

# Package: flip (via r-universe)

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**Type** Package

**Title** Multivariate Permutation Tests

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**Description** It implements many univariate and multivariate permutation (and rotation) tests. Allowed tests: the t one and two samples, ANOVA, linear models, Chi Squared test, rank tests (i.e. Wilcoxon, Mann-Whitney, Kruskal-Wallis), Sign test and Mc Nemar. Test on Linear Models are performed also in presence of covariates (i.e. nuisance parameters). The permutation and the rotation methods to get the null distribution of the test statistics are available. It also implements methods for multiplicity control such as Westfall & Young minP procedure and Closed Testing (Marcus, 1976) and k-FWER. Moreover, it allows to test for fixed effects in mixed effects models.

**License** GPL (>= 2)

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**LazyData** true

**NeedsCompilation** yes

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flip-package	<i>flip-package</i>
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## Description

The library is devoted to permutation-based inferential methods.

It implements many univariate and multivariate permutation (and rotation) tests.

The tests comprised are: the one and two samples, ANOVA, linear models, Chi Squared test, rank tests (i.e. Wilcoxon, Mann-Whitney, Kruskal-Wallis), Kolmogorov-Smirnov and Anderson-Darling.

Test on Linear Models are performed also in presence of covariates (i.e. nuisance parameters).

The permutation and the rotation method to get the null distribution of the test statistic(s) are available.

It also implements methods for multiplicity control such as Westfall-Young min-p procedure and Closed Testing (Marcus, 1976).

```

Package:    flip
Type:      Package
Version:    1.1
Date:      2012-02-05
License:    GPL <=2
LazyLoad:  yes
Depends:    methods, e1071, someMTP, cherry

```

## Author(s)

Livio Finos, with contributions by Florian Klinglmueller, Dario Basso, Aldo Solari, Lucia Benetazzo, Jelle Goeman and Marco Rinaldo. Special thanks are due to Ivan Marin-Franch and Fredrik Nilsson for the debugging and the good questions.

Maintainer: livio finos <livio@stat.unipd.it>

## References

For the general framework of univariate and multivariate permutation tests see: Pesarin, F. (2001) *Multivariate Permutation Tests with Applications in Biostatistics*. Wiley, New York.

#' @references For the general framework of univariate and multivariate permutation tests see: Pesarin, F. (2001) *Multivariate Permutation Tests with Applications in Biostatistics*. Wiley, New York.

Livio Finos and Fortunato Pesarin (2018) On zero-inflated permutation testing and some related problems. *Statistical Papers*.

For analysis of mixed-models see: L. Finos and D. Basso (2014) Permutation Tests for Between-Unit Fixed Effects in Multivariate Generalized Linear Mixed Models. *Statistics and Computing*. Volume 24, Issue 6, pp 941-952. DOI: 10.1007/s11222-013-9412-6 J. J. Goeman and

D. Basso, L. Finos (2011) Exact Multivariate Permutation Tests for Fixed Effects in Mixed-Models. *Communications in Statistics - Theory and Methods*. DOI 10.1080/03610926.2011.627103

For Rotation tests see: Langsrud, O. (2005) Rotation tests, *Statistics and Computing*, 15, 1, 53-60

A. Solari, L. Finos, J.J. Goeman (2014) Rotation-based multiple testing in the multivariate linear model. *Biometrics*, 70 (4), 954-961.

## Examples

```
Y=data.frame(matrix(rnorm(50),10,5))
names(Y)=LETTERS[1:5]
Y[,1:2]=Y[,1:2]
x=rep(0:1,5)
data=data.frame(x=x, Z=rnorm(10))
res = flip(Y+matrix(x*2,10,5),~x,~Z,data=data)
res

plot(res)

p2=npv(res,"fisher",subsets=list(c1=c("A","B"),c2=names(Y)))
p2
```

---

cFlip

*cFlip*

---

## Description

It concatenates flip objects warning("More than one test statistic is imputed, only the first perms space will be stored.")

