Package: powerSurvEpi (via r-universe)

September 9, 2024

Version 0.1.3

Date 2021-02-28

- Title Power and Sample Size Calculation for Survival Analysis of Epidemiological Studies
- Author Weiliang Qiu, Jorge Chavarro, Ross Lazarus, Bernard Rosner, Jing Ma.

Maintainer Weiliang Qiu <weiliang.qiu@gmail.com>

Depends R $(>= 3.6.0)$

Imports stats, survival, pracma

Description Functions to calculate power and sample size for testing main effect or interaction effect in the survival analysis of epidemiological studies (non-randomized studies), taking into account the correlation between the covariate of the interest and other covariates. Some calculations also take into account the competing risks and stratified analysis. This package also includes a set of functions to calculate power and sample size for testing main effect in the survival analysis of randomized clinical trials and conditional logistic regression for nested case-control study.

License GPL $(>= 2)$

NeedsCompilation no

Date/Publication 2021-03-01 06:30:02 UTC

Repository https://weiliang.r-universe.dev

RemoteUrl https://github.com/cran/powerSurvEpi

RemoteRef HEAD

RemoteSha 4d757552133847363b6aa8a3ce2a44ab7eda56d2

Contents

numDEpi *Calculate Number of Deaths Required for Cox Proportional Hazards Regression with Two Covariates for Epidemiological Studies*

Description

Calculate number of deaths required for Cox proportional hazards regression with two covariates for epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates. Some parameters will be estimated based on a pilot data set.

Usage

```
numDEpi(X1,
X2,
power,
theta,
alpha = 0.05
```
numDEpi 3

Arguments

Details

This is an implementation of the calculation of the number of required deaths derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 should be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of deaths required to achieve a power of $1 - \beta$ is

$$
D = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2}{[\log(\theta)]^2 p (1-p)(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution,

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

 p and rho will be estimated from a pilot data set.

Value

D the number of deaths required to achieve the desired power with given type I error rate.

- p proportion of subjects taking $X_1 = 1$.
- rho2 square of the correlation between X_1 and X_2 .

(1) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

(2) When rho2=0, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio θ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

See Also

[numDEpi.default](#page-3-1)

Examples

```
# generate a toy pilot data set
X1 \leftarrow c(\text{rep}(1, 39), \text{rep}(0, 61))set.seed(123456)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
res \leq numDEpi(X1 = X1,
X2 = X2,
 power = 0.8,
 theta = 2,
 alpha = 0.05print(res)
# proportion of subjects died of the disease of interest.
psi <- 0.505
# total number of subjects required to achieve the desired power
ceiling(res$D / psi)
```
numDEpi.default *Calculate Number of Deaths Required for Cox Proportional Hazards Regression with Two Covariates for Epidemiological Studies*

Note

numDEpi.default 5

Description

Calculate number of deaths required for Cox proportional hazards regression with two covariates for epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates.

Usage

```
numDEpi.default(power,
  theta,
  p,
  rho2,
  alpha = 0.05
```
Arguments

Details

This is an implementation of the calculation of the number of required deaths derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 should be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of deaths required to achieve a power of $1 - \beta$ is

$$
D = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2}{[\log(\theta)]^2 p (1-p)(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution,

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1 - q)}{p(1 - p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

Value

The number of deaths required to achieve the desired power with given type I error rate.

Note

(1) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

(2) When rho2=0, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio θ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

See Also

[numDEpi](#page-1-1)

Examples

```
# Example at the end of Section 5.2 of Latouche et al. (2004)
# for a cohort study.
D \le - numDEpi.default(power = 0.8,
       theta = 2,
       p = 0.39,
                     rho2 = 0.132^2.
       alpha = 0.05# proportion of subjects died of the disease of interest.
psi <- 0.505
# total number of subjects required to achieve the desired power
ceiling(D / psi)
```
Oph *Ophthalmology Data*

Description

The Ophthalmology data set is described in Example 14.41 on page 807 in Rosner (2006).

Oph 7

Usage

data(Oph)

Format

A data frame with 354 observations on the following 3 variables.

- times a numeric vector recording the survival/censoring time for each event/censoring.
- status a numeric vector recording if a observed time is event time (status=1) or censoring time (status=0).

group a factor with levels C (indicating control group) and E (indicating experimental group).

