# Package 'FDX'

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Type Package

**Title** False Discovery Exceedance Controlling Multiple Testing Procedures

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Description Multiple testing procedures for heterogeneous and discrete tests as described in Döhler and Roquain (2020) <doi:10.1214/20-EJS1771>. The main algorithms of the paper are available as continuous, discrete and weighted versions. They take as input the results of a test procedure from package 'DiscreteTests', or a set of observed p-values and their discrete support under their nulls. A shortcut function to obtain such p-values and supports is also provided, along with wrappers allowing to apply discrete procedures directly to data.

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Depends R (>= 3.00)

**Imports** Rcpp (>= 1.0.12), PoissonBinomial (>= 1.2.0), pracma, DiscreteFDR (>= 2.0.0), checkmate, lifecycle, methods

**Suggests** DiscreteTests (>= 0.2.1)

LinkingTo Rcpp, RcppArmadillo, PoissonBinomial

URL https://github.com/DISOhda/FDX

BugReports https://github.com/DISOhda/FDX/issues

**RoxygenNote** 7.3.2 **NeedsCompilation** yes

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2 FDX-package

# **Contents**

FDX-package		False Discovery Exceedance (FDX) Control for Heterogeneous and Discrete Tests								
Index									60	
	weighted.PB							• • • • • • • • • •	56	
	weighted.LR								53	
	weighted.GR								50	
	summary.FDX								49	
	rejection.path								47	
	print.FDX								46	
	plot.FDX								44	
	NDPB								41	
	NDLR								38	
	NDGR								35	
	hist.FDX								34	
	fast.Discrete								31	
	DPB.DiscreteTestRes								27	
	DPB								24	
	DLR								21	
	discrete.GR discrete.LR								15 18	
	direct.discrete								12	
									9	
	continuous.LR								6	
	continuous.GR								3	
	FDX-package								2	

# Description

This package implements the [HLR], [HGR] and [HPB] procedures for both heterogeneous and discrete tests (see Reference).

#### Details

The functions are reorganized from the reference paper in the following way. discrete.LR() (for Discrete Lehmann-Romano) implements [DLR], discrete.GR() (for Discrete Guo-Romano) implements [DGR] and discrete.PB() (for Discrete Poisson-Binomial) implements [DPB]. DLR() and NDLR() are wrappers for discrete.LR() to access [DLR] and its non-adaptive version directly. Likewise, DGR(), NDGR(), DPB() and NDPB() are wrappers for discrete.GR() and discrete.PB(), respectively. Their main parameters are a vector of raw observed p-values and a list of the same length, whose elements are the discrete supports of the CDFs of the p-values.

In the same fashion, weighted.LR() (for Weighted Lehmann-Romano), weighted.GR() (for Weighted Guo-Romano) and weighted.PB() (for Weighted Poisson-Binomial) implement [wLR], [wGR] and [wGR], respectively. They also possess wrapper functions, namely wLR.AM(), wGR.AM() and

continuous.GR 3

wPB.AM() for arithmetic weighting, and wLR.GM(), wPB.GM() and wPB.GM() for geometric weighting.

The functions fast.Discrete.LR(), fast.Discrete.GR() and fast.Discrete.PB() are wrappers for DiscreteFDR::fisher.pvalues.support() and discrete.LR(), discrete.GR() and discrete.PB(), respectively, which allow to apply discrete procedures directly to a data set of contingency tables.

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#### References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

Lehmann, E. L. & Romano, J. P. (2005). Generalizations of the familywise error rate. *The Annals of Statistics*, *33*(3), pp. 1138-1154. doi:10.1214/009053605000000084

Guo, W. & Romano, J. P. (2007). A generalized Sidak-Holm procedure and control of generalized error rates under independence. *Statistical Applications in Genetics and Molecular Biology*, *6*(1), Art. 3, 35 pp. (electronic). doi:10.2202/15446115.1247

#### See Also

Useful links:

- https://github.com/DISOhda/FDX
- Report bugs at https://github.com/DISOhda/FDX/issues

continuous.GR

Continuous Guo-Romano procedure

#### **Description**

Apply the usual continuous [GR] procedure, with or without computing the critical values, to a set of p-values. A non-adaptive version is available as well.

4 continuous.GR

#### Usage

```
continuous.GR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  adaptive = TRUE,
  critical.values = FALSE,
  select.threshold = 1
)
GR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
NGR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
```

# Arguments

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

#### **Details**

GR and NGR are wrapper functions for continuous. GR. The first one simply passes all its arguments to continuous. GR with adaptive = TRUE and NGR does the same with adaptive = FALSE.

continuous.GR 5

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num.rejected number of rejections. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

 $\label{eq:data_data} \mbox{Data$Data.name} \ \ \mbox{the respective variable name}(s) \mbox{ of the input data}.$ 

#### References

Guo, W. & Romano, J. P. (2007). A generalized Sidak-Holm procedure and control of generalized error rates under independence. *Statistical Applications in Genetics and Molecular Biology*, 6(1), Art. 3, 35 pp. (electronic). doi:10.2202/15446115.1247

## See Also

kernel, FDX-package, continuous.LR(), discrete.LR(), discrete.GR(), discrete.PB(), weighted.LR(), weighted.GR(), weighted.PB()

6 continuous.LR

#### **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 < - rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# GR without critical values; using extracted p-values
GR.fast <- GR(raw.pvalues)</pre>
summary(GR.fast)
# LR with critical values; using test results object
GR.crit <- GR(test.results, critical.values = TRUE)</pre>
summary(GR.crit)
# Non-adaptive GR without critical values; using test results object
NGR.fast <- NGR(test.results)
summary(NGR.fast)
# Non-adaptive GR with critical values; using extracted p-values
NGR.crit <- NGR(raw.pvalues, critical.values = TRUE)
summary(NGR.crit)
```

continuous.LR

Continuous Lehmann-Romano procedure

#### **Description**

Apply the usual (continuous) [LR] procedure, with or without computing the critical values, to a set of p-values. A non-adaptive version is available as well.

# Usage

```
continuous.LR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  adaptive = TRUE,
  critical.values = FALSE,
```

continuous.LR 7

```
select.threshold = 1
)
LR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
NLR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
```

### **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

#### Details

LR and NLR are wrapper functions for continuous.LR. The first one simply passes all its arguments to continuous.LR with adaptive = TRUE and NLR does the same with adaptive = FALSE.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values. Indices indices of rejected p-values. 8 continuous.LR

Num. rejected number of rejections. Adjusted adjusted p-values.

Critical.values

 $critical\ values\ (only\ exists\ if\ computations\ where\ performed\ with\ \verb|critical.values|$ 

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

## References

Lehmann, E. L. & Romano, J. P. (2005). Generalizations of the familywise error rate. *The Annals of Statistics*, *33*(3), pp. 1138-1154. doi:10.1214/009053605000000084

#### See Also

kernel(), FDX, continuous.GR(), discrete.LR(), discrete.GR(), discrete.PB(), weighted.LR(), weighted.GR(), weighted.PB()

## **Examples**

