

# Package ‘MHTdiscrete’

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**Type** Package

**Title** Multiple Hypotheses Testing for Discrete Data

**Version** 1.0.0

**Suggests** multcomp, FixSeqMTP

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**BugReports** <https://github.com/allenzhuaz/MHTdiscrete/issues>

**URL** <https://allen.shinyapps.io/MTPs/>

**Description** A comprehensive tool for almost all existing multiple testing methods for discrete data. The package also provides some novel multiple testing procedures controlling FWER/FDR for discrete data. Given discrete p-values and their domains, the [method].p.adjust function returns adjusted p-values, which can be used to compare with the nominal significant level alpha and make decisions. For users' convenience, the functions also provide the output option for printing decision rules.

**License** GPL (>= 2)

**LazyData** TRUE

**NeedsCompilation** yes

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getPval	<i>Calculating p-values for discrete data</i>
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**Description**

The function for calculating the original available p-values and all attainable p-values for the corresponding hypothesis.

**Usage**

```
getPval(raw.data, test.type, alternative)
```

**Arguments**

<code>raw.data</code>	original data set with count number for treatment group and study group. The data set type could be <a href="#">matrix</a> or <a href="#">data.frame</a> .
<code>test.type</code>	there are two discrete test available now, must be one of "FET" for Fisher's Exact Test and "BET" for Binomial Exact Test.
<code>alternative</code>	indicates the alternative hypothesis and must be one of "two.sided", "greater" or "less".

**Value**

A numeric vector of the adjusted p-values (of the same length as *p*).

**Author(s)**

Yalin Zhu

**References**

Zhu, Y., & Guo, W. (2017). Familywise error rate controlling procedures for discrete data *arXiv preprint* arXiv:1711.08147.

Clopper, C. J. & Pearson, E. S. (1934). The use of confidence or fiducial limits illustrated in the case of the binomial. *Biometrika*, **26**: 404-413.

Fisher, R. A. (1922). On the Interpretation of  $\chi^2$  from Contingency Tables, and the Calculation of P. *Journal of the Royal Statistical Society*, **85**: 87-94.

**Examples**

```
## Using Fisher's Exact Test to get the available and attainable p-values
# import raw data set as data.frame type
df <- data.frame(X1=c(4, 2, 2, 13, 6, 8, 4, 0, 1), N1 = rep(148, 9),
  X2 = c(0, 0, 1, 3, 2, 1, 2, 2, 2), N2 = rep(132, 9))
# obtain the available p-values and attainable p-values using two-sided Fisher's Exact Test
getPval(raw.data=df, test.type = "FET", alternative = "two.sided")
# store the available p-values
p <- getPval(raw.data=df, test.type = "FET", alternative = "two.sided")[[1]]; p
# store the attainable p-values
p.set <- getPval(raw.data=df, test.type = "FET", alternative = "two.sided")[[2]]; p.set
```

---

GTBH.p.adjust	<i>The adjusted p-values for Gilbert-Tarone-BH step-up FDR controlling procedure.</i>
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---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
GTBH.p.adjust(p, p.set, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p).

## Author(s)

Yalin Zhu

## References

Gilbert, P. B. (2005). A modified false discovery rate multiple-comparisons procedure for discrete data, applied to human immunodeficiency virus genetics. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **54**: 143-158.

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society Series B*, **57**: 289-300.

## See Also

[GTBY.p.adjust](#), [MBH.p.adjust](#), [MBY.p.adjust](#)

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
GTBH.p.adjust(p,p.set)
```

---

GTBY.p.adjust	<i>The adjusted p-values for Gilbert-Tarone-BY step-up FDR controlling procedure.</i>
---------------	---

---

### Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

### Usage

```
GTBY.p.adjust(p, p.set, alpha, make.decision)
```

### Arguments

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

### Value

A numeric vector of the adjusted p-values (of the same length as p).

### Author(s)

Yalin Zhu

### References

Gilbert, P. B. (2005). A modified false discovery rate multiple-comparisons procedure for discrete data, applied to human immunodeficiency virus genetics. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **54**: 143-158.

Benjamini, Y., and Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. *Annals of Statistics*, **29**: 1165-1188.

### See Also

[GTBH.p.adjust](#), [MBH.p.adjust](#), [MBY.p.adjust](#)

### Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
GTBY.p.adjust(p,p.set)
```

---

MBH.p.adjust	<i>The adjusted p-values for Modified Benjamini-Hochberg (BH) step-up FDR controlling procedure.</i>
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---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
MBH.p.adjust(p, p.set, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with NAs). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p).

