Package 'DiscreteFWER'

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Title FWER-Based Multiple Testing Procedures with Adaptation for

Type Package

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Discrete Tests

```
Description Implementations of several multiple testing procedures that control
      the family-wise error rate (FWER) designed specifically for discrete tests.
      Included are discrete adaptations of the Bonferroni, Holm, Hochberg and
      Sidák procedures as described in the papers Döhler (2010) ``Validation of
      credit default probabilities using multiple-testing procedures"
      <doi:10.21314/JRMV.2010.062> and Zhu & Guo (2019) ``Family-Wise Error Rate
      Controlling Procedures for Discrete Data"
      <doi:10.1080/19466315.2019.1654912>. The main procedures of this package
      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 apply discrete procedures directly to data is also
      provided.
License GPL (>= 2)
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Encoding UTF-8
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Suggests DiscreteDatasets, DiscreteTests (>= 0.2.1)
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```

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Contents

	DiscreteFWER-pack	cage .																									2
	DBonferroni																										3
	DHochberg																										6
	DHolm																										9
	direct_discrete_FWI	ER																									12
	discrete_FWER																										14
	DSidak																										18
	hist.DiscreteFWER																										21
	plot.DiscreteFWER																										22
	print.DiscreteFWER																										24
	summary.DiscreteF																										
Index																											27
	reteFWER-package	FWER	-Ba	sed	 lult	inl	e 7	Tes	tin	.0	Pr	oc	ed	'ur	es	w	itk	ı /	\delta	an	tai	tio	n	fo	r	Di.	S-
	parameter parame	crete T			 	Τ.				0	-							_		r				, -		-	-

Description

This package implements adaptions for discrete tests of the Bonferroni, Holm, Hochberg and Šidák procedures for control of the family-wise error rate (FWER).

Details

The main function discrete_FWER() makes all four procedures available to the user. DBonferroni(), DHolm(), DHochberg() and DSidak() are wrapper functions that enable the user to access them directly. Their main parameters are either a DiscreteTestResults object from package DiscreteTests or a vector of raw observed p-values and a list whose elements are the discrete supports of the CDFs of the p-values.

The function direct_discrete_FWER() is a wrapper for DiscreteFDR::generate.pvalues() and discrete_FWER(), which applies discrete procedures directly to data.

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DBonferroni 3

References

Döhler, S. (2010). Validation of credit default probabilities using multiple-testing procedures. *Journal of Risk Model Validation*, *4*(4), 59-92. doi:10.21314/JRMV.2010.062

Zhu, Y., & Guo, W. (2019). Family-Wise Error Rate Controlling Procedures for Discrete Data. *Statistics in Biopharmaceutical Research*, 12(1), 117-128. doi:10.1080/19466315.2019.1654912

See Also

Useful links:

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

DBonferroni

Discrete Bonferroni Procedure

Description

DBonferroni() is a wrapper function of discrete_FWER() for computing the discrete Bonferroni procedure for tests with an arbitrary dependency structure. It simply passes its arguments to discrete_FWER() with fixed independence = FALSE and single_step = TRUE.

Usage

```
DBonferroni(test_results, ...)
## Default S3 method:
DBonferroni(
  test_results,
  pCDFlist,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
  pCDFlist_indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
DBonferroni(
  test_results,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
)
```

4 DBonferroni

Arguments

test_results either a numeric vector with p-values or an R6 object of class DiscreteTestResults

from package DiscreteTests for which a discrete FWER 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 indicating the target FWER level.

critical_values

single boolean specifying 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

 $select_threshold = 1$ (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(s) each support in pCDFlist belongs; if NULL (the default) the lengths of test_results and pCDFlist **must** be 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 set in a plot or other theoretical reasons. Setting (critical_values = FALSE) is sufficient for obtaining rejection decisions and adjusted *p*-values.

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections. Adjusted adjusted p-values.

Critical_constants

critical values (only exists if computations where performed with critical_values

= TRUE).

Data list with input data.

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed p-values.

DatapCDFlist list of the p-value supports.

DBonferroni 5