```
X1 <- c(4, 2, 2, 14, 6, 9, 4, 0, 1)

X2 <- c(0, 0, 1, 3, 2, 1, 2, 2, 2)

N1 <- rep(148, 9)

N2 <- rep(132, 9)

Y1 <- N1 - X1

Y2 <- N2 - X2
```

DGR 9

```
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# LR without critical values; using extracted p-values
LR.fast <- LR(raw.pvalues)</pre>
summary(LR.fast)
# LR with critical values; using test results object
LR.crit <- LR(test.results, critical.values = TRUE)</pre>
summary(LR.crit)
# Non-adaptive LR without critical values; using test results object
NLR.fast <- NLR(test.results)
summary(NLR.fast)
# Non-adaptive LR with critical values; using extracted p-values
NLR.crit <- NLR(raw.pvalues, critical.values = TRUE)
summary(NLR.crit)
```

DGR

Wrapper Functions for the Discrete Guo-Romano Procedure

### **Description**

DGR() is a wrapper function of discrete.GR() for computing [DGR]. It simply passes its arguments to discrete.GR() with fixed adaptive = TRUE.

## Usage

```
DGR(test.results, ...)
## Default S3 method:
DGR(
  test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
  ...
)
```

DGR

```
## S3 method for class 'DiscreteTestResults'
DGR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1,
  ...
)
```

## **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

further arguments to be passed to or from other methods. They are ignored here.

pCDF1ist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

## Details

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

DGR 11

Indices indices of rejected *p*-values.

Num.rejected number of rejections.

Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected p-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$pCDFlist list of the *p*-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

 $\label{eq:defData} \mbox{Data$Data.name} \ \ \mbox{the respective variable name}(s) \mbox{ of the input data}.$ 

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
discrete.GR(), NDGR(), discrete.LR(), DLR(), NDLR(), discrete.PB(), DPB(), NDPB()
```

## **Examples**

```
X1 <- c(4, 2, 2, 14, 6, 9, 4, 0, 1)

X2 <- c(0, 0, 1, 3, 2, 1, 2, 2, 2)

N1 <- rep(148, 9)

N2 <- rep(132, 9)

Y1 <- N1 - X1
```

12 direct.discrete

```
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)
df

# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)
raw.pvalues <- test.results$get_pvalues()
pCDFlist <- test.results$get_pvalue_supports()

# DGR without critical values; using extracted p-values and supports
DGR.fast <- DGR(raw.pvalues, pCDFlist)
summary(DGR.fast)

# DGR with critical values; using test results object
DGR.crit <- DGR(test.results, critical.values = TRUE)
summary(DGR.crit)</pre>
```

direct.discrete

Direct Application of Multiple Testing Procedures to Dataset

## **Description**

Apply the [DLR], [NDLR], [DGR], [NDGR], [PB] or [NPB] procedure, with or without computing the critical constants, to a data set of 2x2 contingency tables using a hypothesis test function from package DiscreteTests.

#### Usage

```
direct.discrete.LR(
  dat.
  test.fun,
  test.args = NULL,
  alpha = 0.05,
  zeta = 0.5,
  direction = "su",
  adaptive = FALSE,
  critical.values = FALSE,
  select.threshold = 1,
  preprocess.fun = NULL,
  preprocess.args = NULL
)
direct.discrete.GR(
  dat,
  test.fun,
  test.args = NULL,
```

direct.discrete 13

```
alpha = 0.05,
  zeta = 0.5,
  adaptive = FALSE,
  critical.values = FALSE,
  select.threshold = 1,
  preprocess.fun = NULL,
 preprocess.args = NULL
)
direct.discrete.PB(
  dat,
  test.fun,
  test.args = NULL,
  alpha = 0.05,
  zeta = 0.5,
  adaptive = FALSE,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1,
  preprocess.fun = NULL,
 preprocess.args = NULL
)
```

## **Arguments**

dat	input data; must be suitable for the first parameter of the provided preprocess. fun
	function or, if preprocess. fun = NULL, for the first parameter of the test. fun

function.

test.fun function **from package** DiscreteTests, i.e. one whose name ends with \*\_test\_pv

and which performs hypothesis tests and provides an object with p-values and their support sets; can be specified by a single character string (which is automatically checked for being a suitable function **from that package** and may be

abbreviated) or a single function object.

test.args optional named list with arguments for test.fun; the names of the list fields

must match the test function's parameter names. The first parameter of the test

function (i.e. the data) MUST NOT be included!

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

direction single character string specifying whether to perform the step-up ("su) or step-

down ("sd"; the default) version of the Lehmann-Romano procedure.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

critical.values

single boolean indicating whether critical constants are to be computed.

14 direct.discrete

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

preprocess.fun optional function for pre-processing the input data; its result must be suitable for the first parameter of the test.fun function.

preprocess.args

optional named list with arguments for preprocess. fun; the names of the list fields must match the pre-processing function's parameter names. The first parameter of the test function (i.e. the data) **MUST NOT** be included!

exact

single boolean indicating whether to compute the Poisson-Binomial distribution exactly or by normal approximation.

## **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 < - rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DLR
DLR.sd <- direct.discrete.LR(df, "fisher")
summary(DLR.sd)
# Non-adaptive DLR (step-up variant; adjusted p-values do not exist here!)
NDLR.su <- direct.discrete.LR(df, "fisher", direction = "su", adaptive = FALSE)
summary(NDLR.su)
# DGR
DGR <- direct.discrete.GR(df, "fisher")</pre>
summary(DGR)
# Non-adaptive DGR
NDGR <- direct.discrete.GR(df, "fisher", adaptive = FALSE)
summary(NDGR)
# DPB (normal approximation)
PB.approx <- direct.discrete.PB(df, "fisher", exact = FALSE)
summary(DGR)
```

discrete.GR 15

```
# Non-adaptive DPB
NPB.exact <- direct.discrete.GR(df, "fisher", adaptive = FALSE)
summary(NDGR)</pre>
```

discrete.GR

Discrete Guo-Romano procedure

## **Description**

Apply the [DGR] procedure, with or without computing the critical values, to a set of p-values and their discrete support. A non-adaptive version is available as well.

# Usage

```
discrete.GR(test.results, ...)
## Default S3 method:
discrete.GR(
  test.results,
  pCDFlist,
 alpha = 0.05,
 zeta = 0.5,
  adaptive = TRUE,
 critical.values = FALSE,
 select.threshold = 1,
 pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
discrete.GR(
  test.results,
  alpha = 0.05,
 zeta = 0.5,
  adaptive = TRUE,
 critical.values = FALSE,
 select.threshold = 1,
)
```

## **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults from package DiscreteTests for which a discrete FDR procedure is to be performed.

further arguments to be passed to or from other methods. They are ignored here.

16 discrete.GR

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

#### **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num. rejected number of rejections. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select. threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

discrete.GR 17

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

DatapCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

#### References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
FDX, discrete.LR(), discrete.PB(), continuous.LR(), continuous.GR(), weighted.LR(), weighted.GR(), weighted.PB()
```