Details

This data set was from a clinical trial (Berson et al., 1993) conducted to test the efficacy of different vitamin supplements in preventing visual loss in patients with retinitis pigmentosa. Rosner (2006) used the data from this clinical trial to illustrate the analysis of survival data (Sections 14.9-14.12 of Rosner (2006)).

The data set consists of two groups of participants: (1) the experimental group (i.e., group E in which participants receiving 15,000 IU of vitamin A per day) and (2) the control group (i.e., group C in which participants receiving 75 IU of vitamin A per day).

The participants were enrolled over a 2-year period (1984-1987) and followed for a maximum of 6 years. The follow-up was terminated in September 1991. Some participants dropped out of the study before September 1991 and had not failed. Dropouts were due to death, other diseases, or side effects possibly due to the study medications, or unwillingness to comply (take study medications). There are 6 time points (at 1st year, 2nd year, 3rd year, 4th year, 5-th year, and 6-th year) in this data set.

Rosner (2006, page 786) defined the participants who do not reach a disease endpoint during their period of follow-up as censored observations. A participant has been censored at time t if the participant has been followed up to time t and has not failed. Noninformative censoring is assumed. That is, participants who are censored have the same underlying survival curve after their censoring time as patients who are not censored.

Source

Created based on Table 14.12 on page 787 of Rosner (2006).

References

Berson, E.L., Rosner, B., Sandberg, M.A., Hayes, K.C., Nicholson, B.W., Weigel-DiFranco, C., and Willett, W.C. (1993). A randomized trial of vitamin A and vitamin E supplementation for retinitis pigmentosa. *Archives of Ophthalmology*. 111:761-772.

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

Examples

data(Oph)

Description

Power calculation for survival analysis with binary predictor and exponential survival function.

Usage

```
power.stratify(
    n,
    timeUnit,
    gVec,
   PVec,
    HR,
    lambda0Vec,
    power.ini = 0.8,
    power.low = 0.001,
    power.upp = 0.999,
    alpha = 0.05,
    verbose = TRUE)
```


power.stratify 9

Details

We assume (1) there is only one predictor and no covariates in the survival model (exponential survival function); (2) there are m strata; (3) the predictor x is a binary variable indicating treatment group 1 $(x = 1)$ or treatment group 0 $(x = 0)$; (3) the treatment effect is constant over time (proportional hazards); (4) the hazard ratio is the same in all strata, and (5) the data will be analyzed by the stratified log rank test.

The sample size formula is Formula (1) on page 801 of Palta M and Amini SB (1985):

$$
n=(Z_{\alpha}+Z_{\beta})^2/\mu^2
$$

where α is the Type I error rate, β is the Type II error rate (power= $1 - \beta$), Z_{α} is the $100(1 - \alpha)$ -th percentile of standard normal distribution, and

$$
\mu = \log(\delta) \sqrt{\sum_{s=1}^{m} g_s P_s (1 - P_s) V_s}
$$

and

$$
V_s = P_s \left[1 - \frac{1}{\lambda_{1s}} \left\{ \exp\left[-\lambda_{1s}(T-1)\right] - \exp(-\lambda_{1s}T) \right\} \right] + (1 - P_s) \left[1 - \frac{1}{\lambda_{0s}} \left\{ \exp\left[-\lambda_{0s}(T-1)\right] - \exp(-\lambda_{0s}T) \right\} \right]
$$

In the above formulas, m is the number of strata, T is the total study length, δ is the hazard ratio, g_s is the proportion of the total sample size in stratum s, P_s is the proportion of stratum s, which is in treatment group 1, and λ_{is} is the hazard for the *i*-th treatment group in stratum *s*.

Value

A list of 2 elments.

References

Palta M and Amini SB. (1985). Consideration of covariates and stratification in sample size determination for survival time studies. *Journal of Chronic Diseases*. 38(9):801-809.

See Also

[ssize.stratify](#page-36-1)

```
# example on page 803 of Palta M and Amini SB. (1985).
res.power <- power.stratify(
 n = 146,
 timeUnit = 1.25,
 gVec = c(0.5, 0.5),
 PVec = c(0.5, 0.5),
```

```
HR = 1 / 1.91,lambda0Vec = c(2.303, 1.139),
power.ini = 0.8,
power.low = 0.001,
power.upp = 0.999,
alpha = 0.05,
verbose = TRUE
)
```
powerConLogistic.bin *Sample Size Calculation for Conditional Logistic Regression with Binary Covariate*

Description

Sample Size Calculation for Conditional Logistic Regression with Binary Covariate, such as matched logistic regression or nested case-control study.