```
Data$FWER_level
```

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

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

Select\$Threshold

p-value selection threshold (select_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.

References

Döhler, S. (2010). Validation of credit default probabilities using multiple-testing procedures. *Journal of Risk Model Validation*, *4*(4), 59-92. doi:10.21314/JRMV.2010.062

Zhu, Y., & Guo, W. (2019). Family-Wise Error Rate Controlling Procedures for Discrete Data. *Statistics in Biopharmaceutical Research*, 12(1), 117-128. doi:10.1080/19466315.2019.1654912

See Also

```
discrete_FWER(), DHolm(), DSidak(), DHochberg()
```

```
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

# Computation of 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()

# d-Bonferroni without critical values; using extracted p-values and supports
DBonferroni_fast <- DBonferroni(raw_pvalues, pCDFlist)
summary(DBonferroni_fast)</pre>
```

6 DHochberg

```
# d-Bonferroni with critical values; using test results object
DBonferroni_crit <- DBonferroni(test_results, critical_values = TRUE)
summary(DBonferroni_crit)</pre>
```

DHochberg

Discrete Hochberg Procedure

Description

DHochberg() is a wrapper function of discrete_FWER() for computing the discrete Hochberg stepup procedure for independent or positively correlated discrete tests. It simply passes its arguments to discrete_FWER() with fixed independence = TRUE and single_step = FALSE.

Usage

```
DHochberg(test_results, ...)
## Default S3 method:
DHochberg(
  test_results,
  pCDFlist,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
  pCDFlist_indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
DHochberg(
  test_results,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
)
```

Arguments

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

... 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.

DHochberg 7

alpha single real number strictly between 0 and 1 indicating the target FWER level.

critical_values

single boolean specifying 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 select_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(s) each support in pCDFlist belongs; if NULL (the default) the lengths of test_results and pCDFlist **must** be 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 set in a plot or other theoretical reasons. Setting (critical_values = FALSE) is sufficient for obtaining rejection decisions and adjusted p-values.

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections. Adjusted adjusted p-values.

Critical_constants

critical values (only exists if computations where performed with critical_values

= TRUE).

Data list with input data.

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed *p*-values.

DatapCDFlist list of the p-value supports.

Data\$FWER_level

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

8 DHochberg

References

Zhu, Y., & Guo, W. (2019). Family-Wise Error Rate Controlling Procedures for Discrete Data. *Statistics in Biopharmaceutical Research*, *12*(1), 117-128. doi:10.1080/19466315.2019.1654912

See Also

```
discrete_FWER(), DSidak(), DBonferroni(), DHolm
```

```
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
# Computation of 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>
# d-Hochberg without critical values; using test results object
DHoch_fast <- DHochberg(test_results)</pre>
summary(DHoch_fast)
# d-Hochberg with critical values; using extracted p-values and supports
DHoch_crit <- DHochberg(raw_pvalues, pCDFlist, critical_values = TRUE)</pre>
summary(DHoch_crit)
```

DHolm 9

DHolm

Discrete Holm Procedure

Description

DHolm() is a wrapper function of discrete_FWER() for computing the discrete Holm step-down procedure for tests with an arbitrary dependency structure. It simply passes its arguments to discrete_FWER() with fixed independence = FALSE and single_step = FALSE.

Usage