# Examples

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 < - rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DGR without critical values; using test results object
DGR.fast <- discrete.GR(test.results)</pre>
summary(DGR.fast)
# DGR with critical values; using extracted p-values and supports
DGR.crit <- discrete.GR(raw.pvalues, pCDFlist, critical.values = TRUE)</pre>
summary(DGR.crit)
```

18 discrete.LR

```
# Non-Adaptive DGR without critical values; using extracted p-values and supports
NDGR.fast <- discrete.GR(raw.pvalues, pCDFlist, adaptive = FALSE)
summary(NDGR.fast)

# Non-Adaptive DGR without critical values; using test results object
NDGR.crit <- discrete.GR(test.results, adaptive = FALSE, critical.values = TRUE)
summary(NDGR.crit)</pre>
```

discrete.LR

Discrete Lehmann-Romano procedure

#### **Description**

Apply the [DLR] procedure, with or without computing the critical values, to a set of p-values and their discrete support. Both step-down and step-up procedures can be computed and non-adaptive versions are available as well.

# Usage

```
discrete.LR(test.results, ...)
## Default S3 method:
discrete.LR(
  test.results.
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  adaptive = TRUE,
  critical.values = FALSE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
discrete.LR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  adaptive = TRUE,
  critical.values = FALSE,
  select.threshold = 1,
)
```

discrete.LR 19

#### **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

... further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

direction single character string specifying whether to perform the step-up ("su) or step-

down ("sd"; the default) version of the Lehmann-Romano procedure.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

#### **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected p-values.

Num.rejected number of rejections.

Adjusted adjusted p-values (only for step-down direction).

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

20 discrete.LR

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

DatapCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

 $X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)$ 

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
FDX, discrete.GR(), discrete.PB(), continuous.LR(), continuous.GR(), weighted.LR(), weighted.GR(), weighted.PB()
```

## **Examples**

```
X2 <- c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 <- rep(148, 9)
N2 <- rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)
df

# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)
raw.pvalues <- test.results$get_pvalues()</pre>
```

DLR 21

```
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DLR without critical values; using results object
DLR.sd.fast <- discrete.LR(test.results)</pre>
summary(DLR.sd.fast)
# DLR with critical values; using extracted p-values and supports
DLR.sd.crit <- discrete.LR(raw.pvalues, pCDFlist, critical.values = TRUE)
summary(DLR.sd.crit)
# DLR (step-up) without critical values; using extracted p-values and supports
DLR.su.fast <- discrete.LR(raw.pvalues, pCDFlist, direction = "su")</pre>
summary(DLR.su.fast)
# DLR (step-up) with critical values; using results object
DLR.su.crit <- discrete.LR(test.results, direction = "su",
                           critical.values = TRUE)
summary(DLR.su.crit)
# Non-adaptive DLR without critical values; using results object
NDLR.sd.fast <- discrete.LR(test.results, adaptive = FALSE)
summary(NDLR.sd.fast)
# Non-adaptive DLR with critical values; using extracted p-values and supports
NDLR.sd.crit <- discrete.LR(raw.pvalues, pCDFlist, adaptive = FALSE,
                            critical.values = TRUE)
summary(NDLR.sd.crit)
# Non-adaptive DLR (step-up) without critical values; using extracted p-values and supports
NDLR.su.fast <- discrete.LR(raw.pvalues, pCDFlist, direction = "su",
                            adaptive = FALSE)
summary(NDLR.su.fast)
# Non-adaptive DLR (step-up) with critical values; using results object
NDLR.su.crit <- discrete.LR(test.results, direction = "su",
                            adaptive = FALSE, critical.values = TRUE)
summary(NDLR.su.crit)
```

DLR

Wrapper Functions for the Discrete Guo-Romano Procedure

#### **Description**

DLR() is a wrapper function of discrete.LR() for computing [DLR]. It simply passes its arguments to discrete.LR() with fixed adaptive = TRUE.

## Usage

```
DLR(test.results, ...)
```

DLR DLR

```
## Default S3 method:
DLR(
  test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  critical.values = FALSE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  critical.values = FALSE,
  select.threshold = 1,
)
```

# Arguments

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

... further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

direction single character string specifying whether to perform the step-up ("su) or step-

down ("sd"; the default) version of the Lehmann-Romano procedure.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

DLR 23

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

#### **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num.rejected number of rejections.

Adjusted adjusted *p*-values (only for step-down direction).

Critical.values

 $critical\ values\ (only\ exists\ if\ computations\ where\ performed\ with\ \verb|critical.values|$ 

= TRUE).

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected *p*-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

DatapCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

24 DPB

#### References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, 14(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
discrete.LR(), NDLR(), discrete.GR(), DGR(), NDGR(), discrete.PB(), DPB(), NDPB()
```

## **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DLR without critical values; using results object
DLR.sd.fast <- DLR(test.results)</pre>
summary(DLR.sd.fast)
# DLR with critical values; using extracted p-values and supports
DLR.sd.crit <- DLR(raw.pvalues, pCDFlist, critical.values = TRUE)</pre>
summary(DLR.sd.crit)
# DLR (step-up) without critical values; using extracted p-values and supports
DLR.su.fast <- DLR(raw.pvalues, pCDFlist, direction = "su")</pre>
summary(DLR.su.fast)
# DLR (step-up) with critical values; using test results object
DLR.su.crit <- DLR(test.results, direction = "su", critical.values = TRUE)
summary(DLR.su.crit)
```

DPB

Wrapper Functions for the Discrete Guo-Romano Procedure

#### **Description**

DPB() is a wrapper function of discrete.PB() for computing [DPB]. It simply passes its arguments to discrete.PB() with fixed adaptive = TRUE.

DPB 25

#### Usage

```
DPB(test.results, ...)
## Default S3 method:
DPB(
   test.results,
   pCDFlist,
   alpha = 0.05,
   zeta = 0.5,
   critical.values = FALSE,
   exact = TRUE,
   select.threshold = 1,
   pCDFlist.indices = NULL,
   ...
)
```

#### **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

... further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

critical.values

single boolean indicating whether critical constants are to be computed.

exact single boolean indicating whether to compute the Poisson-Binomial distribution

exactly or by normal approximation.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if

threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

#### **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have

26 DPB

them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num. rejected number of rejections. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected *p*-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$pCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

#### References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

### See Also

```
discrete.PB(), NDPB(), discrete.GR(), DGR(), NDGR(), discrete.LR(), DLR(), NDLR()
```

#### **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 < - rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DPB (exact) without critical values; using results object
DPB.exact.fast <- DPB(test.results)</pre>
summary(DPB.exact.fast)
# DPB (exact) with critical values; using extracted p-values and supports
DPB.exact.crit <- DPB(raw.pvalues, pCDFlist, critical.values = TRUE)</pre>
summary(DPB.exact.crit)
# DPB (normal approximation) without critical values; using extracted p-values and supports
DPB.norm.fast <- DPB(raw.pvalues, pCDFlist, exact = FALSE)</pre>
summary(DPB.norm.fast)
# DPB (normal approximation) with critical values; using test results object
DPB.norm.crit <- DPB(test.results, critical.values = TRUE, exact = FALSE)</pre>
summary(DPB.norm.crit)
```

DPB.DiscreteTestResults

Discrete Poisson-Binomial procedure

## Description

Apply the [DPB] procedure, with or without computing the critical values, to a set of p-values and their discrete support. A non-adaptive version is available as well. Additionally, the user can choose between exact computation of the Poisson-Binomial distribution or a refined normal approximation.

#### Usage