Usage

```
powerConLogistic.bin(
 N = NULL,power = 0.8,
 OR,
 pE,
 nD,
 nH,
 R2 = 0,alpha = 0.05,
 nTests = 1,OR.low = 1.01,
 OR.upp = 100)
```


Details

The power and sample size calculation formulas are provided by Lachin (2008, Section 3.3, Formula (38))

$$
power = \Phi\left(\sqrt{Nc} - z_{\alpha/(2nTests)}\right)
$$

and

$$
N = (z_{power} + z_{\alpha/(2nTests)})^2/c
$$

where Φ is the cumulative distribution function of the standard normal distribution $N(0, 1)$, z_a is the upper $100a$ -th percentile of $N(0, 1)$,

$$
c = \theta^2 pE(1 - pE)(1 - R^2)nD * nH/(nD + nH)
$$

and $R²$ is the coefficient of determination for linear regression linking the exposure with other covariates.

Value

If the inputs is.null(N) = TRUE and is.null(power) = FALSE, then the function returns the number N of sets.

If the inputs is.null(N) = FALSE and is.null(power) = TRUE, then the function returns the power. Otherwise, an error message is output.

References

Lachin, JM Sample Size Evaluation for a Multiply Matched Case-Control Study Using the Score Test From a Conditional Logistic (Discrete Cox PH) Regression Model. Stat Med. 2008 27(14): 2509-2523

```
# estimate power
power = powerConLogistic.bin(
 N = 59,
 power = NULL,
 OR = 3.5,
 pE = 0.15,
 nD = 1,
```

```
nH = 2,
  R2 = 0,alpha = 0.05,
  nTests = 1)print(power) # 0.80
# estimate N (number of sets)
N = powerConLogistic.bin(
  N = NULL,power = 0.80,
  OR = 3.5,
  pE = 0.15,
  nD = 1,
  nH = 2,
  R2 = 0,alpha = 0.05,
  nTests = 1)
print(ceiling(N)) # 59
# estimate OR
OR = powerConLogistic.bin(
  N = 59,power = 0.80,
  OR = NULL,pE = 0.15,
  nD = 1,
  nH = 2,
  R2 = 0,alpha = 0.05,
  nTests = 1,
  OR.low = 1.01,
  OR.upp = 100)
print(OR) # 3.49
```
powerConLogistic.con *Sample Size Calculation for Conditional Logistic Regression with Continuous Covariate*

Description

Sample Size Calculation for Conditional Logistic Regression with Continuous Covariate, such as matched logistic regression or nested case-control study.

powerConLogistic.con 13

Usage

```
powerConLogistic.con(
 N = NULL,power = 0.8,
 OR,
 sigma,
 nD,
 nH,
 R2 = 0,alpha = 0.05,
 nTests = 1,
 OR.low = 1.01,
 OR.upp = 100
```
)

Arguments

Details

The power and sample size calculation formulas are provided by Lachin (2008, Section 3.1, Formulas (24) and (25))

$$
power = \Phi\left(\sqrt{Nc} - z_{\alpha/(2nTests)}\right)
$$

and

$$
N = (z_{power} + z_{\alpha/(2nTests)})^2/c
$$

where Φ is the cumulative distribution function of the standard normal distribution $N(0, 1)$, z_a is the upper $100a$ -th percentile of $N(0, 1)$,

$$
c = \theta^2 \sigma^2 n D(1 - 1/b)(1 - R^2)
$$

and b is the Binomial coefficient (*n* chooses *nD*), $n = nD + nH$, and R^2 is the coefficient of determination for linear regression linking the exposure with other covariates.

Value

If the inputs is.null(N) = TRUE and is.null(power) = FALSE, then the function returns the number N of sets.

If the inputs is.null(N) = FALSE and is.null(power) = TRUE, then the function returns the power. Otherwise, an error message is output.