## Author(s)

Yalin Zhu

## References

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society Series B*, **57**: 289-300.

## See Also

[MBY.p.adjust](#), [MBL.p.adjust](#)

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MBH.p.adjust(p,p.set)
```

---

MBL.p.adjust	<i>The adjusted p-values for Modified Benjamini-Liu (BL) step-down FDR controlling procedure.</i>
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---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
MBL.p.adjust(p, p.set, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with NAs). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p).

## Note

The MBL procedure for discrete data controls FDR under the specific dependence assumption where the joint distribution of statistics from true nulls are independent of the joint distribution of statistics from false nulls.

## Author(s)

Yalin Zhu

## References

Benjamini, Y., and Liu, W. (1999). A step-down multiple hypotheses testing procedure that controls the false discovery rate under independence. *Journal of Statistical Planning and Inference*, **82**: 163-170.

## See Also

[MBH.p.adjust](#), [MBY.p.adjust](#)

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MBL.p.adjust(p,p.set)
```

---

MBonf.p.adjust	<i>The adjusted p-values for Modified Bonferroni single-step FWER controlling procedure.</i>
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---

**Description**

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values

**Usage**

```
MBonf.p.adjust(p, p.set, alpha, make.decision)
```

**Arguments**

p	numeric vector of p-values (possibly with NAs). Any other R is coerced by <code>as.numeric</code> . Same as in <code>p.adjust</code> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

**Value**

A numeric vector of the adjusted p-values (of the same length as p) if `make.decision = FALSE`, or a list including original p-values, adjusted p-values and decision rules if `make.decision = TRUE`.

**Note**

The attainable p-value refers to the element of domain set of p-value for the corresponding hypothesis. For continuous test statistics, the p-value under true null are uniform distributed in (0,1), thus the p-values are attainable everywhere between 0 and 1. But for discrete test statistics, the p-value can only take finite values bewtween 0 and 1, that is the attainable p-values for discrete case are finite and countable, so we can assign them in a finite list p.set.

**Author(s)**

Yalin Zhu

**References**

Zhu, Y., & Guo, W. (2017). Familywise error rate controlling procedures for discrete data *arXiv preprint* arXiv:1711.08147.

**See Also**

[Tarone.p.adjust](#), [MixBonf.p.adjust](#), [p.adjust](#).

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MBonf.p.adjust(p,p.set)
## Compare with the traditional Bonferroni adjustment
p.adjust(p,method = "bonferroni")
## Compare with the Tarone adjustment
Tarone.p.adjust(p,p.set)
```

---

MBY.p.adjust

*The adjusted p-values for Modified Benjamini-Yekutieli (BY) step-up FDR controlling procedure.*

---

**Description**

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

**Usage**

```
MBY.p.adjust(p, p.set, alpha, make.decision)
```

**Arguments**

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

**Value**

A numeric vector of the adjusted p-values (of the same length as p).

**Author(s)**

Yalin Zhu

**References**

Benjamini, Y., and Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. *Annals of Statistics*, **29**: 1165-1188.



**See Also**

[MBH.p.adjust](#), [MBL.p.adjust](#)

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MBY.p.adjust(p,p.set)
```

---

MHoch.p.adjust

*The adjusted p-values for Modified Hochberg step-up FWER controlling procedure.*

---

**Description**

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

**Usage**

```
MHoch.p.adjust(p, p.set, alpha, make.decision)
```

**Arguments**

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis..
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

**Value**

A numeric vector of the adjusted p-values (of the same length as p).

**Author(s)**

Yalin Zhu

**References**

Zhu, Y., & Guo, W. (2017). Familywise error rate controlling procedures for discrete data *arXiv preprint* arXiv:1711.08147.

Hochberg, Y. (1988). A sharper Bonferroni procedure for multiple tests of significance. *Biometrika*, **75**: 800-803.

**See Also**

[Roth.p.adjust](#), [p.adjust](#).

