(W, Z)
```

Arguments

W a vector of responses, should have some zeros and all rest positive

Z a vector of group membership, should be either 0 or 1

Value

A vector of three types of p-values: p.lower, p.upper, and p.2sided.

Author(s)

M.P. Fay

See Also

choplump

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