References

Lachin, JM Sample Size Evaluation for a Multiply Matched Case-Control Study Using the Score Test From a Conditional Logistic (Discrete Cox PH) Regression Model. Stat Med. 2008 27(14): 2509-2523

```
library(pracma)
```

```
# Section 4.1 in Lachin (2008)
# estimate number of sets
N = powerConLogistic.com(N = NULL,power = 0.85,
                                OR = 1.39,
                                sigma = 1,
                                nD = 1,
                                nH = 2,
                                R2 = 0,
                                alpha = 0.05,
                                nTests = 1)
print(ceiling(N)) # 125
# estimate power
power = powerConLogistic.con(N = 125,
                                power = NULL,
                                OR = 1.39,
                                sigma = 1,
                                nD = 1,
                                nH = 2,
                                R2 = 0,
                                alpha = 0.05,
                                nTests = 1)
print(power) # 0.85
```
 $powerCT$ 15

```
# estimate OR
OR = powerConLogistic.con(N = 125,
                                power = 0.85,
                                OR = NULL,sigma = 1,
                                nD = 1,
                                nH = 2,
                                R2 = 0,alpha = 0.05,
                                nTests = 1)
```
print(OR) # 1.39

powerCT *Power Calculation in the Analysis of Survival Data for Clinical Trials*

Description

Power calculation for the Comparison of Survival Curves Between Two Groups under the Cox Proportional-Hazards Model for clinical trials. Some parameters will be estimated based on a pilot data set.

Usage

```
powerCT(formula,
dat,
nE,
nC,
RR,
alpha = 0.05
```


Details

This is an implementation of the power calculation method described in Section 14.12 (page 807) of Rosner (2006). The method was proposed by Freedman (1982).

The movitation of this function is that some times we do not have information about m or p_E and p_C available, but we have a pilot data set that can be used to estimate p_E and p_C hence m, where $m = n_E p_E + n_C p_C$ is the expected total number of events over both groups, n_E and n_C are numbers of participants in group E (experimental group) and group C (control group), respectively. p_E is the probability of failure in group E (experimental group) over the maximum time period of the study (t years). p_C is the probability of failure in group C (control group) over the maximum time period of the study (t years).

Suppose we want to compare the survival curves between an experimental group (E) and a control group (C) in a clinical trial with a maximum follow-up of t years. The Cox proportional hazards regression model is assumed to have the form:

$$
h(t|X_1) = h_0(t) \exp(\beta_1 X_1).
$$

Let n_E be the number of participants in the E group and n_C be the number of participants in the C group. We wish to test the hypothesis $H0:RR = 1$ versus $H1:RR$ not equal to 1, where $RR = \exp(\beta_1)$ =underlying hazard ratio for the E group versus the C group. Let RR be the postulated hazard ratio, α be the significance level. Assume that the test is a two-sided test. If the ratio of participants in group E compared to group $C = n_E/n_C = k$, then the power of the test is

power =
$$
\Phi(\sqrt{k*m} * |RR - 1|/(k * RR + 1) - z_{1-\alpha/2}),
$$

where

$$
m = n_E p_E + n_C p_C, \quad
$$

and $z_{1-\alpha/2}$ is the 100(1 – $\alpha/2$)-th percentile of the standard normal distribution $N(0, 1)$, Φ is the cumulative distribution function (CDF) of $N(0, 1)$.

 p_C and p_E can be calculated from the following formulaes:

$$
p_C = \sum_{i=1}^{t} D_i, p_E = \sum_{i=1}^{t} E_i,
$$

where $D_i = \lambda_i A_i C_i$, $E_i = RR\lambda_i B_i C_i$, $A_i = \prod_{j=0}^{i-1} (1 - \lambda_j)$, $B_i = \prod_{j=0}^{i-1} (1 - RR\lambda_j)$, $C_i =$ $\prod_{j=0}^{i-1}(1-\delta_j)$. And λ_i is the probability of failure at time i among participants in the control group, given that a participant has survived to time $i - 1$ and is not censored at time $i - 1$, i.e., the approximate hazard time i in the control group, $i = 1, ..., t$; $RRlambda_i$ is the probability of failure at time i among participants in the experimental group, given that a participant has survived to time $i-1$ and is not censored at time $i-1$, i.e., the approximate hazard time i in the experimental group, $i = 1, ..., t$; delta is the probability that a participant is censored at time i given that he was followed up to time i and has not failed, $i = 0, 1, ..., t$, which is assumed the same in each group.

