

# Package: someMTP (via r-universe)

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**Author** livio finos  
**Maintainer** livio finos <livio@stat.unipd.it>  
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someMTP-package

*Some Multiple Testing Procedures*

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## Description

It is a collection of functions for Multiplicity Correction and Multiple Testing.

## Details

Package: someMTP  
Type: Package  
Version: 1.2  
Date: 2011-01-10  
License: GPL (>= 2)  
LazyLoad: yes

## Author(s)

livio finos

Maintainer: <livio@stat.unipd.it>

## References

For weighted methods:

Benjamini, Hochberg (1997). Multiple hypotheses testing with weights. *Scand. J. Statist.* 24, 407-418.

Finos, Salmaso (2007). FDR- and FWE-controlling methods using data-driven weights. *Journal of Statistical Planning and Inference*, 137,12, 3859-3870.

For LSD test:

J. Lauter, E. Glimm and S. Kropf (1998). Multivariate test based on Left-Spherically Distributed Linear Scores. *The Annals of Statistics*, Vol. 26, No. 5, 1972-1988

L. Finos (2011). A note on Left-Spherically Distributed Test with covariates, *Statistics and Probability Letters*, Volume 81, Issue 6, June 2011, Pages 639-641

## Examples

```
set.seed(13)
y <- matrix(rnorm(5000),5,1000) #create toy data
y[,1:100] <- y[,1:100]+3 #create toy data

p <- apply(y,2,function(y) t.test(y)$p.value) #compute p-values
M2 <- apply(y^2,2,mean) #compute ordering criterion
```

```
fdr <- p.adjust(p,method="BH") #(unweighted) procedure, fdr control
sum(fdr<.05)
fdr.w <- p.adjust.w(p,method="BH",w=M2) #weighted procedure, weighted fdr control
sum(fdr.w<.05)

fwer <- p.adjust(p,method="holm") #(unweighted) procedure, fwer control
sum(fwer<.05)
fwer.w <- p.adjust.w(p,method="BHfwe",w=M2) #weighted procedure, weighted fwer (=fwer) control
sum(fwer.w<.05)

plot(M2,-log10(p))
```

---

*OrNULL-class	<i>Class *OrNULL</i>
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**Description**

class \* or Null

**Objects from the Class**

A virtual Class: No objects may be created from it.

**Methods**

No methods defined with class "\*OrNULL" in the signature.

**Examples**

```
showClass("call10rNULL")
```

---

draw	<i>Plots results of fdrOrd()</i>
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**Description**

Plots results of fdrOrd()

**Usage**

```
draw(object, what = c("all", "ordVsP", "stepVsR"), pdfName = NULL)
```

**Arguments**

- object            a someMTP.object resulting from fdrOrd()
- what             what to plot; "all" is the default
- pdfName          it is the pdf filename where the plot will be saved. If pdfName is null (the default) the plot will show as window.

**Value**

No value is returned

**Author(s)**

Livio Finos

**See Also**

See Also [fdrOrd](#).

**Examples**

```
set.seed(17)
x=matrix(rnorm(60),3,20)
x[,1:10]=x[,1:10]+2 ##variables 1:10 have tests under H1
ts=apply(x,2,function(x) t.test(x)$statistic)
ps=apply(x,2,function(x) t.test(x)$p.value)
m2=apply(x^2,2,mean)
p0rd <- fdrOrd(ps,q=.05,ord=m2)
draw(p0rd)
```

---

fdrOrd/kfweOrd

*Controlling the False Discovery Rate and the Generalized FWER  
in ordered Test*

---

**Description**

Ordinal procedure controlling the FDR and the Generalized FWER

**Usage**

```
fdrOrd(p, q = .01, ord = NULL, GD=FALSE)
kfweOrd(p, k = 1, alpha = 0.01, ord = NULL, alpha.prime = alpha,
        J = qnbinom(alpha, k, alpha.prime), GD = FALSE)
```

**Arguments**

p	vector of p-values
ord	Values on the basis of which the procedure select the hypotheses (following decreasing order). The vector have the same length of p. If NULL the natural ordering is considered.
q	average FDR level
alpha	global significance level
k	number of allowed errors in kFWE controls
J	number of allowed jumps befor stopping

alpha.prime      univariate alpha for single step Guo and Romano procedure  
 GD                Logic value. Should the correction for general dependence be applied?