```
## S3 method for class 'DiscreteTestResults'
DPB(
   test.results,
   alpha = 0.05,
```

```
zeta = 0.5,
 critical.values = FALSE,
 exact = TRUE,
  select.threshold = 1,
)
discrete.PB(test.results, ...)
## Default S3 method:
discrete.PB(
  test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  adaptive = TRUE,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
discrete.PB(
  test.results,
 alpha = 0.05,
 zeta = 0.5,
 adaptive = TRUE,
  critical.values = FALSE,
 exact = TRUE,
  select.threshold = 1,
)
```

#### **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

critical.values

single boolean indicating whether critical constants are to be computed.

exact single boolean indicating whether to compute the Poisson-Binomial distribution

exactly or by normal approximation.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

adaptive single boolean indicating whether to conduct an adaptive procedure or not.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

#### Details

DPB and NDPB are wrapper functions for discrete.PB. The first one simply passes all its arguments to discrete.PB with adaptive = TRUE and NDPB does the same with adaptive = FALSE.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num. rejected number of rejections. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUF)

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected *p*-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

#### References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
kernel, FDX, continuous.LR(), continuous.GR(), discrete.LR(), discrete.GR(), weighted.LR(), weighted.GR(), weighted.PB()
```

#### **Examples**

```
X1 <- c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 < - N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DPB (exact) without critical values; using results object
DPB.exact.fast <- discrete.PB(test.results)</pre>
summary(DPB.exact.fast)
# DPB (exact) with critical values; using extracted p-values and supports
DPB.exact.crit <- discrete.PB(raw.pvalues, pCDFlist, critical.values = TRUE)
summary(DPB.exact.crit)
# DPB (normal approximation) without critical values; using extracted p-values and supports
DPB.norm.fast <- discrete.PB(raw.pvalues, pCDFlist, exact = FALSE)</pre>
summary(DPB.norm.fast)
# DPB (normal approximation) with critical values; using results object
DPB.norm.crit <- discrete.PB(test.results, critical.values = TRUE,</pre>
                               exact = FALSE)
summary(DPB.norm.crit)
```

fast.Discrete 31

fast.Discrete

Fast application of discrete procedures

#### **Description**

#### [Deprecated]

Applies the [DLR], [DGR] or [DPB] procedures, **without** computing the critical values, to a data set of 2 x 2 contingency tables using Fisher's exact test.

Note: These functions are deprecated and will be removed in a future version. Please use direct.discrete.\*()
with test.fun = DiscreteTests::fisher.test.pv and (optional) preprocess.fun = DiscreteDatasets::reconstruct
or preprocess.fun = DiscreteDatasets::reconstruct\_four instead. Alternatively, use a pipeline
like
data |>
DiscreteDatasets::reconstruct\_\*(<args>) |>
DiscreteTests::\*.test.pv(<args>) |>

## Usage

```
fast.Discrete.LR(
  counts,
  alternative = "greater",
  input = "noassoc",
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  adaptive = TRUE
```

discrete.\*(<args>).

32 fast.Discrete

```
)
fast.Discrete.GR(
  counts,
  alternative = "greater",
  input = "noassoc",
  alpha = 0.05,
 zeta = 0.5,
  adaptive = TRUE
)
fast.Discrete.PB(
  counts,
  alternative = "greater",
  input = "noassoc",
  alpha = 0.05,
  zeta = 0.5,
  adaptive = TRUE,
  exact = FALSE
```

## **Arguments**

counts	a data frame of 2 or 4 columns and any number of lines, each line representing a 2 x 2 contingency table to test. The number of columns and what they must contain depend on the value of the input argument, see Details of DiscreteFDR::fisher.pvalues.support
alternative	same argument as in fisher.test(). The three possible values are "greater" (default), "two.sided" or "less"; may be abbreviated.
input	the format of the input data frame, see Details of DiscreteFDR::fisher.pvalues.support(). The three possible values are "noassoc" (default), "marginal" or "HG2011"; may be abbreviated.
alpha	single real number strictly between 0 and 1 specifying the target FDP.
zeta	single real number strictly between 0 and 1 specifying the target probability of not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen equal to alpha.
direction	single character string specifying whether to perform the step-up ("su) or step-down ("sd"; the default) version of the Lehmann-Romano procedure.
adaptive	single boolean indicating whether to conduct an adaptive procedure or not.
exact	single boolean indicating whether to compute the Poisson-Binomial distribution exactly or by normal approximation.

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values. Indices indices of rejected p-values. fast.Discrete 33

Num.rejected number of rejections.

Adjusted adjusted *p*-values (only for step-down direction).

Critical.values

 $critical\ values\ (only\ exists\ if\ computations\ where\ performed\ with\ \verb|critical.values|$ 

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

## **Examples**

```
X1 <- c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 <- c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 <- rep(148, 9)
N2 <- rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)
df

# DLR
DLR.sd <- fast.Discrete.LR(counts = df, input = "noassoc")
summary(DLR.sd)

# DLR
DLR.su <- fast.Discrete.LR(counts = df, input = "noassoc", direction = "su")
summary(DLR.su)</pre>
```

34 hist.FDX

```
# Non-adaptive DLR
NDLR.sd <- fast.Discrete.LR(counts = df, input = "noassoc", adaptive = FALSE)
summary(NDLR.sd)
# Non-adaptive DLR
NDLR.su <- fast.Discrete.LR(counts = df, input = "noassoc", direction = "su", adaptive = FALSE)
summary(NDLR.su)
# DGR
DGR <- fast.Discrete.GR(counts = df, input = "noassoc")</pre>
summary(DGR)
# Non-adaptive DGR
NDGR <- fast.Discrete.GR(counts = df, input = "noassoc", adaptive = FALSE)
summary(NDGR)
# DPB
DPB <- fast.Discrete.PB(counts = df, input = "noassoc")</pre>
summary(DPB)
# Non-adaptive DPB
NDPB <- fast.Discrete.PB(counts = df, input = "noassoc", adaptive = FALSE)
summary(NDPB)
```

hist.FDX

Histogram of Raw P-Values

## **Description**

Computes a histogram of the raw p-values of a FDX object.

## Usage

```
## S3 method for class 'FDX'
hist(x, breaks = "FD", mode = c("raw", "selected", "weighted"), ...)
```

## **Arguments**

X	object of class FDX.
breaks	as in <code>graphics::hist()</code> ; here, the Friedman-Diaconis algorithm ("FD") is used as default.
mode	single character string specifying for which \$p\$-values the histogram is to be generated; must be one of "raw", "selected" or "weighted".
• • •	further arguments to graphics::hist() or graphics::plot.histogram(), respectively.