Value

mat.lambda a matrix with 9 columns and nTimes+1 rows, where nTimes is the number of observed time points for the control group in the data set. The 9 columns are (1) time - observed time point for the control group; (2) lambda; (3) RRlambda; (4)

Note

(1) The estimates of $RRlambda_i = RR * \lambda_i$. That is, RR1 ambda is not directly estimated based on data from the experimental group; (2) The power formula assumes that the central-limit theorem is valid and hence is appropriate for large samples.

References

Freedman, L.S. (1982). Tables of the number of patients required in clinical trials using the log-rank test. *Statistics in Medicine*. 1: 121-129

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

See Also

[powerCT.default0](#page-19-1), [powerCT.default](#page-17-1)

```
# Example 14.42 in Rosner B. Fundamentals of Biostatistics.
# (6-th edition). (2006) page 809
```

```
library(survival)
```

```
data(Oph)
res \leq powerCT(formula = Surv(times, status) \sim group,
dat = 0ph,nE = 200,
nC = 200,
 RR = 0.7,
 alpha = 0.05)
```

```
print(round(res$mat.lambda, 4))
# Table 14.12 on page 787 of Rosner (2006)
print(round(res$mat.event, 4))
# the power
print(round(res$power, 2))
```
powerCT.default *Power Calculation in the Analysis of Survival Data for Clinical Trials*

Description

Power calculation for the Comparison of Survival Curves Between Two Groups under the Cox Proportional-Hazards Model for clinical trials.

Usage

```
powerCT.default(nE,
 nC,
 pE,
 pC,
 RR,
  alpha = 0.05
```
Arguments

Details

This is an implementation of the power calculation method described in Section 14.12 (page 807) of Rosner (2006). The method was proposed by Freedman (1982).

Suppose we want to compare the survival curves between an experimental group (E) and a control group (C) in a clinical trial with a maximum follow-up of t years. The Cox proportional hazards regression model is assumed to have the form:

$$
h(t|X_1) = h_0(t) \exp(\beta_1 X_1).
$$

powerCT.default 19

Let n_E be the number of participants in the E group and n_C be the number of participants in the C group. We wish to test the hypothesis $H0 : RR = 1$ versus $H1 : RR$ not equal to 1, where $RR = \exp(\beta_1)$ =underlying hazard ratio for the E group versus the C group. Let RR be the postulated hazard ratio, α be the significance level. Assume that the test is a two-sided test. If the ratio of participants in group E compared to group $C = n_E/n_C = k$, then the power of the test is

power =
$$
\Phi(\sqrt{k*m} * |RR - 1|/(k * RR + 1) - z_{1-\alpha/2}),
$$

where

$$
m = n_E p_E + n_C p_C,
$$

and $z_{1-\alpha/2}$ is the 100(1 – $\alpha/2$)-th percentile of the standard normal distribution $N(0, 1)$, Φ is the cumulative distribution function (CDF) of $N(0, 1)$.

Value

The power of the test.

Note

The power formula assumes that the central-limit theorem is valid and hence is appropriate for large samples.

References

Freedman, L.S. (1982). Tables of the number of patients required in clinical trials using the log-rank test. *Statistics in Medicine*. 1: 121-129

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

See Also

[powerCT.default0](#page-19-1), [powerCT](#page-14-1)

```
# Example 14.42 in Rosner B. Fundamentals of Biostatistics.
# (6-th edition). (2006) page 809
powerCT.default(nE = 200,
 nC = 200,
 pE = 0.3707,
 pC = 0.4890,RR = 0.7,
  alpha = 0.05)
```


Description

Power calculation for the Comparison of Survival Curves Between Two Groups under the Cox Proportional-Hazards Model for clinical trials.

Usage

```
powerCT.default0(k,
   m,
   RR,
   alpha = 0.05)
```
Arguments

Details

This is an implementation of the power calculation method described in Section 14.12 (page 807) of Rosner (2006). The method was proposed by Freedman (1982).