### Value

The function returns an object of class `someMTP.object`.

rej:                a logical vector indicating whenever the related hypothesis have been rejected.  
 p:                 the vector of p-values used in the call  
 ord:                The vector used to sort the p-values (decreasing).  
 MTP:               "fdrOrd" or "kfweOrd"  
 GD:                A logical value incating if the correction for General Dependence have been used or not.  
 q:                 The level of controlled FDR.  
 alpha:             The level of controlled k-FWER  
 alphaprime:       The significance level of individual tests  
 k:                 Number of allowed Errors  
 J:                 Number of allowed Jumps

### Author(s)

L. Finos and A. Farcomeni

### References

L. Finos, A. Farcomeni (2011). k-FWER Control without p-value Adjustment, with Application to Detection of Genetic Determinants of Multiple Sclerosis in Italian Twins. *Biometrics*.  
 A. Farcomeni, L. Finos (2013). FDR Control with Pseudo-Gatekeeping Based on a Possibly Data Driven Order of the Hypotheses. *Biometrics*.

### See Also

See also [draw](#)

### Examples

```
set.seed(17)
x=matrix(rnorm(60),3,20)
x[,1:10]=x[,1:10]+2 ##variables 1:10 have tests under H1
ts=apply(x,2,function(x) t.test(x)$statistic)
ps=apply(x,2,function(x) t.test(x)$p.value) #compute p-values
m2=apply(x^2,2,mean) #compute ordering criterion

p0rd <- fdrOrd(ps,q=.05,ord=m2) #ordinal Procedure
p0rd
draw(p0rd)
```

```

sum(p.adjust(ps,method="BH")<=.05) #rejections with BH

kOrd <- kfweOrd(ps,k=5,ord=m2)#ordinal procedure
kOrd
kOrdGD <- kfweOrd(ps,k=5,ord=m2,GD=TRUE)#ord. proc. (any dependence)
kOrdGD

```

---

lsd.object class      *Class "lsd.object" for storing the result of the function lsd*

---

### Description

The class lsd.object is the output of a call to [lsd.test](#)

### Slots

**F** : the test statistic  
**df** : the degrees of freedom of F  
**globalP**: the associated p-value  
**D**: the matrix used in the test (it provides the influence of columns in resp to the test statistic)  
**call**: The matched call to [lsd](#).  
**MTP**: The procedure used ("fdrOrd", "kfweOrd" or others).

### Methods

**p.value** (lsd.object): Extracts the p-values.  
**show** lsd.object: Prints the test results: p-value, test statistic, expected value of the test statistic under the null hypothesis, standard deviation of the test statistic under the null hypothesis, and number of covariates tested.  
**summary** lsd.object: Prints the test results: p-value, test statistic, expected value of the test statistic under the null hypothesis, standard deviation of the test statistic under the null hypothesis, and number of covariates tested.  
**weights** lsd.object: diagonal of matrix D used in the test (i.e. the influence of columns in resp to the test statistic)

### Author(s)

Livio Finos: <livio@stat.unipd.it>

### See Also

[lsd](#)

**Examples**

```
# Simple examples with random data here
set.seed(1)
#Standard multivariate LSD test for one sample case
X=matrix(rnorm(50),5,10)+5
res <- lsd.test(resp=X,alternative=~1)
print(res)
p.value(res)
summary(res,showD=TRUE)
```

---

lsd.test	<i>Multivariate Left Spherically Distributed (LSD) linear scores test.</i>
----------	--

---

**Description**

It performs the multivariate Left Spherically Distributed linear scores test of L\auter et al. (The Annals of Statistics, 1998) (see also details below).