## Usage

```
cFlip(...)
```

**Arguments**

... arguments

**Value**

a flip.object

---

flip

*flip*

---

**Description**

The main function for univariate and multivariate testing under a permutation (and rotation) framework + some utilities.

flip is the main function for permutation (or rotation) test.

It allows for multivariate one sample,  $C \geq 2$  samples and any regression tests. Also the use of covariates (to be fitted in the model but) not under test is allowed.

statTest="t" is the t statistic derived from the correlation among each Xs and each Ys (i.e. a linear model for each couples of Xs and Ys). This is different from the fit of a multiple (multivariate) linear models, since the correlation does not consider the other covariates). The test t is valid only under the assumption that each variable in X is independent of each variable in Y. To get adequate test while adjusting for covariates, use Z (see example below) The test statistic "sum" is the sum of values (or frequencies) of the given sample centered on the expected (i.e. computed on the overall sample). "coeff" is the statistic based on the estimated coefficient of an lm. It produces a test for every possible combination of (columns of) X and Y (p-values can be combined using npc). "cor" is the correlation (i.e. not partial correlation) between each column of X and each of Y. "cor.Spearman" (or "cor.rank") is the analogous for Spearman's rank correlation coefficient.

"ANOVA" is synonyms of "F". Only valid for dependence tests (i.e. non constant X). "Mann-Whitney" is synonyms of "Wilcoxon". "rank" choose among "Wilcoxon" and "Kruskal-Wallis" depending if the samples are two or more (respectively).

The "Wilcoxon" statistic is based on the 'sum of ranks of second sample minus  $n1*(n+1)/2$ ' instead of 'sum of ranks of smallest sample minus  $nSmallest*(n+1)/2$ '. Therefore the statistic is centered on 0 and allow for two sided alternatives. Despite the p-value are ok, it requires the X to be a two-levels factor in order to compute the right test statistic. When the X is not a two-levels factor, it measures the codeviance among X and ranks of Y.

For paired samples (see also the argument Strata and the example below) the Signed Rank test is performed. To perform the Sign Test use option Sign (i.e. same as Signed Rank but without using magnitude of ranks).

The "Fisher" test is allowed only with dichotomous Ys. The reported statistic is the bottom-right cell of the 2 by 2 frequencies table. The "chisq.separated" test perform cell-wise chi squared (see also Finos and Salmaso (2004) Communications in Statistics - Theory and methods).

The "McNemar" test is based on the signs of the differences, hence it can be used also with ordinal or continuous responses. Only valid for symmetry tests (i.e. X is constant or NULL). The reported

statistic for "McNemar" test is the signed squared root of the McNemar statistic. Hence it allows for tailed alternatives.

For ordered X, a stochastic ordering test can be performed using "t", "Wilcoxon", "sum" and then combining the separated test using npc.

When statTest is a function, the first argument must be Y. This same function is ran to observed data Y and to a number of permuted rows of Y. The returned value must be a vector of test statistics. Please note that argument tail must be defined accordingly. The default way the rows of Y are rearranged is through permutation (without strata). More complex permutation strategies can be defined through proper definition of argument perm (see also [permutationSpace](#)).

For testType="rotation": As long as the number of orthogonalized residuals (i.e. the number of observations minus the number of columns in Z) is lower than 50, the function rom is used. The the number is larger, the faster version romFast is used instead. Although the latter is less accurate, for such a big sample size, it is not expected to affect the control of the type I error.

## Usage

```
flip(Y, X = NULL, Z = NULL, data = NULL, tail = 0, perms = 1000,
     statTest = NULL, Strata = NULL, flipReturn, testType = NULL,
     returnGamma = TRUE, ...)
```

## Arguments

Y	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.
X	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.
Z	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 formula. 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.
tail	Vector of values -1, 0 or 1 indicating the tail to be used in the test for each column of Y. tail=1 (-1) means that greater (smaller) values bring more evidence to the alternative hypothesis. tail=0 indicates a two sided alternative. If the length of tail is smaller than number of columns of Y, the values are recycled.
perms	The number of permutations to use. The default is perms = 1000. Alternatively it can be a matrix (i.e. the permutation space) or a list with elements number and seed.
statTest	Choose a test statistic from flip.statTest. See also Details section.