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MHoch.p.adjust(p,p.set)
## Compare with the traditional Hochberg adjustment
p.adjust(p,method = "hochberg")
## Compare with the Roth adjustment
Roth.p.adjust(p,p.set)
```

---

MHolm.p.adjust	<i>The adjusted p-values for Modified Holm step-down FWER controlling procedure.</i>
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---

**Description**

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

**Usage**

```
MHolm.p.adjust(p, p.set, alpha, make.decision)
```

**Arguments**

p	numeric vector of p-values (possibly with NAs). Any other R is coerced by <code>as.numeric</code> . Same as in <code>p.adjust</code> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

**Value**

A numeric vector of the adjusted p-values (of the same length as p).

**Author(s)**

Yalin Zhu

**References**

Zhu, Y., & Guo, W. (2017). Familywise error rate controlling procedures for discrete data *arXiv preprint* arXiv:1711.08147.

Holm, S. (1979). A simple sequentially rejective multiple test procedure. *Scandinavian Journal of Statistics*, 6: 65-70.

**See Also**

[TH.p.adjust](#), [p.adjust](#).

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
MHolm.p.adjust(p,p.set)
## Compare with the traditional Holm adjustment
p.adjust(p,method = "holm")
## Compare with the Tarone-Holm adjustment
TH.p.adjust(p,p.set)
```

---

MHTdiscrete

---

*MHTdiscrete: A package for Multiple Hypotheses Testing for Discrete Data.*


---

## Description

The MHTdiscrete package provides two categories of important functions for discrete data multiple hypothesis testing:

### FWER controlling procedures

Single-step: MBonf.p.adjust, MixBonf.p.adjust, Tarone.p.adjust.

Step-down: MHolm.p.adjust, TH.p.adjust.

Step-up: MHoch.p.adjust, Roth.p.adjust .

### FDR controlling procedures

Step-down: MBL.p.adjust.

Step-up: MBH.p.adjust, GTBH.p.adjust, MBY.p.adjust, GTBY.p.adjust.

## Author(s)

Yalin Zhu

## References

- Tarone, R. E. (1990). A modified Bonferroni method for discrete data. *Biometrics*, **46**: 515-522.
- Hommel, G., & Krummenauer, F. (1998). Improvements and modifications of Tarone's multiple test procedure for discrete data. *Biometrics*, **54**: 673-681.
- Holm, S. (1979). A simple sequentially rejective multiple test procedure. *Scandinavian Journal of Statistics*, **6**: 65-70.
- Roth, A. J. (1999). Multiple comparison procedures for discrete test statistics. *Journal of statistical planning and inference*, **82**: 101-117.
- Gilbert, P. B. (2005). A modified false discovery rate multiple-comparisons procedure for discrete data, applied to human immunodeficiency virus genetics. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **54**: 143-158.
- Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society Series B*, **57**: 289-300.
- Benjamini, Y., and Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. *Annals of Statistics*, **29**: 1165-1188.

Benjamini, Y., and Liu, W. (1999). A step-down multiple hypotheses testing procedure that controls the false discovery rate under independence. *Journal of Statistical Planning and Inference*, **82**: 163-170.

---

MixBonf.p.adjust	<i>The adjusted p-values for Mixed Bonferroni single-step FWER controlling procedure.</i>
------------------	---

---

## Description

The function for calculating the adjusted p-values based on original available p-values and the attainable p-values for the discrete test statistics.

## Usage

```
MixBonf.p.adjust(pc, pd, pd.set, alpha, make.decision)
```

## Arguments

pc	numeric vector of the available p-values (possibly with NAs) for the continuous test statistics. Any other R is coerced by <code>as.numeric</code> . Same as in <code>p.adjust</code> .
pd	numeric vector of the available p-values (possibly with NAs) for the discrete test statistics. Any other R is coerced by <code>as.numeric</code> . Same as in <code>p.adjust</code> .
pd.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis for discrete data.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p) if `make.decision = FALSE`, or a list including original p-values, adjusted p-values and decision rules if `make.decision = TRUE`.

## Note

The arguments include three parts, the available p-values need to be reorganized in advance. Gather all available p-values for continuous data as pc, and all available p-values for discrete data as pd. The attainable p-value refers to the element of domain set of p-value for the corresponding hypothesis for discrete test statistics, the p-value can only take finite values between 0 and 1, that is, the attainable p-values for discrete case are finite and countable, so we can assign them in a finite list pd.set. The function returns the adjusted p-values with the first part for continuous data of the same length as pc, and second part for discrete data of the same length as pd

## Author(s)

Yalin Zhu

## References

Zhu, Y., & Guo, W. (2017). Familywise error rate controlling procedures for discrete data *arXiv preprint* arXiv:1711.08147.

## See Also

[Tarone.p.adjust](#), [MBonf.p.adjust](#), [p.adjust](#).