**Usage**

```
lsd.test(resp, alternative = 1, null = NULL, D = NULL, data=NULL)
```

**Arguments**

resp	The response vector of the regression model. May be supplied as a vector or as a <a href="#">formula</a> object. In the latter case, the right hand side of Y is passed on to alternative if that argument is missing, or otherwise to null.
alternative	The part of the design matrix corresponding to the alternative hypothesis. The covariates of the null model do not have to be supplied again here. May be given as a half <a href="#">formula</a> object (e.g. ~a+b). In that case the intercept is always suppressed.
null	The part of the design matrix corresponding to the null hypothesis. May be given as a design matrix or as a half <a href="#">formula</a> object (e.g. ~a+b). The default for Z is ~1, i.e. only an intercept. This intercept may be suppressed, if desired, with Z = ~0.
data	Only used when Y, X, or Z is given in formula form. An optional data frame, list or environment containing the variables used in the formulae. If the variables in a formula are not found in data, the variables are taken from environment(formula), typically the environment from which gt is called.
D	is q x p matrix or it is a function with arguments resp and null returning the q x p transformation matrix. When D = NULL, then $D = \text{diag}(t(\text{resp})\%*\%IP0\%*\%\text{resp})$ with $IP0 = \text{diag}(n) - \text{null}\%*\%\text{solve}(t(\text{null})\%*\%\text{null})\%*\%t(\text{null})$

**Value**

The function returns an object of class `lsd.object`.

F	the test statistic
df	the degrees of freedom of F
p	the associated p-value
D	the matrix used in the test (it provide information on the influence of columns in resp to the test)
call:	The matched call to <code>lsd.test</code> .

**Author(s)**

Livio Finos

**References**

J. Laeuter, E. Glimm and S. Kropf (1998) Multivariate test based on Left-Spherically Distributed Linear Scores. The Annals of Statistics, Vol. 26, No. 5, 1972-1988

L. Finos (2011). A note on Left-Spherically Distributed Test with covariates, Statistics and Probability Letters, Volume 81, Issue 6, June 2011, Pages 639-641

**Examples**

```
set.seed(1)
#Standard multivariate LSD test for one sample case
X=matrix(rnorm(50),5,10)+2
lsd.test(resp=X,alternative=~1)

#Standard multivariate LSD test for two sample case
X2=X+matrix(c(0,0,1,1,1),5,10)*10
lsd.test(resp=X2,null=~1,alternative=c(0,0,1,1,1))

#General multivariate LSD test for linear predictor with covariates
lsd.test(resp=X2,null=cbind(rep(1,5),c(0,0,1,1,1)),alternative=1:5)
```

---

p.adjust.w

*Adjust P-values for Multiple Comparisons*

---

**Description**

Given a set of p-values, returns p-values adjusted using one of several (weighted) methods. It extends the method of `p.adjust{stats}`

**Usage**

```
p.adjust.w(p, method = c("bonferroni", "holm", "BHfwe", "BH", "BY"), n = length(p), w=NULL)
```



**Arguments**

p	vector of p-values (possibly with NAs)
method	correction method
n	number of comparisons, must be at least length(p); only set this (to non-default) when you know what you are doing!
w	weights to be used. p.adjust.w(..., rep(1, length(p))) produces the same results as in p.adjust(...) (i.e. the unweighted counterpart).

**Value**

A vector of corrected p-values (same length as p) having two attributes: attributes(...)\$w is the vector of used weights and attributes(...)\$method is the method used.

**Author(s)**

Livio Finos

**References**

Benjamini, Hochberg (1997). Multiple hypotheses testing with weights. *Scand. J. Statist.* 24, 407-418.

Finos, Salmaso (2007). FDR- and FWE-controlling methods using data-driven weights. *Journal of Statistical Planning and Inference*, 137,12, 3859-3870.

**See Also**

[p.adjust](#)

**Examples**

```
set.seed(13)
y <- matrix(rnorm(5000),5,1000) #create toy data
y[,1:100] <- y[,1:100]+3 #create toy data

p <- apply(y,2,function(y) t.test(y)$p.value) #compute p-values
M2 <- apply(y^2,2,mean) #compute ordering criterion

fdr <- p.adjust(p,method="BH") #(unweighted) procedure, fdr control
sum(fdr<.05)
fdr.w <- p.adjust.w(p,method="BH",w=M2) #weighted procedure, weighted fdr control
sum(fdr.w<.05)

fwer <- p.adjust(p,method="holm") #(unweighted) procedure, fwer control
sum(fwer<.05)
fwer.w <- p.adjust.w(p,method="BHfwe",w=M2) #weighted procedure, weighted fwer (=fwer) control
sum(fwer.w<.05)

plot(M2,-log10(p))
```

---

someMTP.object class *Class "someMTP.object" for storing the result of the function fdrOrd*

---

### Description

The class someMTP.object is the output of a call to [fdrOrd](#). It also stores the information needed for related plots.

