# Package 'BRACE' 

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Type Package
Title Bias Reduction Through Analysis of Competing Events (BRACE)
Version 0.1.0
Maintainer Tuo Lin[tulin@health.ucsd.edu](mailto:tulin@health.ucsd.edu)
Description Adjusting the bias due to residual confounding (often called
treatment selection bias) in estimating the treatment effect in a
proportional hazard model, as described in Williamson et al.
(2022) [doi:10.1158/1078-0432.ccr-21-2468](doi:10.1158/1078-0432.ccr-21-2468).
License GPL (>=3)
Depends survival, survminer
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Author Tuo Lin [aut, cre],
Jingjing Zou [aut],
Loren Mell [aut]
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## $R$ topics documented:

brace ..... 2
gendat ..... 4
gendat2 ..... 5
Index ..... 6

## Description

brace is used to estimate the treatment effect with adjusted confounders on the composite hazard for primary or competing events, and adjust for bias from residual confounding in non-randomized data by BRACE method

## Usage

brace(
ftime, fstatus, covs = NA, trt,
failcode $=1$,
cencode $=0$,
PS = 0,
$B=1000$
)

## Arguments

| ftime | vector of failure/censoring times |
| :--- | :--- |
| fstatus | vector with a unique code for each failure type and a separate code for censored <br> observations (default is primary event $=1$, competing event $=2$, censored $=0)$ <br> matrix (nobs x ncovs) of fixed covariates. If no covariates, set covs = NA (default <br> is NA) |
| covs | vector of treatment indicator (1 for treatment group) <br> trt |
| failcode | code of fstatus that denotes the failure type of interest <br> cencode of fstatus that denotes censored observations |
| PS | whether to use propensity score method for adjusting the confounding effect (1 <br> for propensity score method, default is 0) <br> bootstrap sample size for calculating the Confidence interval, default is 1000 |
| B |  |

## Value

a list of class brace, with components:
\$Summary summary table of BRACE method
\$‘BRACE HR Distribution‘
the estimated regression coefficients in each bootstrap sample
\$‘Omega Estimate‘
estimate of relative hazards for primary events vs. combined events


## References

Williamson, Casey W., et al. "Bias Reduction through Analysis of Competing Events (BRACE) Correction to Address Cancer Treatment Selection Bias in Observational Data." Clinical Cancer Research 28.9 (2022): 1832-1840.

## Examples

```
nsims = 1; nobs = 1500
f = 0.5; g = 0.333; b = 8; w1 = w2 = 0.667
theta1 = 0.5; theta2 = 1; omegaplus = 1; k3 = 0.333
sim1 = gendat(nsims, nobs,f,g,b,w1,w2,omegaplus,theta1, theta2,k3)
ftime = sim1$time
fstatus = sim1$pfs_ci
covs = NA
trt = sim1$group
braceoutput = brace(ftime, fstatus, covs, trt, PS=0, B=10)
nsims = 1; nobs = 1500
f1 = f2 = 0.5; g = 0.333; b1 = 8; b2 = 4; w1 = w2 = 0.667
theta1 = 0.5; theta2 = 1; omegaplus = 1; k3 = 0.333
sim1 = gendat2(nsims,nobs,f1,f2,g,b1,b2,w1,w2,omegaplus,theta1,theta2,k3)
ftime = sim1$time
fstatus = sim1$pfs_ci
covs = sim1$factor2
trt = sim1$group
braceoutput = brace(ftime, fstatus, covs, trt, PS=1, B=10)
```


## Description

generating the simulation data to apply in brace

## Usage

gendat(nsims, nobs, f, g, b, w1, w2, omegaplus, theta1, theta2, k3)

## Arguments

nsims number of simulation datasets
nobs number of observations for one dataset
f parameter for generating unmeasured binary confounder
g parameter for generating group assignment
b confounder effect on group assignment
w1 shape parameter in generating survival time for event 1 from weibull distribution
w2 shape parameter in generating survival time for event 2 from weibull distribution
omegaplus multiplier on the baseline hazard for event 1
theta1 multiplier on the baseline hazard for event 1
theta2 multiplier on the baseline hazard for event 2
k3 multiplier on the baseline hazard for event 2

## Value

a matrix of nsims* nobs row, which consists of nsims datasets

## Examples

```
nsims = 1; nobs = 1500
f = 0.5; g = 0.333; b = 8; w1 = w2 = 0.667
theta1 = 0.5; theta2 = 1; omegaplus = 1; k3 = 0.333
sim1 = gendat(nsims, nobs,f,g,b,w1,w2,omegaplus,theta1, theta2,k3)
```

gendat2 simulation data generating function (Adding a measured confounder)

## Description

generating the simulation data to apply in brace

## Usage

gendat2(nsims, nobs, f1, f2, g, b1, b2, w1, w2, omegaplus, theta1, theta2, k3)

## Arguments

nsims number of simulation datasets
nobs number of observations for one dataset
f1 parameter for generating unmeasured binary confounder
f2 parameter for generating measured binary confounder
$\mathrm{g} \quad$ parameter for generating group assignment
b1 unmeasured confounder effect on group assignment
b2 measured confounder effect on group assignment
w1 shape parameter in generating survival time for event 1 from weibull distribution
w2 shape parameter in generating survival time for event 2 from weibull distribution
omegaplus multiplier on the baseline hazard for event 1
theta1 multiplier on the baseline hazard for event 1
theta2 multiplier on the baseline hazard for event 2
k3 multiplier on the baseline hazard for event 2

## Value

a matrix of nsims*nobs row, which consists of nsims datasets

## Examples

```
nsims = 1; nobs = 1500
f1 = f2 = 0.5; g = 0.333; b1 = 8; b2 = 4; w1 = w2 = 0.667
theta1 = 0.5; theta2 = 1; omegaplus = 1; k3 = 0.333
sim1 = gendat2(nsims,nobs,f1,f2,g,b1,b2,w1,w2,omegaplus,theta1,theta2,k3)
```


## Index

brace, 2
gendat, 4
gendat2, 5

