Package 'medScan'

March 6, 2024

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Title Large Scale Single Mediator Hypothesis Testing	
Version 1.0.2	
<pre>URL https://github.com/umich-cphds/medScan</pre>	
<pre>BugReports https://github.com/umich-cphds/medScan/issues</pre>	
Description A collection of methods for large scale single mediator hypothesis testing. The six included methods for testing the mediation effect are Sobel's test, Max P test, joint significance test under the composite null hypothesis, high dimensional mediation testing, divide-aggregate composite null test, and Sobel's test under the composite null hypothesis. Du et al (2023) <doi:10.1002 gepi.22510="">.</doi:10.1002>	
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medScan

Large Scale Single Mediator Hypothesis Testing

Description

A collection of methods for large scale single mediator hypothesis testing. The six included methods for testing the mediation effect are Sobel's test, Max P test, joint significance test under the composite null hypothesis, high dimensional mediation testing, divide-aggregate composite null test, and Sobel's test under the composite null hypothesis. Du, J., Zhou, X., Hao, W., Liu, Y., Smith, J. A., & Mukherjee, B. (2022). "Methods for Large-scale Single Mediator Hypothesis Testing: Possible Choices and Comparisons."

Usage

```
medScan(z.alpha, z.beta, method)
```

Arguments

z.alpha the z-test statistic for alpha (exposure effect on the mediator).

z.beta the z-test statistic for beta (mediator effect on the outcome).

method the method to use for testing the mediation effect. It should belong to one of the

six methods: "Sobel", "MaxP", "JT_comp", "HDMT", "DACT", and "Sobel_comp". (1) Sobel's test (method = "Sobel"), (2) Max P test (method = "MaxP"), (3) joint significance test under the composite null hypothesis (method = "JT_comp"), (4) high dimensional mediation testing (method = "HDMT"), (5) Divide-Aggregate Composite-null Test (method = "DACT"), and (6) Sobel's test under the compos-

ite null hypothesis (method = "Sobel_comp").

Details

The available methods are: (1) Sobel's test (method = "Sobel"), (2) Max P test (method = "MaxP"), (3) joint significance test under the composite null hypothesis (method = "JT_comp"), (4) high dimensional mediation testing (method = "HDMT"), (5) Divide-Aggregate Composite-null Test (method = "DACT"), and (6) Sobel's test under the composite null hypothesis (method = "Sobel_comp").

We incorporated code from the DACT R package formerly on CRAN. Author: Zhonghua Liu Maintainer: Zhonghua Liu, zhhliu@hku.hk.

Value

A list that contains

pvalues: p-values for all mediators from the chosen method.

pi: the estimated proportions of the three null cases from the HDMT method. pi00

is the proportion of alpha=beta=0; pi01 is the proportion of alpha=0 and beta!=0;

and pi10 is the proportion of alpha!=0 and beta=0.

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References

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Huang, Y. T. (2019). Genome-wide analyses of sparse mediation effects under composite null hypotheses. The Annals of Applied Statistics, 13(1), 60-84.

Liu, Z., Shen, J., Barfield, R., Schwartz, J., Baccarelli, A. A., & Lin, X. (2022). Large-scale hypothesis testing for causal mediation effects with applications in genome-wide epigenetic studies. Journal of the American Statistical Association, 117(537), 67-81.

Dai, J. Y., Stanford, J. L., & LeBlanc, M. (2022). A multiple-testing procedure for high-dimensional mediation hypotheses. Journal of the American Statistical Association, 117(537), 198-213.

Du, Jiacong, et al. "Methods for large-scale single mediator hypothesis testing: Possible choices and comparisons." Genetic Epidemiology 47.2 (2023): 167-184.

Examples

```
# simulate data under the mixture null
n=10000
u = runif(n, 0, 1)
z.alpha = z.beta = rep(NA,0)
pi00 = 0.98
pi10 = 0.01
pi01 = 0.01
for(i in 1:n){
  if(u[i]<=pi00){
   z.alpha[i] = rnorm(1, 0, 1)
    z.beta[i] = rnorm(1, 0, 1)
  } else if (u[i]<= pi00+pi10){</pre>
    z.alpha[i] = rnorm(1, 1, 1)
    z.beta[i] = rnorm(1, 0, 1)
  } else {
    z.alpha[i] = rnorm(1, 0, 1)
    z.beta[i] = rnorm(1, 1, 1)
}
# obtain p-values
# method = "Sobel", "MaxP", "HDMT", "Sobel_comp", "JT_comp", "DACT"
obj = medScan(z.alpha = z.alpha, z.beta = z.beta, method = "Sobel")
qqman::qq(obj\$pvalues, xlim = c(0,4), ylim = c(0,4), main = "Sobel")
```

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