Suppose we want to compare the survival curves between an experimental group (E) and a control group (C) in a clinical trial with a maximum follow-up of t years. The Cox proportional hazards regression model is assumed to have the form:

$$
h(t|X_1) = h_0(t) \exp(\beta_1 X_1).
$$

Let n_E be the number of participants in the E group and n_C be the number of participants in the C group. We wish to test the hypothesis $H0 : RR = 1$ versus $H1 : RR$ not equal to 1, where $RR = \exp(\beta_1)$ =underlying hazard ratio for the E group versus the C group. Let RR be the postulated hazard ratio, α be the significance level. Assume that the test is a two-sided test. If the ratio of participants in group E compared to group $C = n_E/n_C = k$, then the power of the test is

power =
$$
\Phi(\sqrt{k*m} * |RR - 1|/(k * RR + 1) - z_{1-\alpha/2}),
$$

where $z_{1-\alpha/2}$ is the 100(1 – $\alpha/2$)-th percentile of the standard normal distribution $N(0, 1)$, Φ is the cumulative distribution function (CDF) of $N(0, 1)$.

Value

The power of the test.

powerEpi 21

Note

The power formula assumes that the central-limit theorem is valid and hence is appropriate for large samples.

References

Freedman, L.S. (1982). Tables of the number of patients required in clinical trials using the log-rank test. *Statistics in Medicine*. 1: 121-129

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

See Also

[powerCT.default](#page-17-1), [powerCT](#page-14-1)

Examples

Example 14.42 in Rosner B. Fundamentals of Biostatistics. # (6-th edition). (2006) page 809 powerCT.default0($k = 1$, $m = 171.9$, $RR = 0.7$, alpha = 0.05)

powerEpi *Power Calculation for Cox Proportional Hazards Regression with Two Covariates for Epidemiological Studies*

Description

Power calculation for Cox proportional hazards regression with two covariates for epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates. Some parameters will be estimated based on a pilot data set.

Usage

```
powerEpi(X1, X2, failureFlag, n, theta, alpha = 0.05)
```


Details

This is an implementation of the power calculation formula derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 should be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\beta_1) = \theta$ is

$$
power = \Phi\left(-z_{1-\alpha/2} + \sqrt{n[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)}\right),\,
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1 - q)}{p(1 - p)}}
$$

,

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

 p, ρ^2 , and ψ will be estimated from a pilot data set.

Value

Note

(1) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

(2) When $\rho^2 = 0$, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio θ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

powerEpi.default 23

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

See Also

[powerEpi.default](#page-22-1)

Examples

```
# generate a toy pilot data set
X1 <- c(rep(1, 39), rep(0, 61))
set.seed(123456)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.5, 0.5), replace = TRUE)
powerEpi(X1 = X1, X2 = X2, failureFlag = failureFlag,
 n = 139, theta = 2, alpha = 0.05)
```


Description

Power calculation for Cox proportional hazards regression with two covariates for epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates.

Usage

```
powerEpi.default(n,
   theta,
   p,
   psi,
   rho2,
   alpha = 0.05
```


,

Details

This is an implementation of the power calculation formula derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 should be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\beta_1) = \theta$ is

$$
power = \Phi\left(-z_{1-\alpha/2} + \sqrt{n[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)}\right)
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

Value

The power of the test.

Note

(1) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

(2) When rho2=0, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio θ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

powerEpiCont 25

See Also

[powerEpi](#page-20-1)

Examples

```
# Example at the end of Section 5.2 of Latouche et al. (2004)
# for a cohort study.
powerEpi.default(n = 139,
  theta = 2,
   p = 0.39,
  psi = 0.505,
                 rho2 = 0.132^2,
   alpha = 0.05)
```


powerEpiCont *Power Calculation for Cox Proportional Hazards Regression with Nonbinary Covariates for Epidemiological Studies*

Description

Power calculation for Cox proportional hazards regression with nonbinary covariates for Epidemiological Studies. Some parameters will be estimated based on a pilot data set.

Usage

```
powerEpiCont(formula,
      dat,
      var.X1,
      var.failureFlag,
     n,
      theta,
      alpha = 0.05
```


Details

This is an implementation of the power calculation formula derived by Hsieh and Lavori (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is a nonbinary variable and X_2 is a vector of other covariates.