NDGR 35

#### **Details**

If x does not contain results of a weighting or selection approach, a warning is issued and a histogram of the raw p-values is drawn.

#### Value

An object of class histogram.

# Examples

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DGR
DGR <- DGR(raw.pvalues, pCDFlist)</pre>
# histogram of raw p-values
hist(DGR)
# arithmetic-weighted GR (using 1 - raw.pvalues as weights)
wGR <- wGR.AM(raw.pvalues, 1 - raw.pvalues)
# histogram of raw p-values
hist(wGR)
# histogram of weighted p-values
hist(wGR, mode = "weighted")
```

NDGR

Wrapper Functions for the Non-Adaptive Discrete Guo-Romano Procedure

#### **Description**

NDGR() is a wrapper function of discrete.GR() for computing non-adaptive [DGR]. It simply passes its arguments to discrete.GR() with fixed adaptive = FALSE.

NDGR NDGR

#### Usage

```
NDGR(test.results, ...)
## Default S3 method:
NDGR(
  test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1,
)
```

#### **Arguments**

 $test.results \qquad either a numeric vector with p-values or an R6 object of class {\tt DiscreteTestResults}$ 

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

... further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw

NDGR 37

p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

## **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected p-values.

Num.rejected number of rejections.

Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

SelectScaled scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

DatapCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

NDLR NDLR

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

## See Also

```
discrete.GR(), DGR(), discrete.LR(), DLR(), NDLR(), discrete.PB(), DPB(), NDPB()
```

## **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# Non-adaptive DGR without critical values; using extracted p-values and supports
NDGR.fast <- NDGR(raw.pvalues, pCDFlist)</pre>
summary(NDGR.fast)
# Non-adaptive DGR with critical values; using test results object
NDGR.crit <- NDGR(test.results, critical.values = TRUE)</pre>
summary(NDGR.crit)
```

NDLR

Wrapper Functions for the Non-Adaptive Discrete Guo-Romano Procedure

# Description

NDLR() is a wrapper function of discrete.LR() for computing non-adaptive [DLR]. It simply passes its arguments to discrete.LR() with fixed adaptive = FALSE.

```
NDLR(test.results, ...)
## Default S3 method:
NDLR(
```

NDLR 39

```
test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  critical.values = FALSE,
  select.threshold = 1,
 pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
NDLR(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  direction = "sd",
  critical.values = FALSE,
  select.threshold = 1,
)
```

## **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

... further arguments to be passed to or from other methods. They are ignored here.

pCDFlist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

direction single character string specifying whether to perform the step-up ("su) or step-

down ("sd"; the default) version of the Lehmann-Romano procedure.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

40 NDLR

## **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected p-values.

Num. rejected number of rejections.

Adjusted adjusted p-values (only for step-down direction).

Critical.values

 $critical\ values\ (only\ exists\ if\ computations\ where\ performed\ with\ \verb|critical.values|$ 

= TRUE).

Select list with data related to p-value selection; only exists if select.threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$pCDFlist list of the *p*-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

# References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

NDPB 41

## See Also

```
discrete.GR(), DGR(), discrete.LR(), DLR(), NDLR(), discrete.PB(), DPB(), NDPB()
```

## **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# Non-adaptive DLR without critical values; using results object
NDLR.sd.fast <- NDLR(test.results)</pre>
summary(NDLR.sd.fast)
# Non-adaptive DLR with critical values; using extracted p-values and supports
NDLR.sd.crit <- NDLR(raw.pvalues, pCDFlist, critical.values = TRUE)
summary(NDLR.sd.crit)
# Non-adaptive DLR (step-up) without critical values; using extracted p-values and supports
NDLR.su.fast <- NDLR(raw.pvalues, pCDFlist, direction = "su")</pre>
summary(NDLR.su.fast)
# Non-adaptive DLR (step-up) with critical values; using test results object
NDLR.su.crit <- NDLR(test.results, direction = "su", critical.values = TRUE)
summary(NDLR.su.crit)
```

**NDPB** 

Wrapper Functions for the Non-Adaptive Discrete Guo-Romano Procedure

## **Description**

NDPB() is a wrapper function of discrete.PB() for computing non-adaptive [DPB]. It simply passes its arguments to discrete.PB() with fixed adaptive = FALSE.

42 NDPB

## Usage

```
NDPB(test.results, ...)
## Default S3 method:
NDPB(
  test.results,
  pCDFlist,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1,
  pCDFlist.indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
NDPB(
  test.results,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1,
)
```

## **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

.. further arguments to be passed to or from other methods. They are ignored here.

pCDF1ist list of the supports of the CDFs of the p-values; each list item must be a numeric

vector, which is sorted in increasing order and whose last element equals 1.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

critical.values

single boolean indicating whether critical constants are to be computed.

exact single boolean indicating whether to compute the Poisson-Binomial distribution

exactly or by normal approximation.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and

NDPB 43

the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

pCDFlist.indices

list of numeric vectors containing the test indices that indicate to which raw p-value each **unique** support in pCDFlist belongs; ignored if the lengths of test.results and pCDFlist are equal.

## **Details**

Computing critical constants (critical.values = TRUE) requires considerably more execution time, especially if the number of unique supports is large. We recommend that users should only have them calculated when they need them, e.g. for illustrating the rejection area in a plot or other theoretical reasons.

#### Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected p-values.

Num.rejected number of rejections. Adjusted p-values.

Critical.values

 $critical\ values\ (only\ exists\ if\ computations\ where\ performed\ with\ \verb|critical.values|$ 

= TRUE).

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

p-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled scaled selected p-values.

Select\$Number number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$pCDFlist list of the p-value supports.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Adaptive boolean indicating whether an adaptive procedure was conducted or not.

Data\$Data.name the respective variable name(s) of the input data.

44 plot.FDX

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

## See Also

```
discrete.PB(), DPB(), discrete.GR(), DGR(), NDGR(), discrete.LR(), DLR(), NDLR()
```

# **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# Non-adaptive DPB (exact) without critical values; using results object
NDPB.exact.fast <- NDPB(test.results)</pre>
summary(NDPB.exact.fast)
# Non-adaptive DPB (exact) with critical values; using extracted p-values and supports
NDPB.exact.crit <- NDPB(raw.pvalues, pCDFlist, critical.values = TRUE)</pre>
summary(NDPB.exact.crit)
# Non-adaptive DPB (normal approx.) without critical values; using extracted p-values and supports
NDPB.norm.fast <- NDPB(raw.pvalues, pCDFlist, exact = FALSE)</pre>
summary(NDPB.norm.fast)
# Non-adaptive DPB (normal approx.) with critical values; using test results object
NDPB.norm.crit <- NDPB(test.results, critical.values = TRUE, exact = FALSE)
summary(NDPB.norm.crit)
```

plot.FDX

Plot Method for FDX objects

## **Description**

Plots raw \$p\$-values of a FDX object and highlights rejected and non-rejected \$p\$-values. If present, the critical values are plotted, too.