Strata	A vector, which unique values identifies strata. This option is used only with <code>testType="permutation"</code> ; parameter <code>Z</code> is not considered in this case. Also note that when only two levels with one observation per each level are present in each stratum, the problem becomes a paired two-samples problem and hence simplified to a one-sample test.
flipReturn	list of objects indicating what will be included in the output. e.g. <code>list(permP=TRUE, permT=TRUE, data=TRUE)</code> .
testType	by default <code>testType="permutation"</code> . The use of option <code>"combination"</code> is more efficient when <code>X</code> is indicator of groups (i.e. <code>C&gt;1</code> samples testing). When the total number of possible combinations exceeds 10 thousand, <code>"permutation"</code> is performed. As an alternative, if you choose <code>"rotation"</code> , resampling is performed through random linear combinations (i.e. a rotation test is performed). This option is useful when only few permutations are available, that is, minimum reachable significance is high. See also the details section for the algorithm used. The old syntax <code>rotationTest=TRUE</code> is maintained for compatibility but is deprecated, use <code>testType="rotation"</code> instead.
returnGamma	logical. Should be the eigenvectors (with corresponding non-null eigenvalues) of the anti-projection matrix of <code>Z</code> (i.e. $I - Z(Z'Z)^{-1}Z'$ ) returned?
...	Further parameters. The followings are still valid but deprecated: <code>permT.return = TRUE, permP.return = FALSE,</code> <code>permSpace.return = FALSE, permY.return = FALSE.</code> Use <code>flipReturn</code> instead. dummyfy a named list of logical values (eg. <code>list(X=TRUE, Y=TRUE)</code> ) <code>rotationTest= TRUE.</code> Deprecated, use <code>testType='rotation'</code> instead.

### Value

An object of class `flip.object`. Several operations and plots can be made from this object. See also [flip.object-class](#).

### Author(s)

livio finos, Florian Klinglmueller

### References

For the general framework of univariate and multivariate permutation tests see: Pesarin, F. (2001) *Multivariate Permutation Tests with Applications in Biostatistics*. Wiley, New York.

For Rotation tests see: Langsrud, O. (2005) Rotation tests, *Statistics and Computing*, 15, 1, 53-60

A. Solari, L. Finos, J.J. Goeman (2014) Rotation-based multiple testing in the multivariate linear model. *Biometrics*, 70 (4), 954-961.

Livio Finos and Fortunato Pesarin (2018) On zero-inflated permutation testing and some related problems. *Statistical Papers*.

**See Also**

The permutation spaces on which the test is based: [permutationSpace](#) function and useful functions associated with that object.

Multiplicity correction: [flip.adjust](#) and Global test: [npc](#).

**Examples**

```

Y=matrix(rnorm(50),10,5)
colnames(Y)=LETTERS[1:5]
Y[,1:2]=Y[,1:2] +1
res = flip(Y)
res
plot(res[[1]])
plot(res[2:3])
plot(res)

X=rep(0:1,5)
Y=Y+matrix(X*2,10,5)

data=data.frame(Y,X=X, Z=rnorm(10))
#testing dependence among Y's and X
(res = flip(Y,~X,data=data))
#same as:
#res = flip(A+B+C+D+E~X,data=data)

#testing dependence among Y's and X, also using covariates
res = flip(Y,~X,~Z,data=data)
res
#Note that
#flip(Y,X=~X,Z=~1,data=data)
#is different from
#flip(Y,~X,data=data)
#since the former is based on orthogonalized residuals of Y and X by Z.

## Not run:
#Rotation tests:
rot=flip(Y,X,Z=~1,testType="rotation")
# note the use Z=~1.

## End(Not run)

#Using rank tests:
res = flip(Y,~X,data=data,statTest="Wilcoxon")
res

#testing symmetry of Y around 0
Y[,1:2]=Y[,1:2] +2
res = flip(Y)
res
plot(res)