```
DHolm(test_results, ...)
## Default S3 method:
DHolm(
  test_results,
  pCDFlist,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
  pCDFlist_indices = NULL,
)
## S3 method for class 'DiscreteTestResults'
DHolm(
  test_results,
  alpha = 0.05,
  critical_values = FALSE,
  select_threshold = 1,
)
```

Arguments

test_results	either a numeric vector with p -values or an R6 object of class <code>DiscreteTestResults</code> from package <code>DiscreteTests</code> for which a discrete FWER procedure is to be performed.
	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 indicating the target FWER level.
critical_values	3
	single boolean specifying whether critical constants are to be computed.

DHolm

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 select_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(s) each support in pCDFlist belongs; if NULL (the default) the lengths of test_results and pCDFlist **must** be 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 set in a plot or other theoretical reasons. Setting (critical_values = FALSE) is sufficient for obtaining rejection decisions and adjusted *p*-values.

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections. Adjusted adjusted p-values.

Critical_constants

critical values (only exists if computations where performed with critical_values

= TRUE).

Data list with input data.

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed p-values.

 ${\tt Data\$pCDFlist} \quad {\tt list of the } \textit{p-value supports}.$

Data\$FWER_level

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

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

Select\$Threshold

p-value selection threshold (select_threshold).

DHolm 11

```
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.
```

References

Döhler, S. (2010). Validation of credit default probabilities using multiple-testing procedures. *Journal of Risk Model Validation*, *4*(4), 59-92. doi:10.21314/JRMV.2010.062

Zhu, Y., & Guo, W. (2019). Family-Wise Error Rate Controlling Procedures for Discrete Data. *Statistics in Biopharmaceutical Research*, 12(1), 117-128. doi:10.1080/19466315.2019.1654912

See Also

```
discrete_FWER(), DBonferroni(), DSidak(), DHochberg()
```

```
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>
# Computation of 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>
# d-Holm without critical values; using extracted p-values and supports
DHolm_fast <- DHolm(raw_pvalues, pCDFlist)</pre>
summary(DHolm_fast)
# d-Holm with critical values; using test results object
DHolm_crit <- DHolm(test_results, critical_values = TRUE)</pre>
summary(DHolm_crit)
```

Description

Apply one of the various FWER adaptation procedures, with or without computing the critical constants, to a data set of 2x2 contingency tables using statistical test functions from package DiscreteTests. If necessary, functions for pre-processing can be passed as well.

Usage

```
direct_discrete_FWER(
  dat,
  test_fun,
  test_args = NULL,
  alpha = 0.05,
  independence = FALSE,
  single_step = TRUE,
  critical_values = FALSE,
  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 is 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 indicating the target FWER level.				
independence	single boolean specifying whether the p -values are statistically independent or not.				
single_step	single boolean specifying whether to perform a single-step (TRUE) or step-down (FALSE; the default) procedure.				
critical_values					
	single boolean specifying 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 select_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!

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections. adjusted p-values. Adjusted

Critical_constants

critical values (only exists if computations where performed with critical_values

= TRUE).

list with input data. Data

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed p-values.

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

Data\$FWER_level

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

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

Select\$Threshold

p-value selection threshold (select_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.

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
# Computation of 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>
DBonf <- direct_discrete_FWER(df, "fisher")</pre>
summary(DBonf)
DHolm <- direct_discrete_FWER(df, "fisher_test_pv", single_step = FALSE)</pre>
summary(DHolm)
DBonf_bin <- direct_discrete_FWER(X1 + X2, "binom_test_pv",
                                     list(n = N1 + N2, p = 0.05))
summary(DBonf_bin)
DHolm_bin <- direct_discrete_FWER(X1 + X2, "binom",</pre>
                                     list(n = N1 + N2, p = 0.05),
                                     single_step = TRUE)
summary(DHolm_bin)
```

discrete_FWER

Discrete Family-wise Error Rate (FWER) Adaptation Procedures

Description

Apply a discrete FWER adaptation procedure, with or without computing the critical values, to a set of p-values and their discrete support.

Usage

```
discrete_FWER(test_results, ...)
## Default S3 method:
discrete_FWER(
  test_results,
  pCDFlist,
  alpha = 0.05,
```

```
independence = FALSE,
single_step = FALSE,
critical_values = FALSE,
select_threshold = 1,
pCDFlist_indices = NULL,
...
)