## Examples

```
pd <- c(pbinom(1,8,0.5),pbinom(1,5,0.75)); pc <- c(0.04, 0.1)
pd.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75))
MixBonf.p.adjust(pc,pd,pd.set)
## Compare with the traditional Bonferroni adjustment
p.adjust(c(pc,pd),method = "bonferroni")
```

---

Roth.p.adjust	<i>The adjusted p-values for Roth's step-up FWER controlling procedure.</i>
---------------	---

---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
Roth.p.adjust(p, p.set, digits, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with NAs). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis..
digits	minimal number of significant digits for the adjusted p-values, the default value is 4, see <a href="#">print.default</a> .
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p).

## Author(s)

Yalin Zhu

## References

Roth, A. J. (1999). Multiple comparison procedures for discrete test statistics. *Journal of statistical planning and inference*, **82**: 101-117.

## See Also

[MHoch.p.adjust](#), [p.adjust](#).

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
Roth.p.adjust(p,p.set,digits=5)
```

---

Roth.rej	<i>The number of rejected hypotheses for Roth's step-up FWER controlling procedure.</i>
----------	---

---

## Description

The function for calculating the number of rejected hypotheses (rejection region) based on original available p-values, all attainable p-values and the given significant level.

## Usage

```
Roth.rej(p,p.set,alpha)
```

## Arguments

- |       |   |
|-------|---|
| p     | numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> . |
| p.set | a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis..     |
| alpha | the given significant level for Roth's procedure, the default value is 0.05.  |

## Value

An integer value of the number of rejected hypotheses.

## Author(s)

Yalin Zhu

## References

Roth, A. J. (1999). Multiple comparison procedures for discrete test statistics. *Journal of statistical planning and inference*, **82**: 101-117.

Hochberg, Y. (1988). A sharper Bonferroni procedure for multiple tests of significance. *Biometrika*, **75**: 800-803.

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
Roth.rej(p,p.set,0.05)
```

---

Sidak.p.adjust	<i>The adjusted p-values for Sidak single-step FWER controlling procedure.</i>
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---

**Description**

The function for calculating the adjusted p-values based on original available p-values.

**Usage**

```
Sidak.p.adjust(p, alpha, make.decision)
```

**Arguments**

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

**Value**

A numeric vector of the adjusted p-values (of the same length as p) if make.decision = FALSE, or a list including original p-values, adjusted p-values and decision rules if make.decision = TRUE.

**Author(s)**

Yalin Zhu

**See Also**

[p.adjust](#).

**Examples**

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
Sidak.p.adjust(p)
```

---

Tarone.p.adjust	<i>The adjusted p-values for Tarone's single-step FWER controlling procedure.</i>
-----------------	---

---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
Tarone.p.adjust(p, p.set, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p) if make.decision = FALSE, or a list including original p-values, adjusted p-values and decision rules if make.decision = TRUE.

## Author(s)

Yalin Zhu

## References

Tarone, R. E. (1990). A modified Bonferroni method for discrete data. *Biometrics*, **46**: 515-522.

## See Also

[MBonf.p.adjust](#), [MixBonf.p.adjust](#), [p.adjust](#).

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
Tarone.p.adjust(p,p.set)
```



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TH.p.adjust	<i>The adjusted p-values for Tarone-Holm step-down FWER controlling procedure.</i>
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---

## Description

The function for calculating the adjusted p-values based on original available p-values and all attainable p-values.

## Usage

```
TH.p.adjust(p, p.set, alpha, make.decision)
```

## Arguments

p	numeric vector of p-values (possibly with <a href="#">NAs</a> ). Any other R is coerced by <a href="#">as.numeric</a> . Same as in <a href="#">p.adjust</a> .
p.set	a list of numeric vectors, where each vector is the vector of all attainable p-values containing the available p-value for the corresponding hypothesis.
alpha	significant level used to compare with adjusted p-values to make decisions, the default value is 0.05.
make.decision	logical; if TRUE, then the output include the decision rules compared adjusted p-values with significant level $\alpha$

## Value

A numeric vector of the adjusted p-values (of the same length as p).

## Author(s)

Yalin Zhu

## References

Hommel, G., & Krummenauer, F. (1998). Improvements and modifications of Tarone's multiple test procedure for discrete data. *Biometrics*, **54**: 673-681.

Holm, S. (1979). A simple sequentially rejective multiple test procedure. *Scandinavian Journal of Statistics*, **6**: 65-70.

## See Also

[MHolm.p.adjust](#), [p.adjust](#).

## Examples

```
p <- c(pbinom(1,8,0.5),pbinom(1,5,0.75),pbinom(1,6,0.6))
p.set <-list(pbinom(0:8,8,0.5),pbinom(0:5,5,0.75),pbinom(0:6,6,0.6))
TH.p.adjust(p,p.set)
```

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