```

```

#use of strata (in this case equal to paired samples)
data$S=rep(1:5,rep(2,5))
#paired t
flip(A+B+C+D+E~X,data=data,statTest="t",Strata=~S)
#signed Rank test
flip(A+B+C+D+E~X,data=data,statTest="Wilcox",Strata=~S)

# tests for categorical data
data=data.frame(X=rep(0:2,10))
data=data.frame(X=factor(data$X),Y=factor(rbinom(30,2,.2+.2*data$X)))
flip(~Y,~X,data=data,statTest="chisq")
# separated chisq (Finos and Salmaso, 2004. Nonparametric multi-focus analysis
# for categorical variables. CommStat - T.M.)
(res.sep=flip(~Y,~X,data=data,statTest="chisq.separated"))
npc(res.sep,"sumT2") #note that combined test statistic is the same as chisq

## Not run:
# User-defined test statistic:
my.fun <- function(Y){
summary(lm(Y~X))$coeff[2,"Estimate"]
}
X <- matrix(rep(0:2,5))
Y <- matrix(rnorm(mean=X,n=15))
res=flip(Y=Y,X=X,statTest=my.fun,tail=0)
res
hist(res)

## End(Not run)

```

---

flip.adjust

*Functions for multiplicity corrections*


---

## Description

npc provides overall tests (i.e. weak FWER control), while flip.adjust provides adjusted p-values (i.e. strong FWER control).

## Usage

```
flip.adjust(permTP, method = flip.npc.methods, maxalpha = 1,
  weights = NULL, stdSpace = FALSE, ...)
```

```
npc(permTP, comb.funct = c(flip.npc.methods, p.adjust.methods),
  subsets = NULL, weights = NULL, stdSpace = FALSE, ...)
```

## Arguments

permTP            A permutation space (B times m matrix) or an flip.object as produced by [flip](#). Alternatively it can be a [flip.object-class](#) resulting, for example from a call of function [flip](#).



method	A method among <code>flip.npc.methods</code> or <code>p.adjust.methods</code> . By default "maxT" is used. See also the section Details.
maxalpha	Adjusted p-values greater than maxalpha are forced to 1. It saves computational time when there are many hypotheses under test.
weights	Optional argument that can be used to give certain variables greater weight in the combined test. Can be a vector or a list of vectors. In the latter case, a separate test will be performed for each weight vector. If both subsets and weights are specified as a list, they must have the same length. In that case, weights vectors may have either the same length as the number of covariates in alternative, or the same length as the corresponding subset vector. Weights can be negative; the sign has no effect unless directional is TRUE. It works for npc and flip.adjust with method="maxT", "maxTstd" or "minP"
stdSpace	Ask if the permutation distribution of the test statistic should be standardized or not. The default is FALSE. The option is applied only if comb.funct or method is equal to "maxT" or "sumT", it becomes useful when test statistics are of different nature (e.g. chisquare and t-test).
...	further arguments. Among them, tail can be used to set the tail of the alternative for the permTP (see also <code>flip</code> ). The arguments statTest, fastSumCombination and linComb are used in objects flipMix and comb.funct="data.sum", "data.linComb", "data.pc" or "data.trace".
comb.funct	A combining function <code>flip.npc.methods</code> (all but "kfwer"): "Fisher", "Liptak", "MahalanobisT", "MahalanobisP" (i.e. related to Hotelling T2), "minP" (i.e. Tippet), "maxT", "sumT" (i.e. direct), "sumT2" (sum of T <sup>2</sup> ). "Fisher" combining function is the default. See also the section Details.
subsets	Optional argument that can be used to test one or more subsets of variables. Can be a vector of column names or indices of a <code>flip.object-class</code> ( <code>names(flipObject)</code> ), or a list of such vectors. In the latter case, a separate test will be performed for each subset. Only for comb.funct %in% c("data.sum", "data.linComb", "data.pc", "data.trace") the names refers to the columns of Y data <code>colnames(flipObject@data\$Y)</code> .

## Details

npc combines the p-values using the combining functions (and the method) described in Pesarin (2001). It makes use of the join space of the permutations. This is usually derived from a call of flip function or flipMixWithin.