### Slots

**rej**: a logical vector indicating whenever the related hypothesis have been rejected.

**p**: The vector of (raw) p-values used in the procedure.

**ord**: The vector used to sort the p-values (decreasing).

**idOrd**: The vector of indices used in sorting.

**MTP**: The type of procedure used.

**GD**: A logical value incating if the correction for General Dependence have been used or not.

**q**: The level of controlled FDR when MTP=="fdrOrd".

**k**: The number of false rejection when MTP=="kfweOrd"

**J**: The number of allowed Jumps when MTP=="kfweOrd"

**alpha**: The significance level when MTP=="kfweOrd"

**alphaprime**: The significance level of individual tests.

**call**: The cal that generates the object.

### Methods

**show** someMTP.object: Prints the test results.

**summary** someMTP.object: Prints the test results (as show).

**draw** someMTP.object: Plots results; what = c("all", "ordVsP", "stepVsR")

**sort** signature(x = "someMTP.object"): Sorts the p-values to decreasing order of ord.

**length** signature(x = "someMTP.object"): The number of tests performed.

**names** signature(x = "someMTP.object"): Extracts the row names of the results matrix.

**names<-** signature(x = "someMTP.object"): Changes the row names of the results matrix. Duplicate names are not allowed, but see alias.

### Author(s)

Livio Finos: <livio@stat.unipd.it>

### See Also

[someMTP.object](#)

**Examples**

```

# Simple examples with random data
set.seed(17)
x=matrix(rnorm(60),3,20)
x[,1:10]=x[,1:10]+2 ##variables 1:10 have tests under H1
ts=apply(x,2,function(x) t.test(x)$statistic)
ps=apply(x,2,function(x) t.test(x)$p.value)
m2=apply(x^2,2,mean)
p0rd <- fdr0rd(ps,q=.05,ord=m2)
p0rd
  length(p0rd)
names(p0rd) <- paste("V",1:20,sep="")
names(p0rd)

```

step.adj

*Multiplicity correction for Stepwise Selected models***Description**

Corrects the p-value due to model selection. It works with models of class `glm` and selected with function `step {stats\}`.

**Usage**

```

step.adj(object, MC = 1000, scope = NULL, scale = 0,
         direction = c("both", "backward", "forward"),
         trace = 0, keep = NULL, steps = 1000, k = 2)

```

**Arguments**

object	object of class <code>glm</code> . Note that formula have to write by variables name like <code>y~var1+var2+var3</code> , data is a <code>data.frame</code> (see example below), <code>offset</code> is not yet implemented, avoid its use, <code>glm(formula, data, family=gaussian)</code> produce the same result of <code>lm(formula, data)</code> , then linear model can be allways performed
MC	number of random permutations for the dependent variable
scope	as in function <code>step</code>
scale	as in function <code>step</code>
direction	as in function <code>step</code>
trace	as in function <code>step</code>
keep	as in function <code>step</code>
steps	as in function <code>step</code>
k	as in function <code>step</code> , other arguments are not implemented yet.

**Details**

It performs anova function (stats library) on the model selected by function step vs the null model with the only intercept and it corrects for multiplicity. For lm models and gaussian glm models it computes a F-test, for other models it uses Chi-square-test (see also anova.glm and anova.lm help).

**Value**

An anova table with an extra column reporting the corrected p-value

**Author(s)**

Livio Finos and Chiara Brombin

**References**

L. Finos, C. Brombin, L. Salmaso (2010). Adjusting stepwise p-values in generalized linear models. Communications in Statistics - Theory and Methods.

**See Also**

[glm](#), [anova](#)

**Examples**

```
set.seed(17)
y=rnorm(10)
x=matrix(rnorm(50),10,5)
#define a data.frame to be used in the glm function
DATA=data.frame(y,x)
#fit the model on a toy dataset
mod=glm(y~X1+X2+X3+X4+X5,data=DATA)

#select the model using function step
mod.step=step(mod, trace=0)
#test the selected model vs the null model
anova(glm(y~1, data=DATA),mod.step,test="F")

#step.adj do the same, but it also provides multiplicity control
step.adj(mod,MC=101, trace=0)
```

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