## S3 method for class 'DiscreteTestResults'
discrete_FWER(
  test_results,
   alpha = 0.05,
   independence = FALSE,
   single_step = FALSE,
   critical_values = FALSE,
   select_threshold = 1,
...
)
```

Arguments

 ${\tt test_results} \qquad {\tt either \, a \, numeric \, vector \, with \, p\text{-}values \, or \, an \, R6 \, object \, of \, class \, {\tt DiscreteTestResults}}$

from package DiscreteTests for which a discrete FWER 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 indicating the target FWER level.

independence single boolean specifying whether the p-values are statistically independent or

not.

single_step single boolean specifying whether to perform a single-step (TRUE) or step-down

(FALSE; the default) procedure.

critical_values

single boolean specifying 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

select_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(s) each support in pCDFlist belongs; if NULL (the default) the lengths of test_results and pCDFlist **must** be 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 set in a plot or other theoretical reasons. Setting (critical_values = FALSE) is sufficient for obtaining rejection decisions and adjusted *p*-values.

Depending on the choices of independence and single_step, one of the following procedures, is applied:

single-step stepwise
independent Šidák Hochberg (step-up)
not independent Bonferroni Holm (step-down)

Each procedure is available by its own shortcut function:

single-step stepwise independent DSidak() DHochberg() not independent DBonferroni() DHolm()

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections.

Adjusted adjusted p-values.

Critical_constants

critical values (only exists if computations where performed with critical_values

= TRUE).

Data list with input data.

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed p-values.

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

Data\$FWER_level

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

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

Select\$Threshold

p-value selection threshold (select_threshold).

```
\label{eq:continuous} Select$Effective\_Thresholds \\ results of each $p$-value CDF evaluated at the selection threshold. \\ Select$Pvalues \\ selected $p$-values that are $\le$ selection threshold. \\ Select$Indices \\ indices of $p$-values $\le$ selection threshold. \\ Select$Scaled \\ scaled selected $p$-values. \\ Select$Number \\ number of selected $p$-values $\le$ selection threshold. \\ }
```

References

Döhler, S. (2010). Validation of credit default probabilities using multiple-testing procedures. *Journal of Risk Model Validation*, *4*(4), 59-92. doi:10.21314/JRMV.2010.062

Zhu, Y., & Guo, W. (2019). Family-Wise Error Rate Controlling Procedures for Discrete Data. *Statistics in Biopharmaceutical Research*, 12(1), 117-128. doi:10.1080/19466315.2019.1654912

See Also

DiscreteFWER, DBonferroni(), DHolm(), DSidak(), DHochberg()

```
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)
# Computation of 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>
# d-Holm without critical values; using test results object
DFWER_dep_sd_fast <- discrete_FWER(test_results)</pre>
summary(DFWER_dep_sd_fast)
# d-Holm with critical values; using extracted p-values and supports
DFWER_dep_sd_crit <- discrete_FWER(raw_pvalues, pCDFlist,</pre>
                                     critical_values = TRUE)
summary(DFWER_dep_sd_crit)
# d-Bonferroni without critical values; using test results object
DFWER_dep_fast <- discrete_FWER(test_results, single_step = TRUE)</pre>
summary(DFWER_dep_fast)
# d-Bonferroni with critical values; using extracted p-values and supports
DFWER_dep_crit <- discrete_FWER(raw_pvalues, pCDFlist, single_step = TRUE,</pre>
```

18 DSidak

```
critical_values = TRUE)
summary(DFWER_dep_crit)
# d-Hochberg without critical values; using test results object
DFWER_ind_su_fast <- discrete_FWER(test_results, independence = TRUE)</pre>
summary(DFWER_ind_su_fast)
# d-Hochberg with critical values; using extracted p-values and supports
DFWER_ind_su_crit <- discrete_FWER(raw_pvalues, pCDFlist,</pre>
                                    independence = TRUE,
                                    critical_values = TRUE)
summary(DFWER_ind_su_crit)
# d-Šidák without critical values; using extracted p-values and supports
DFWER_ind_fast <- discrete_FWER(raw_pvalues, pCDFlist,</pre>
                                 independence = TRUE,
                                 single_step = TRUE)
summary(DFWER_ind_fast)
# d-Šidák with critical values; using test results object
DFWER_ind_crit <- discrete_FWER(test_results, independence = TRUE,</pre>
                                 single_step = TRUE,
                                 critical_values = TRUE)
summary(DFWER_ind_crit)
```

DSidak

Discrete Šidák Procedure for Independent Tests

Description

DSidak() is a wrapper function of discrete_FWER() for computing the discrete Sidák procedure for independent discrete tests. It simply passes its arguments to discrete_FWER() with fixed independence = TRUE and single_step = TRUE.

Usage