Very shortly: "Fisher" = -sum log(p-values) "Liptak" = sum qnorm(p-values) "MahalanobisT" = Mahalanobis distance of centered matrix permTP (or permTP@permT) "MahalanobisP" = same as above, but using scores defined by qnorm(p-values) (tails are forced to be one-sided) "minP" = "Tippet" = min(p-values) \ "maxT" = max(test statistics) "maxTstd" = max(standardized test statistics) "sumT" = sum (test statistics) "sumTstd" = sum (standardized test statistics) "sumT2" = sum (test statistics)<sup>2</sup>. The followings have to be used carefully and only with objects from function flipMix: "data.sum" = sum of all columns of Y, "data.linComb" = sum of all columns of Y (includes a vector or matrix linComb among the arguments), "data.pc" = extracts the first Principal component from the covariance matrix (you may also include a vector whichPCs indicating which PCs you want to consider) \ "data.trace" = Extends the Pillai Trace, use parametric bootstrap to asses the significance. "kfwer" = can be only used with flip.adjust (not in npc). It requires an extra parameter k (k=11 by default).

flip.adjust adjusts the p-value for multiplicity (FamilyWise Error Rate -FWER- and kFWER). When method is equal to "maxT", "maxTstd" (i.e. max T on scale(permTP)) or "minP" (i.e. Tippett) it performs the step-down method of Westfall and Young (1993). For any other element of flip.npc.methods (i.e. "Fisher", "Liptak", "sumT" (i.e. direct) or "sumT2" (sum of T^2)) a call to npc together with a closed testing procedure is used (it make use of [cherry:closed](#)). When method is any among p.adjust.methods the function stats.p.adjust or -if weights are provided- someMTP:p.adjust.w is used. To perform control of the kFWER use flip.adjust with method="kfwer" and extra parameter k.

### Value

The function returns an object of class [flip.object-class](#) (and the use of `getFlip(obj, "Adjust")`).

### Author(s)

livio finos, Florian Klinglmueller and Aldo Solari.

### References

Pesarin (2001) Multivariate Permutation Tests with Applications in Biostatistics. Wiley, New York.  
 P. H. Westfall and S. S. Young (1993). Resampling-based multiple testing: Examples and methods for p-value adjustment. John Wiley & Sons.

### Examples

```
Y=data.frame(matrix(rnorm(50),10,5))
names(Y)=LETTERS[1:5]
Y[,1:2]=Y[,1:2]+1.5
res = flip(Y,perms=10000)

#####npc
p2=npv(res) # same as p2=npv(res,"Fisher")
summary(p2)
p2=npv(res,"minP")
summary(p2)
p2=npv(res,"Fisher",subsets=list(c1=c("A","B"),c2=names(Y)))
summary(p2)
p2=npv(res,"Fisher",subsets=list(c1=c("A","B"),c2=names(Y)),weights=1:5)
summary(p2)

res=flip.adjust(res, method="maxT")

#res=flip.adjust(res,"BH")
##same as
#p.adjust(res,"BH")

## now try
getFlip(res,"Adjust")
```

---

```
flip.object-class      Class flip.object
```

---

## Description

The class `flip.object` is the output of a call to `flip`, `flipMix`, `npc`, `flip.adjust` etc. The following are functions to extract and manipulate relevant information from a `flip.object`.

`show` prints the `flip.object`.

`summary`: Prints information about the `flip-object`.

`dim`: Size of permutation space, i.e. the number of permutations X number of variables.

`[` and `[[`: Extract parts of `flip.object`

`names`: it deals the names of the tests (the variables, usually)

`sort`: it sorts the tests stored in `flip.objects` by their p-values.

`hist`: it produces the histogram of the distribution of the test statistic. When there are more test, it produces one histogram for each test statistic.

`plot`: it produces the scatter plot of the (joint) distribution of the test statistics. When there are more than 2 tests, the plot is based on the first two principal components.