plot.FDX 45

# Usage

```
## S3 method for class 'FDX'
plot(
    x,
    col = c(2, 4, 1),
    pch = c(20, 20, 17),
    lwd = rep(par()$lwd, 3),
    cex = rep(par()$cex, 3),
    type.crit = "b",
    legend = NULL,
    ...
)
```

# Arguments

X	an object of class "FDX".
col	numeric or character vector of length 3 indicating the colors of the
	<ol> <li>rejected \$p\$-values</li> <li>non-rejected \$p\$-values</li> <li>critical values (if present).</li> </ol>
pch	numeric or character vector of length 3 indicating the point characters of the
	<ol> <li>rejected \$p\$-values</li> <li>non-rejected \$p\$-values</li> <li>critical values (if present and type.crit is a plot type like 'p', 'b' etc.).</li> </ol>
lwd	numeric vector of length 3 indicating the thickness of the points and lines; defaults to current par()\$1wd setting for all components.
cex	numeric vector of length 3 indicating the size of point characters or lines of the
	<ol> <li>rejected p-values</li> <li>accepted p-values</li> <li>critical values (if present).</li> </ol>
	defaults to current par()\$cex setting for all components.
type.crit	single character giving the type of plot desired for the critical values (e.g.: 'p', 'l' etc; see graphics::plot.default()).
legend	if NULL, no legend is plotted; otherwise expecting a character string like "topleft" etc. or a numeric vector of two elements indicating $(x, y)$ coordinates.
	further arguments to graphics::plot.default().

# **Details**

If x contains results of a weighted approach, the Y-axis of the plot is derived from the weighted p-values. Otherwise, it is constituted by the raw ones.

46 print.FDX

# **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 < - rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DLR without critical values; using extracted p-values and supports
DLR.sd.fast <- DLR(raw.pvalues, pCDFlist)</pre>
# plot with default settings
plot(DLR.sd.fast)
# DLR (step-up) with critical values; using test results object
DLR.su.crit <- DLR(test.results, direction = "su", critical.values = TRUE)
# limited plot range
plot(DLR.su.crit, xlim = c(1, 5), ylim = c(0, 0.4))
# DPB without critical values; using test results object
DPB.fast <- DPB(test.results)</pre>
# limited plot range, custom colors, line widths and point symbols, top-left legend
plot(DPB.fast, col = c(2, 4), pch = c(2, 3), lwd = c(2, 2),
     legend = "topleft", xlim = c(1, 5), ylim = c(0, 0.4))
# DGR with critical values; using extracted p-values and supports
DGR.crit <- DGR(raw.pvalues, pCDFlist, critical.values = TRUE)</pre>
# additional customized plot parameters
plot(DGR.crit, col = c(2, 4, 1), pch = c(1, 1, 4), lwd = c(1, 1, 2),
     type.crit = 'o', legend = c(1, 0.4), lty = 1, xlim = c(1, 5),
     ylim = c(0, 0.4), cex = c(3, 3, 2)
```

print.FDX

Printing FDX results

## Description

Prints the results of discrete FDX analysis, stored in a FDX S3 class object.

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

rejection.path 47

## **Arguments**

x object of class FDX.

... further arguments to be passed to or from other methods. They are ignored in this function.

## Value

The respective input object is invisibly returned via invisible(x).

# **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 \leftarrow N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DPB with critical values; using test results object
DPB.crit <- DPB(test.results, critical.values = TRUE)</pre>
# print results
print(DPB.crit)
```

rejection.path

Rejection Path Plot (for FDX objects)

# **Description**

Displays the number of rejections of the raw p-values in a FDX object in dependence of the exceedance probability zeta.

```
rejection.path(
    x,
    xlim = NULL,
    ylim = NULL,
    main = NULL,
    xlab = expression(zeta),
```

48 rejection.path

```
ylab = "Number of Rejections",
verticals = FALSE,
pch = 19,
ref.show = FALSE,
ref.col = "gray",
ref.lty = 2,
ref.lwd = 2,
...
)
```

# **Arguments**

X	object of class "FDX".		
xlim	x axis limits of the plot. If NULL (default), the [0, 1] range is used.		
ylim	the y limits of the plot. If NULL (default), the double of the median of the number of possible rejections is used as upper limit.		
main	main title. If NULL (default), a description string is used.		
xlab, ylab	labels for x and y axis.		
verticals	logical; if TRUE, draw vertical lines at steps.		
pch	jump point character.		
ref.show	logical; if TRUE a vertical reference line is plotted, whose height is the number of rejections of the original Benjamini-Hochberg (BH) procedure.		
ref.col	color of the reference line.		
ref.lty, ref.lwd			
	line type and thickness for the reference line.		
	further arguments to stats::plot.stepfun().		

# Value

Invisibly returns a stepfun object that computes the number of rejections in dependence on the exceedance probability zeta.

# **Examples**

```
X1 <- c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 <- c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 <- rep(148, 9)
N2 <- rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)
df

# Construction of the p-values and their supports with Fisher's exact test library(DiscreteTests) # for Fisher's exact test test.results <- fisher_test_pv(df)
raw.pvalues <- test.results$get_pvalues()</pre>
```

summary.FDX 49

```
pCDFlist <- test.results$get_pvalue_supports()

# DLR without critical values; using extracted p-values and supports
DLR <- DLR(raw.pvalues, pCDFlist)

# plot number of rejections dependent on the exceedance probability zeta
rejection.path(DLR, xlim = c(0, 1), ref.show = TRUE, ref.col = "green", ref.lty = 4)

# None-adaptive DLR without critical values; using test results object
NDLR <- NDLR(test.results)

# add plot for non-adaptive procedure (in red)
rejection.path(NDLR, col = "red", add = TRUE)</pre>
```

summary.FDX

Summarizing Discrete FDX Results

# Description

summary method for class FDX

## Usage

```
## S3 method for class 'FDX'
summary(object, ...)
## S3 method for class 'summary.FDX'
print(x, max = NULL, ...)
```

# Arguments

object of class "FDX".

. . . further arguments passed to or from other methods.

x object of class "summary.FDX".

max numeric or NULL, specifying the maximum number of *rows* of the p-value table

to be printed; if NULL (the default), getOption("max.print") is used.

## **Details**

summary. FDX objects include all data of an FDX class object, but also include an additional table which includes the raw p-values, their indices, the respective critical values (if present), the adjusted p-values (if present) and a logical column to indicate rejection. The table is sorted in ascending order by the raw p-values.

print.summary.FDX simply prints the same output as print.FDX, but also prints the p-value table.

50 weighted.GR

## Value

summary. FDX computes and returns a list that includes all the data of an input FDX, plus

Table

a data.frame, sorted by the raw p-values, that contains the indices, that raw p-values themselves, their respective critical values (if present), their adjusted p-values (if present) and a logical column to indicate rejection.

print.summary.FDX returns that object invisibly.