```
DSidak(test_results, ...)
## Default S3 method:
DSidak(
   test_results,
   pCDFlist,
   alpha = 0.05,
   critical_values = FALSE,
   select_threshold = 1,
   pCDFlist_indices = NULL,
   ...
)
```

DSidak 19

```
## S3 method for class 'DiscreteTestResults'
DSidak(
  test_results,
  alpha = 0.05,
  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 FWER 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 indicating the target FWER level.

critical_values

single boolean specifying 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 select_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(s) each support in pCDFlist belongs; if NULL (the default) the lengths of test_results and pCDFlist **must** be 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 set in a plot or other theoretical reasons. Setting (critical_values = FALSE) is sufficient for obtaining rejection decisions and adjusted *p*-values.

Value

A DiscreteFWER S3 class object whose elements are:

Rejected rejected raw p-values.

Indices indices of rejected hypotheses.

Num_rejected number of rejections. Adjusted adjusted p-values.

20 DSidak

Critical_constants

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

= TRUE).

Data list with input data.

Data\$Method character string describing the performed algorithm, e.g. 'Discrete Bonferroni

procedure'.

Data\$Raw_pvalues

observed p-values.

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

Data\$FWER_level

FWER level alpha.

Data\$Independence

boolean indicating whether the p-values were considered as independent.

Data\$Single_step

boolean indicating whether a single-step or step-down procedure was performed.

Data\$Data_name the respective variable names of the input data.

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

Select\$Threshold

p-value selection threshold (select_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.

References

Döhler, S. (2010). Validation of credit default probabilities using multiple-testing procedures. *Journal of Risk Model Validation*, *4*(4), 59-92. doi:10.21314/JRMV.2010.062

See Also

```
discrete_FWER(), DHochberg(), DBonferroni(), DHolm()
```

```
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
```

hist.DiscreteFWER 21

```
# Computation of 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()

# d-Šidák without critical values; using extracted p-values and supports
DSidak_fast <- DSidak(raw_pvalues, pCDFlist)
summary(DSidak_fast)

# d-Šidák with critical values; using test results object
DSidak_crit <- DSidak(test_results, critical_values = TRUE)
summary(DSidak_crit)</pre>
```

hist.DiscreteFWER

Histogram of Raw P-Values

Description

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

Usage

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

Arguments

x	an object of class DiscreteFWER.
breaks	as in graphics::hist(); 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 either be "raw" or "selected".
	<pre>further arguments to graphics::hist() or graphics::plot.histogram(), respectively.</pre>

Details

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

Value

An object of class histogram.

22 plot.DiscreteFWER

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
# Computation of 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>
# d-Holm with critical values; using test results object
DHolm_crit <- DHolm(test_results, critical_values = TRUE)</pre>
hist(DHolm_crit)
```

plot.DiscreteFWER

Plot Method for DiscreteFWER objects

Description

Plots raw p-values of a DiscreteFWER object and highlights rejected and accepted p-values. If calculated, the critical values are plotted, too.

Usage

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

Arguments

x object of class DiscreteFWER.

numeric or character vector of length 3 indicating the colours of the

1. rejected p-values

plot.DiscreteFWER 23