## Usage

```
## S4 method for signature 'flip.object'
show(object)

summary(object, ...)

## S4 method for signature 'flip.object'
summary(object, star.signif = TRUE,
         only.p.leq = NULL, ...)

## S4 method for signature 'flip.object'
dim(x)

## S4 method for signature 'flip.object,ANY,ANY,ANY'
x[i]

## S4 method for signature 'flip.object'
x[[i]]

## S4 method for signature 'flip.object'
length(x)

## S4 method for signature 'flip.object'
names(x)
```

```
## S4 replacement method for signature 'flip.object'
names(x) <- value

## S4 method for signature 'flip.object'
sort(x, decreasing = FALSE)

## S4 method for signature 'flip.object'
hist(x, ...)

## S4 method for signature 'flip.object'
plot(x, y, ...)
```

### Arguments

object	a flip-object
...	additional arguments to be passed
star.signif	If TRUE (default), it puts stars on the significant tests
only.p.leq	Shows only tests with a p-value lower than only.p.leq. The default NULL is equivalent to set only.p.leq=1.
x	a flip.object.
i	indices specifying elements to extract or replace. Indices are numeric, character or logical vectors or empty (missing) or NULL.
value	character vector of up to the same length as x, or NULL.
decreasing	logical. Should the sort be increasing or decreasing?
y	not used.

### Examples

```
Y=matrix(rnorm(50),10,5)
colnames(Y)=LETTERS[1:5]
Y[,1:2]=Y[,1:2] +2
res = flip(Y)
res
summary(res)
sort(res)
names(res)
length(res)
dim(res)
res=res[2:3]
res
hist(res)
plot(res)
```

---

flipMix	<i>The main function for testing mixed models under a permutation (and rotation) framework</i>
---------	--

---

## Description

It allows to test fixed effect in mixed models. You can test within-unit effects, between-unit and interactions of the two. The response can be uni- or multi-variate. See also examples below.

## Usage

```
flipMix(modelWithin, X = NULL, Z = NULL, units, perms = 1000,
        data = NULL, tail = 0, statTest = NULL, flipReturn,
        testType = "permutation", Su = NULL, equal.se = FALSE, se = NA,
        replaceNA.coeffWithin = "coeffMeans",
        replaceNA.coeffWithin.se = replaceNA.coeffWithin, ...)
```

## Arguments

modelWithin	When it is a <a href="#">formula</a> object, a (possibly multivariate) multiple linear model is fitted. Responses are on the left, while the right part contains ONLY within-unit variables. In this case data must be supplied. Alternatively, it can be a glm, a lm or vgam (library(VGAM)) (i.e. vglm) object. The modelWithin have to be performed using only variables within-unit, without using units indicator (in this case the argument data is not used). It can be also a list of models. It can be null if data is provided in the right format (see below).
X	The part of the design matrix corresponding to the between-unit effect that are not null under the alternative hypothesis. If it is a matrix or a data.frame it must have a number of rows equal to the number of units or equal to the total number of observations (in the latest case all elements of the same units must have the same values since they are between-unit effects). The non-null between-unit covariates of null model are defined in Z (see argument below) and do not have to be supplied again here. See also the function <a href="#">flip</a> NOTE: When called from flipMixWithin, W is used only if statTest="TBTWest".
Z	The part of the design matrix corresponding to the non-null between-unit covariates of the model under the null hypothesis. May be given as a design matrix or as a half <a href="#">formula</a> object (e.g. ~a+b). See also the function <a href="#">flip</a> . If it is a matrix or a data.frame it must have a number of rows equal to the number of units or equal to the total number of observations (in the latest case all elements of the same units must have the same values since they are between-unit effects).
units	Vector of units IDs. May be given as a vector or as a half <a href="#">formula</a> object (e.g. ~subj).
perms	The number of permutations to use. The default is perms = 1000. Alternatively it can be a matrix (i.e. the permutation space) or a list with elements number and seed. See also the function <a href="#">flip</a> .

