

Package: RHSDB (via r-universe)

September 14, 2024

Type Package

Title Ryan-Holm Step-Down Bonferroni or Sidak Procedure

Version 0.2.0

Description The Ryan-Holm step-down Bonferroni or Sidak procedure is to control the family-wise (experiment-wise) type I error rate in the multiple comparisons. This procedure provides the adjusting p-values and adjusting CIs. The methods used in this package are referenced from John Ludbrook (2000) [doi:10.1046/j.1440-1681.2000.03223.x](https://doi.org/10.1046/j.1440-1681.2000.03223.x).

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NeedsCompilation no

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Repository <https://marsdu1989.r-universe.dev>

RemoteUrl <https://github.com/cran/RHSDB>

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rh.sd.bonferroni *Ryan-Holm Step-Down Bonferroni Procedure*

Description

This procedure provides the adjusting p-values and adjusting CIs.

Usage

```
rh.sd.bonferroni(p, effect, effect.se, df, type, sig, side, digits)
```

Arguments

| | |
|-----------|---|
| p | the raw p values |
| effect | the effect size from the multiple comparisons, e.g. the mean difference from t test or paried t test |
| effect.se | the standard error of effect size from the multiple comparisons, e.g. the standard error of mean difference from t test or paried t test) |
| df | the degree of freedom of hypothesis test, e.g. n_1+n_2-2 for t test, $n-1$ for paried t test |
| type | the type of the effect size, default is "mean" |
| sig | the significance level, default is 0.05 |
| side | "one" or "two" sided hypothesis test |
| digits | the number of decimal digits |

Value

| | |
|----------|---|
| p.adj | the adjusted p value |
| ci.adj.l | the lower limit of adjusted confidence interval |
| ci.adj.u | the upper limit of adjusted confidence interval |

Note

Please feel free to contact us, if you have any advice and find any bug!

Reference:

1. John Ludbrook (2000). MULTIPLE INFERENCES USING CONFIDENCE INTERVALS. Clinical and Experimental Pharmacology and Physiology. 27: 212-215.

Update:

Version 0.1.0: The first version.

Version 0.2.0: Fix the bug for maintaining monotonicity of the ranking p-values.

See Also

[rh.sd.sidak](#)

Examples

```
p=c(0.217,0.00028,0,0.001,0.024,0.719,0.00033)
effect=c(16,74,-85,-38,29,5,91)
effect.se=c(12,16,14,9,12,16,20)
df=16
rh.sd.bonferroni(p,effect,effect.se,df)
```

rh.sd.sidak

*Ryan-Holm Step-Down Sidak Procedure***Description**

This procedure provides the adjusting p-values and adjusting CIs.

Usage

```
rh.sd.sidak(p,effect,effect.se,df,type,sig,side,digits)
```

Arguments

| | |
|-----------|---|
| p | the raw p values |
| effect | the effect size from the multiple comparisons, e.g. the mean difference from t test or paried t test |
| effect.se | the standard error of effect size from the multiple comparisons, e.g. the standard error of mean difference from t test or paried t test) |
| df | the degree of freedom of hypothesis test, e.g. n_1+n_2-2 for t test, $n-1$ for paried t test |
| type | the type of the effect size, default is "mean" |
| sig | the significance level, default is 0.05 |
| side | "one" or "two" sided hypothesis test |
| digits | the number of decimal digits |

Value

| | |
|----------|---|
| p.adj | the adjusted p value |
| ci.adj.l | the lower limit of adjusted confidence interval |
| ci.adj.u | the upper limit of adjusted confidence interval |

Note

Please feel free to contact us, if you have any advice and find any bug!

Reference:

1. John Ludbrook (2000). MULTIPLE INFERENCES USING CONFIDENCE INTERVALS. *Clinical and Experimental Pharmacology and Physiology*. 27: 212-215.

Update:

Version 0.1.0: The first version.

Version 0.2.0: Fix the bug for maintaining monotonicity of the ranking p-values.

See Also

[rh.sd.bonferroni](#)

Examples

```
p=c(0.217,0.00028,0,0.001,0.024,0.719,0.00033)
effect=c(16,74,-85,-38,29,5,91)
effect.se=c(12,16,14,9,12,16,20)
df=16
rh.sd.sidak(p,effect,effect.se,df)
```

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