```
2. accepted p-values
                     3. critical values (if present).
pch
                   numeric or character vector of length 3 indicating the point characters of the
                     1. rejected p-values
                     2. accepted p-values
                     3. critical values (if present and type_crit is a plot type like 'p', 'b' etc.).
                   numeric vector of length 3 indicating the thickness of the points and lines; de-
lwd
                   faults to current par()$1wd setting.
cex
                   numeric vector of length 3 indicating the size of point characters or lines of the
                     1. rejected p-values
                     2. accepted p-values
                     3. critical values (if present).
                   defaults to current par()$cex setting.
                    1-character string giving the type of plot desired for the critical values (e.g.: 'p',
type_crit
                    'l' etc; see plot()).
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.
```

Value

A plot is created, but no value is returned.

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
# Computation of 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>
DBonf_fast <- DBonferroni(raw_pvalues, pCDFlist)</pre>
DBonf_crit <- DBonferroni(test_results, critical_values = TRUE)</pre>
DHolm_fast <- DHolm(test_results)</pre>
DHolm_crit <- DHolm(raw_pvalues, pCDFlist, critical_values = TRUE)</pre>
plot(DBonf_fast)
plot(DBonf_crit, xlim = c(1, 5), ylim = c(0, 0.4))
```

further arguments to plot.default().

24 print.DiscreteFWER

print.DiscreteFWER

Printing discrete FWER results

Description

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

Usage

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

Arguments

x object of class DiscreteFWER.

... 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).

```
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 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
# Computation of 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>
# d-Holm with critical values; using test results object
DHolm_crit <- DHolm(test_results, critical.values = TRUE)</pre>
# print results
print(DHolm_crit)
```

summary.DiscreteFWER Summarizing Discrete FWER Results

Description

summary method for class DiscreteFWER.

Usage

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

Arguments

object an object of class DiscreteFWER.

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

x an object of class summary.DiscreteFWER.

max numeric or NULL, specifying the maximal number of rows of the p-value table to

be printed. By default, when NULL, getOption("max.print") is used.

Details

summary.DiscreteFWER objects contain all data of an DiscreteFWER 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.DiscreteFWER simply prints the same output as print.DiscreteFWER, but also prints the p-value table.

Value

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

Table data.frame, sorted by the raw p-values, that contains the indices, the raw p-

values themselves, their respective critical values (if present), their adjusted p-

values (if present) and a logical column to indicate rejection.

```
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 \leftarrow rep(132, 9)
Y1 <- N1 - X1
Y2 <- N2 - X2
df <- data.frame(X1, Y1, X2, Y2)</pre>
df
# Computation of 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>
# d-Holm procedure without critical values; using test results object
DFWER_dep_sd_fast <- discrete_FWER(test_results)</pre>
summary(DFWER_dep_sd_fast)
# d-Bonferroni procedure with critical values; using test results object
DFWER_dep_crit <- discrete_FWER(test_results, single_step = TRUE,</pre>
critical_values = TRUE)
summary(DFWER_dep_crit)
```

Index

```
DBonferroni, 3
DBonferroni(), 2, 8, 11, 17, 20
DHochberg, 6
DHochberg(), 2, 5, 11, 17, 20
DHolm, 8, 9
DHolm(), 2, 5, 17, 20
direct_discrete_FWER, 12
direct_discrete_FWER(), 2
discrete_FWER, 14
discrete_FWER(), 2, 3, 5, 6, 8, 9, 11, 18, 20
DiscreteFDR::generate.pvalues(), 2
DiscreteFWER, 17
DiscreteFWER (DiscreteFWER-package), 2
DiscreteFWER-package, 2
DiscreteTestResults, 2, 4, 6, 9, 15, 19
DiscreteTests, 2, 4, 6, 9, 12, 15, 19
DSidak, 18
DSidak(), 2, 5, 8, 11, 17
graphics::hist(), 21
graphics::plot.histogram(), 21
hist.DiscreteFWER, 21
plot(), 23
plot.default(), 23
plot.DiscreteFWER, 22
print.DiscreteFWER, 24
print.summary.DiscreteFWER
        (summary.DiscreteFWER), 25
summary.DiscreteFWER, 25
```