# Package 'cobenrich'

April 15, 2025

Type Package

**Title** Using Multiple Continuous Biomarkers for Patient Enrichment in Two-Stage Clinical Designs

**Version** 1.0.1

**Description** Enrichment strategies play a critical role in modern clinical trial design, especially as precision medicine advances the focus on patient-specific efficacy. Recent developments in enrichment design have introduced biomarker randomness and accounted for the correlation structure between treatment effect and biomarker, resulting in a two-stage threshold enrichment design. We propose novel two-stage enrichment designs capable of handling two or more continuous biomarkers.

See Zhang, F. and Gou, J. (2025). Using multiple biomarkers for patient enrichment in twostage clinical designs. Technical Report.

License GPL-3

**Encoding** UTF-8

**Depends** R (>= 4.2.0)

**Imports** tmvtnorm (>= 1.2), stats (>= 4.0.0)

RoxygenNote 7.3.2

NeedsCompilation no

Author Jiangtao Gou [aut, cre], Fengqing (Zoe) Zhang [aut]

Maintainer Jiangtao Gou <gouRpackage@gmail.com>

Repository CRAN

Date/Publication 2025-04-15 19:30:12 UTC

# Contents

avetrteff2	 2
findATE2	 3
findSATE2	 4
targetDel	
targetLam	 6
	7

Index

avetrteff2

avetrteff2

Compute the average subpopulation treatment effect and the standardized average subpopulation treatment effect when two biomarkers are involved

# Description

Compute the average subpopulation treatment effect and the standardized average subpopulation treatment effect when two biomarkers are involved

# Usage

avetrteff2(z1z2, kappa, rhovec, sigma, muminusmu0)

#### Arguments

z1z2	a numeric vector of two numbers that are standardized biomarker values
kappa	a number of the correlation coefficient between two biomarkers
rhovec	a numeric vector of two correlation coefficients between the output and two biomarkers
sigma	a number of the standard deviation of outcome
muminusmu0	a number of the difference between the mean of outcome and the minimal clin- ically important treatment effect

# Value

a list of three numbers: delta is the average subpopulation treatment effect, lambda is the standardized average subpopulation treatment effect, and cVar is the variance with respect to the truncated distribution with specified cutoff values

### Author(s)

Jiangtao Gou

#### References

Zhang, F. and Gou, J. (2025). Using multiple biomarkers for patient enrichment in two-stage clinical designs. Technical Report.

#### Examples

```
x1x2 <- c(2, 1)
nu1nu2 <- c(0,0)
tau1tau2 <- c(1,1)
z1z2 <- (x1x2 - nu1nu2)/tau1tau2
muminusmu0 <- 1.8
kappa <- 0.1</pre>
```

#### findATE2

```
sigma <- 1
rhovec <- c(0.1, 0.2)
avetrteff2(z1z2, kappa, rhovec, sigma, muminusmu0)</pre>
```

#### findATE2

Find the cutoff values of biomarkers based on the average subpopulation treatment effect

# Description

Find the cutoff values of biomarkers based on the average subpopulation treatment effect

# Usage

findATE2(z2interval, kkk, muminusmu0, kappa, rhovec, sigma, cDel)

# Arguments

z2interval	a numeric vector of two values, including the lower and upper limits of the initial interval for z2
kkk	the researchers' weighting preference between the two biomarkers
muminusmu0	a number of the difference between the mean of outcome and the minimal clin- ically important treatment effect
kappa	a number of the correlation coefficient between two biomarkers
rhovec	a numeric vector of two correlation coefficients between the output and two biomarkers
sigma	a number of the standard deviation of outcome
cDel	the desired average subpopulation treatment effect

# Value

a numeric vector of two values which are the cutoff values for z1 and z2

# Author(s)

Jiangtao Gou Fengqing Zhang

# References

Zhang, F. and Gou, J. (2025). Using multiple biomarkers for patient enrichment in two-stage clinical designs. Technical Report.

# Examples

```
z2interval <- c(-5, 5)
kkk <- 1
muminusmu0 <- 1.8
kappa <- 0.1
rhovec <- c(0.1, 0.2)
sigma <- 1
cDel <- 2.5
findATE2(z2interval, kkk, muminusmu0, kappa, rhovec, sigma, cDel)</pre>
```

findSATE2	Find the cutoff values of biomarkers based on the standardized aver-
	age subpopulation treatment effect

# Description

Find the cutoff values of biomarkers based on the standardized average subpopulation treatment effect

# Usage

findSATE2(z2interval, kkk, muminusmu0, kappa, rhovec, sigma, cLam)

# Arguments

z2interval	a numeric vector of two values, including the lower and upper limits of the initial interval for z2
kkk	the researchers' weighting preference between the two biomarkers
muminusmu0	a number of the difference between the mean of outcome and the minimal clin- ically important treatment effect
kappa	a number of the correlation coefficient between two biomarkers
rhovec	a numeric vector of two correlation coefficients between the output and two biomarkers
sigma	a number of the standard deviation of outcome
cLam	the desired standardized average subpopulation treatment effect

#### Value

a numeric vector of two values which are the cutoff values for z1 and z2

# Author(s)

Jiangtao Gou Fengqing Zhang

4

# targetDel

#### References

Zhang, F. and Gou, J. (2025). Using multiple biomarkers for patient enrichment in two-stage clinical designs. Technical Report.

# Examples

```
z2interval <- c(-4, 4)
kkk <- 1
muminusmu0 <- 1.8
kappa <- 0.1
rhovec <- c(0.1, 0.2)
sigma <- 1
cLam <- 2.5
findSATE2(z2interval, kkk, muminusmu0, kappa, rhovec, sigma, cLam)</pre>
```

Find the difference between the average subpopulation treatment effect and the desired one

# Description

Find the difference between the average subpopulation treatment effect and the desired one

#### Usage

```
targetDel(z2, kkk, muminusmu0, kappa, rhovec, sigma, cDel)
```

#### Arguments

z2	the standardized biomarker value of the second biomarker
kkk	the researchers' weighting preference between the two biomarkers
muminusmu0	a number of the difference between the mean of outcome and the minimal clin- ically important treatment effect
kappa	a number of the correlation coefficient between two biomarkers
rhovec	a numeric vector of two correlation coefficients between the output and two biomarkers
sigma	a number of the standard deviation of outcome
cDel	the desired average subpopulation treatment effect

#### Value

the difference between the average subpopulation treatment effect and the desired one

targetLam

Find the difference between the standardized average subpopulation treatment effect and the desired one

# Description

Find the difference between the standardized average subpopulation treatment effect and the desired one

# Usage

```
targetLam(z2, kkk, muminusmu0, kappa, rhovec, sigma, cLam)
```

# Arguments

z2	the standardized biomarker value of the second biomarker
kkk	the researchers' weighting preference between the two biomarkers
muminusmu0	a number of the difference between the mean of outcome and the minimal clin- ically important treatment effect
kappa	a number of the correlation coefficient between two biomarkers
rhovec	a numeric vector of two correlation coefficients between the output and two biomarkers
sigma	a number of the standard deviation of outcome
cLam	the desired standardized average subpopulation treatment effect

# Value

the difference between the standardized average subpopulation treatment effect and the desired one

# Index

avetrteff2, 2

findATE2, 3
findSATE2, 4

targetDel,5 targetLam,6