# **Examples**

```
X1 \leftarrow c(4, 2, 2, 14, 6, 9, 4, 0, 1)
X2 \leftarrow c(0, 0, 1, 3, 2, 1, 2, 2, 2)
N1 \leftarrow rep(148, 9)
N2 < - rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Construction of the p-values and their supports with Fisher's exact test
library(DiscreteTests) # for Fisher's exact test
test.results <- fisher_test_pv(df)</pre>
raw.pvalues <- test.results$get_pvalues()</pre>
pCDFlist <- test.results$get_pvalue_supports()</pre>
# DGR with critical values; using test results object
DGR.crit <- DGR(test.results, critical.values = TRUE)</pre>
# create summary
DGR.crit.summary <- summary(DGR.crit)</pre>
# print summary
print(DGR.crit.summary)
```

weighted.GR

Weighted Guo-Romano Procedure

# **Description**

Apply the weighted [wGR] procedure, with or without computing the critical values, to a set of p-values. Both arithmetic and geometric weighting are available.

```
weighted.GR(
  test.results,
  weights = NULL,
  alpha = 0.05,
  zeta = 0.5,
```

weighted.GR 51

```
weighting.method = c("AM", "GM"),
  critical.values = FALSE,
  select.threshold = 1
)
wGR.AM(
  test.results,
  weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
wGR.GM(
  test.results,
  weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
```

## **Arguments**

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

weights numeric vector that contains the weights for the p-values.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

weighting.method

single character string specifying whether to conduct arithmetic (direction = "AM", the default) or geometric weighting (direction = "GM") of p-values.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

# Details

wGR.AM and wGR.GM are wrapper functions for weighted.GR. The first one simply passes all its arguments to weighted.GR with weighting.method = "AM" and wGR.GM does the same with weighting.method

52 weighted.GR

= "GM".

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected p-values.

Num.rejected number of rejections. Weighted weighted p-values. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$Weights the weights for the raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Weighting character string describing the weighting method.

Data\$Data.name the respective variable name(s) of the input data.

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

weighted.LR 53

## See Also

```
kernel, FDX, continuous.LR(), continuous.GR(), discrete.LR(), discrete.GR(), discrete.PB(), weighted.LR(), weighted.PB()
```

## **Examples**

```
# Construction of the p-values and their supports for weighted methods
raw.pvalues.weighted <- c(0.7389727, 0.1882310, 0.1302457, 0.9513677,
                          0.7592122, 0.0100559, 0.0000027, 0.1651034)
weights <- c(0.7947122, 1.2633867, 2.8097858, 2.2112801,
             2.3878654, 1.2389620, 2.3878654, 0.7947122)
# arithmetic-weighted Guo-Romano procedure without critical values
wGR.AM.fast <- wGR.AM(raw.pvalues.weighted, weights)
summary(wGR.AM.fast)
# arithmetic-weighted Guo-Romano procedure with critical values
wGR.AM.crit <- wGR.AM(raw.pvalues.weighted, weights, critical.values = TRUE)
summary(wGR.AM.crit)
# geometric-weighted Guo-Romano procedure without critical values
wGR.GM.fast <- wGR.GM(raw.pvalues.weighted, weights)
summary(wGR.GM.fast)
# geometric-weighted Guo-Romano procedure with critical values
wGR.GM.crit <- wGR.GM(raw.pvalues.weighted, weights, critical.values = TRUE)
summary(wGR.GM.crit)
```

weighted.LR

Weighted Lehmann-Romano Procedure

## **Description**

Apply the weighted [wLR] procedure, with or without computing the critical values, to a set of p-values. Both arithmetic and geometric weighting are available.

```
weighted.LR(
  test.results,
  weights = NULL,
  alpha = 0.05,
  zeta = 0.5,
  weighting.method = c("AM", "GM"),
  critical.values = FALSE,
  select.threshold = 1
)
```

54 weighted.LR

```
wLR.AM(
  test.results,
 weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
wLR.GM(
  test.results,
 weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  select.threshold = 1
)
```

# Arguments

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

weights numeric vector that contains the weights for the p-values.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

weighting.method

single character string specifying whether to conduct arithmetic (direction =

"AM", the default) or geometric weighting (direction = "GM") of p-values.

critical.values

single boolean indicating whether critical constants are to be computed.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

## **Details**

wLR. AM and wLR. GM are wrapper functions for weighted. LR. The first one simply passes all its arguments to weighted. LR with weighting. method = "AM" and wLR. GM does the same with weighting. method = "GM".

weighted.LR 55

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num.rejected number of rejections. Weighted weighted p-values. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to p-value selection; only exists if select. threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$Weights the weights for the raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Weighting character string describing the weighting method.

Data\$Data.name the respective variable name(s) of the input data.

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

```
kernel, FDX, continuous.LR(), continuous.GR(), discrete.LR(), discrete.GR(), discrete.PB(), weighted.GR(), weighted.PB()
```

## **Examples**

```
# Construction of the p-values and their supports for weighted methods
raw.pvalues.weighted <- c(0.7389727, 0.1882310, 0.1302457, 0.9513677,
                          0.7592122, 0.0100559, 0.0000027, 0.1651034)
weights <- c(0.7947122, 1.2633867, 2.8097858, 2.2112801,
             2.3878654, 1.2389620, 2.3878654, 0.7947122)
# arithmetic-weighted Lehmann-Romano procedure without critical values
wLR.AM.fast <- wLR.AM(raw.pvalues.weighted, weights)
summary(wLR.AM.fast)
# arithmetic-weighted Lehmann-Romano procedure with critical values
wLR.AM.crit <- wLR.AM(raw.pvalues.weighted, weights, critical.values = TRUE)
summary(wLR.AM.crit)
# geometric-weighted Lehmann-Romano procedure without critical values
wLR.GM.fast <- wLR.GM(raw.pvalues.weighted, weights)</pre>
summary(wLR.GM.fast)
# geometric-weighted Lehmann-Romano procedure with critical values
wLR.GM.crit <- wLR.GM(raw.pvalues.weighted, weights, critical.values = TRUE)
summary(wLR.GM.crit)
```

weighted.PB

Weighted Poisson-Binomial Procedure

# Description

Apply the weighted [wPB] procedure, with or without computing the critical values, to a set of p-values. Both arithmetic and geometric weighting are available. Additionally, the user can choose between exact computation of the Poisson-Binomial distribution or a refined normal approximation.

```
weighted.PB(
  test.results,
  weights = NULL,
  alpha = 0.05,
  zeta = 0.5,
  weighting.method = c("AM", "GM"),
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1
)

wPB.AM(
  test.results,
```

```
weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1
)
wPB.GM(
  test.results,
  weights,
  alpha = 0.05,
  zeta = 0.5,
  critical.values = FALSE,
  exact = TRUE,
  select.threshold = 1
)
```

# Arguments

test.results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FDR procedure is to be per-

formed.

weights numeric vector that contains the weights for the p-values.

alpha single real number strictly between 0 and 1 specifying the target FDP.

zeta single real number strictly between 0 and 1 specifying the target probability of

not exceeding the desired FDP. If zeta = NULL (the default), then zeta is chosen

equal to alpha.

weighting.method

single character string specifying whether to conduct arithmetic (direction =

"AM", the default) or geometric weighting (direction = "GM") of p-values.

critical.values

single boolean indicating whether critical constants are to be computed.

exact

single boolean indicating whether to compute the Poisson-Binomial distribution

exactly or by normal approximation.

select.threshold

single real number strictly between 0 and 1 indicating the largest raw p-value to be considered, i.e. only p-values below this threshold are considered and the procedures are adjusted in order to take this selection effect into account; if threshold = 1 (the default), all raw p-values are selected.