data	Same as in the function <code>flip</code> . It can also be the results of <code>obs2coeffWithin</code> .
tail	Same as in the function <code>flip</code> .
statTest	For function <code>flipMix</code> choose among "t" and "F" (very similar to <code>statTest</code> in function <code>flip</code> ). For function <code>flipMixWithin</code> choose among "Tnaive" (i.e. no estimate of the variance), "TH0est" (Default, i.e. estimate of the variance under H0), "TH1est" (i.e. estimate of the variance under H1 for each permutation. Slower but some time more powerful) and "TBTWest" (i.e. estimate of the variance using ILS algorithm at each permutation; it allows for Z different from a constant term. This is the same algorithm used for <code>flipMix</code> . MUCH slower but some time even more powerful). Both functions allow for vector arguments.
flipReturn	Same as in the function <code>flip</code> .
testType	See also the function <code>flip</code> . Note that this option used only with function <code>flipMix</code> .
Su	Usually NULL. It is the covariance matrix of the random effects. If not supplied, it is estimated by iterative least square algorithm.
equal.se	Logical. If TRUE it force the unit to have the same variance of errors (like it is usually assumed in the lmer methods).
se	Usually NULL. It is a matrix of unit-specific standard errors. If not supplied it is estimated by the algorithm.
replaceNA.coeffWithin	default is NA i.e. no replacement. You can provide a specific value (or a vector of values). You can also choose among "coeffMeans" and "unitMeans" (i.e. mean along columns or along rows of Y).
replaceNA.coeffWithin.se	default is Inf. Use the same options of <code>replaceNA.coeffWithin</code> (but means are over the variances and then rooted).
...	Further parameters. <code>test.coeffWithin</code> Vector of names or IDs of within-unit variables that have to be tested (and reported). Note that variables not in the list are used in the model (i.e. they play the role of nuisance parameters). <code>fastSumCombination,onlyMANOVA</code> and <code>linComb</code> are used in <code>flipMix</code> to deal with combination of variables/coefficients. See also the function <code>flip</code> for other parameters.

## Value

`flipMix` and `flipMixWithin` return an object of class `flip.object`. Several operations and plots can be made from this object. See also [flip.object-class](#).

Note that function `flipMix` with `statTest="t"` or "F" provides tests for each effect between (and interaction) and also provides the overall test PC1 and sum (i.e. all effects are null, same as `npc` does).

Use `npc` with any `comb.funct=c("data.sum", "data.linComb", "data.pc", "data.trace")` to combine results.

`obs2coeffWithin` return a list of objects that can be used as argument of data in the function `flipMix` and `flipMixWithin`.

**Author(s)**

Livio Finos and Dario Basso

**References**

L. Finos and D. Basso (2013) Permutation Tests for Between-Unit Fixed Effects in Multivariate Generalized Linear Mixed Models. *Statistics and Computing*.

D. Basso, L. Finos (2011) Exact Multivariate Permutation Tests for Fixed Effects in Mixed-Models. *Communications in Statistics - Theory and Methods*.

**See Also**

[flip](#), [npc](#)

**Examples**

```

N=10
toyData= data.frame(subj=rep(1:N,rep(4,N)), Within=rep(1:2,N*2),
                    XBetween= rep(1:2,rep(N/2*4,2)),ZBetween= rep(rnorm(N/2),rep(8,N/2)))
toyData= cbind(Y1=rnorm(n=N*4,mean=toyData$subj+toyData$ZBetween+toyData$XBetween),
              Y2=rnorm(n=N*4,mean=toyData$subj+toyData$ZBetween+toyData$Within*2),toyData)
(toyData)

#####
###Testing Between-unit effects
(res=flipMix(modelWithin=as.matrix(toyData[,c("Y1","Y2")])~Within,data=toyData,
        X=~XBetween,Z=~ZBetween,units=~subj,perms=1000,testType="permutation",statTest="t"))
#same as:
modelWithin <- lm(as.matrix(toyData[,c("Y1","Y2")])~Within,data=toyData)
(flipMix(modelWithin=modelWithin,data=toyData, X=~XBetween,Z=~ZBetween,units= ~subj,
        perms=1000,testType="permutation",statTest="t"))

### Note that this is different from:
modelWithin <- list(Y1=lm(Y1~Within,data=toyData),Y2=lm(Y2~Within,data=toyData))
(flipMix(modelWithin=modelWithin,data=toyData, X=~XBetween,Z=~ZBetween,units= ~subj,
        perms=1000,testType="permutation",statTest="t"))

### combining results
(npc(res,"data.pc"))
(npc(res,"data.trace"))
#####
###Testing Within-unit effects
## The resulting test is approximated. The estimate of the variance within units
## takes in account the presence of effects between units.
(flipMix(modelWithin=as.matrix(toyData[,c("Y1","Y2")])~Within,data=toyData,
        units= ~subj, perms=1000,testType="permutation",statTest="t"))

###The resulting tests are exact. If effects between are presents,
## statTest="Tnaive" or "TBTWest" are more suitable:
(res=flipMixWithin(modelWithin=as.matrix(toyData[,c("Y1","Y2")])~Within,data=toyData,
        units= ~subj, perms=1000,statTest=c("TH1est")))
```