# **Details**

wPB.AM and wPB.GM are wrapper functions for weighted.PB. The first one simply passes all its arguments to weighted.PB with weighting.method = "AM" and wPB.GM does the same with weighting.method = "GM".

## Value

A FDX S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected *p*-values.

Num.rejected number of rejections. Weighted weighted p-values. Adjusted adjusted p-values.

Critical.values

critical values (only exists if computations where performed with critical.values

= TRUE).

Select list with data related to *p*-value selection; only exists if select.threshold < 1.

Select\$Threshold

*p*-value selection threshold.

Select\$Effective.Thresholds

results of each p-value CDF evaluated at the selection threshold.

Select\$Pvalues selected p-values that are  $\leq$  selection threshold.

Select\$Indices indices of p-values  $\leq$  selection threshold.

Select\$Scaled selected p-values.

Select\$Number of selected p-values  $\leq$  selection threshold.

Data list with input data.

Data\$Method character string describing the used algorithm, e.g. 'Discrete Lehmann-Romano

procedure (step-up)'.

Data\$Raw.pvalues

all observed raw p-values.

Data\$Weights the weights for the raw p-values.

Data\$FDP.threshold

FDP threshold alpha.

Data\$Exceedance.probability

probability zeta of FDP exceeding alpha; thus, FDP is being controlled at level

alpha with confidence 1 - zeta.

Data\$Weighting character string describing the weighting method.

Data\$Data.name the respective variable name(s) of the input data.

## References

Döhler, S. & Roquain, E. (2020). Controlling False Discovery Exceedance for Heterogeneous Tests. *Electronic Journal of Statistics*, *14*(2), pp. 4244-4272. doi:10.1214/20EJS1771

#### See Also

kernel, FDX, continuous.LR(), continuous.GR(), discrete.LR(), discrete.GR(), discrete.PB(), weighted.LR(), weighted.GR()

## **Examples**

```
# Construction of the p-values and their supports for weighted methods
raw.pvalues.weighted <- c(0.7389727, 0.1882310, 0.1302457, 0.9513677,
                          0.7592122, 0.0100559, 0.0000027, 0.1651034
weights <- c(0.7947122, 1.2633867, 2.8097858, 2.2112801,
             2.3878654, 1.2389620, 2.3878654, 0.7947122)
# arithmetic-weighted Poisson-binomial procedure without critical values
wPB.AM.fast <- wPB.AM(raw.pvalues.weighted, weights)</pre>
summary(wPB.AM.fast)
# arithmetic-weighted Poisson-binomial procedure with critical values
wPB.AM.crit <- wPB.AM(raw.pvalues.weighted, weights, critical.values = TRUE)</pre>
summary(wPB.AM.crit)
# geometric-weighted Poisson-binomial procedure without critical values
wPB.GM.fast <- wPB.GM(raw.pvalues.weighted, weights)</pre>
summary(wPB.GM.fast)
# geometric-weighted Poisson-binomial procedure with critical values
wPB.GM.crit <- wPB.GM(raw.pvalues.weighted, weights, critical.values = TRUE)</pre>
summary(wPB.GM.crit)
```

# **Index**

continuous.GR, 3 continuous.GR(), 8, 17, 20, 30, 53, 55, 58 continuous.LR, 6 continuous.LR(), 5, 17, 20, 30, 53, 55, 58	<pre>GR (continuous.GR), 3 graphics::hist(), 34 graphics::plot.default(), 45 graphics::plot.histogram(), 34</pre>
DGR, 9	hist.FDX, 34
DGR(), 2, 24, 26, 38, 41, 44 direct.discrete, 12 direct.discrete.*(), 31	kernel, 5, 30, 53, 55, 58 kernel(), 8
discrete.GR, 15 discrete.GR(), 2, 3, 5, 8, 9, 11, 20, 24, 26,	LR (continuous.LR), 6
discrete.GR(), 2, 3, 3, 8, 9, 11, 20, 24, 20, 30, 35, 38, 41, 44, 53, 55, 58 discrete.LR, 18 discrete.LR(), 2, 3, 5, 8, 11, 17, 21, 24, 26, 30, 38, 41, 44, 53, 55, 58	NDGR, 35 NDGR(), 2, 11, 24, 26, 44 NDLR, 38
discrete.PB (DPB.DiscreteTestResults),	NDLR(), 2, 11, 24, 26, 38, 41, 44 NDPB, 41
27	NDPB(), 2, 11, 24, 26, 38, 41
discrete.PB(), 2, 3, 5, 8, 11, 17, 20, 24, 26, 38, 41, 44, 53, 55, 58	NGR (continuous.GR), 3 NLR (continuous.LR), 6
DiscreteFDR::fisher.pvalues.support(),	NEW (CONTENTIONS). ENJ, O
3, 32	plot.FDX,44
DiscreteTestResults, 4, 7, 10, 15, 19, 22,	print.FDX, 46
25, 28, 36, 39, 42, 51, 54, 57	print.summary.FDX(summary.FDX),49
DiscreteTests, 4, 7, 10, 12, 13, 15, 19, 22, 25, 28, 36, 39, 42, 51, 54, 57	rejection.path,47
DLR, 21 DLR(), 2, 11, 26, 38, 41, 44 DPB, 24	stats::plot.stepfun(), 48 summary.FDX, 49
DPB(), 2, 11, 24, 38, 41, 44 DPB.DiscreteTestResults, 27	weighted.GR, 50 weighted.GR(), 2, 5, 8, 17, 20, 30, 55, 58 weighted.LR, 53
fast.Discrete, 31	weighted.LR(), 2, 5, 8, 17, 20, 30, 53, 58
fast.Discrete.GR(), $3$	weighted.PB, 56
fast.Discrete.LR(), 3	weighted.PB(), 2, 5, 8, 17, 20, 30, 53, 55
fast.Discrete.PB(), $3$	wGR.AM (weighted.GR), 50
FDX, 8, 17, 20, 30, 53, 55, 58	wGR.AM(), 2
FDX (FDX-package), 2	wGR.GM (weighted.GR), 50
FDX-package, 2	wLR.AM (weighted.LR), 53
fisher.test(), $32$	wLR.AM(), 2

INDEX 61

```
wLR.GM (weighted.LR), 53 wLR.GM(), 3 wPB.AM (weighted.PB), 56 wPB.AM(), 3 wPB.GM (weighted.PB), 56 wPB.GM(), 3
```