npc(res)

---

getFlip

*getFlip*

---

### Description

It returns a given element of a flip.object.

### Usage

getFlip(obj, element)

### Arguments

obj	a flip.object
element	element to the returned

### Value

values of the element requested.

---

permutationSpace

*These functions handle the orbit of permutation/rotation tests (i.e. permutation/rotation space).*

---

### Description

make.permSpace computes the perms x n matrix of ids used for test of dependence. make.signSpace computes the perms x n vector of +1 and -1 used for symmetry test.

### Arguments

IDs	vector of IDs to be permuted. If IDs is a scalar, it is replaced with 1 : IDs.
return.permIDs	logical. If TRUE, the matrix of permuted IDs is stored and returned. Only used with testType="permutaiton"
N	number of elements of the sample. It is also the dimation of the random orthogonal matrix in rom.
Y	a vector of data. It can also be a vector 1:N referring to the IDs of observations.
perms	number of random permutations. If it is a list, it has two elements number (the number of random permutation requested) and seed (the seed to be set when start generating. it is useful for reproducibility) If perms > number of all possible flips, then compute the complete space.



T	the (possibly multivariate) permutation space as returned, for example by <code>flip</code>
obs.only	logical. If TRUE only the p-value for observed test statistic is returned, otherwise the whole space is computed. Defaults: TRUE if T is a flip-object, FALSE otherwise.
tail	Tail of the distribution being significant for H1. See also argument <code>tail</code> in <code>flip</code> . Defaults: 1 if T is NOT a flip-object, it is taken from T otherwise.
testType	See argument <code>testType</code> in <code>flip</code>
Strata	See argument <code>testType</code> in <code>flip</code>
X	A vector of length N with a different value for each group. Only used together with <code>testType="combination"</code> .
...	other parameters

### Details

`rom` computes a Random Orthogonal Matrix of size  $n \times n$  (C-compiled function, very fast). implements the algorithm of Stewart (1980). The function is compiled in C++. NOTE: this option is not available in the newest versions. This is now equivalent to `romFast`

`romFast` computes a Random Orthogonal Matrix of size  $n \times n$  using the `qr.Q` decomposition. `romFast` is faster than `rom` but can be inaccurate (i.e. providing inaccurate type I error control when used in testing), specially for very small  $n$  (i.e. sample size).

`allpermutations` computes all permutations of a vector Y. Is based on the function `permutations` of the library(e1071).

`t2p` computes the (possibly multivariate) space of p-values from the space of test statistic.

### References

- Pesarin (2001) Multivariate Permutation Tests with Applications in Biostatistics. Wiley, New York.
- Stewart, G. W. (1980). The efficient generation of random orthogonal matrices with an application to condition estimators. *SIAM Journal on Numerical Analysis* 17, 403-409.

### See Also

[flip](#)

### Examples

```
#10 random elements of the orbit of a one-sample test
make.signSpace(5, 10)

#All elements of the orbit of a one-sample test (the size of the space is 2^5 < 1000)
make.signSpace(5, 1000)

## Not run:
#A random rotation matrix of size 3
(r=rom(3))
#verify that it is orthogonal:
r%*%t(r)
```

```
## End(Not run)
```

---

seeds

*Seeds data*

---

**Description**

Famous seeds growing data from Pesarin, F. (2001) *Multivariate Permutation Tests with Applications in Biostatistics*. Wiley, New York.

**Format**

the data.frame contains the three columns: grs, x, y

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