Suppose we want to check if the hazard ratio of the main effect $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\beta_1) = \theta$ is

$$
power = \Phi\left(-z_{1-\alpha/2} + \sqrt{n[\log(\theta)]^2 \sigma^2 \psi(1-\rho^2)}\right),\,
$$

where z_a is the 100a-th percentile of the standard normal distribution, $\sigma^2 = Var(X_1)$, ψ is the proportion of subjects died of the disease of interest, and ρ is the multiple correlation coefficient of the following linear regression:

$$
x_1=b_0+\boldsymbol{b}^T\boldsymbol{x}_2.
$$

That is, $\rho^2 = R^2$, where R^2 is the proportion of variance explained by the regression of X_1 on the vector of covriates X_2 .

rho will be estimated from a pilot study.

Value

Note

(1) Hsieh and Lavori (2000) assumed one-sided test, while this implementation assumed two-sided test. (2) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

References

Hsieh F.Y. and Lavori P.W. (2000). Sample-size calculation for the Cox proportional hazards regression model with nonbinary covariates. *Controlled Clinical Trials*. 21:552-560.

See Also

[powerEpiCont.default](#page-26-1)

powerEpiCont.default 27

Examples

```
# generate a toy pilot data set
set.seed(123456)
X1 \le - rnorm(100, mean = 0, sd = 0.3126)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.25, 0.75), replace = TRUE)
dat \le data.frame(X1 = X1, X2 = X2, failureFlag = failureFlag)
powerEpiCont(formula = X1 ~ ~ ~ X2,
      dat = dat,var.X1 = "X1",var.failureFlag = "failureFlag",
            n = 107,theta = exp(1),
      alpha = 0.05)
```


Description

Power calculation for Cox proportional hazards regression with nonbinary covariates for Epidemiological Studies.

Usage

```
powerEpiCont.default(n,
       theta,
       sigma2,
       psi,
       rho2,
       alpha = 0.05
```


Details

This is an implementation of the power calculation formula derived by Hsieh and Lavori (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is a nonbinary variable and X_2 is a vector of other covariates.

Suppose we want to check if the hazard ratio of the main effect $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\beta_1) = \theta$ is

$$
power = \Phi\left(-z_{1-\alpha/2} + \sqrt{n[\log(\theta)]^2 \sigma^2 \psi(1-\rho^2)}\right),\,
$$

where z_a is the 100a-th percentile of the standard normal distribution, $\sigma^2 = Var(X_1)$, ψ is the proportion of subjects died of the disease of interest, and ρ is the multiple correlation coefficient of the following linear regression:

$$
x_1=b_0+\boldsymbol{b}^T\boldsymbol{x}_2.
$$

That is, $\rho^2 = R^2$, where R^2 is the proportion of variance explained by the regression of X_1 on the vector of covriates X_2 .

Value

The power of the test.

Note

(1) Hsieh and Lavori (2000) assumed one-sided test, while this implementation assumed two-sided test. (2) The formula can be used to calculate power for a randomized trial study by setting rho2=0.

References

Hsieh F.Y. and Lavori P.W. (2000). Sample-size calculation for the Cox proportional hazards regression model with nonbinary covariates. *Controlled Clinical Trials*. 21:552-560.

See Also

[powerEpiCont](#page-24-1)

```
# example in the EXAMPLE section (page 557) of Hsieh and Lavori (2000).
# Hsieh and Lavori (2000) assumed one-sided test,
# while this implementation assumed two-sided test.
# Hence alpha=0.1 here (two-sided test) will correspond
# to alpha=0.05 of one-sided test in Hsieh and Lavori's (2000) example.
powerEpiCont.default(n = 107,
       theta = exp(1),
       sigma2 = 0.3126^2,
                    psi = 0.738,
```
powerEpiInt 29

```
rho2 = 0.1837,alpha = 0.1)
```
powerEpiInt *Power Calculation Testing Interaction Effect for Cox Proportional Hazards Regression with two covariates for Epidemiological Studies (Both covariates should be binary)*

Description

Power calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates. Some parameters will be estimated based on a pilot study.

Usage

```
powerEpiInt(X1,
     X2,
     failureFlag,
     n,
     theta,
     alpha = 0.05)
```


Details

This is an implementation of the power calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemoilogical studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),
$$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\gamma) = \theta$ is:

power =
$$
\Phi\left(-z_{1-\alpha/2} + \sqrt{\frac{n}{\delta}[\log(\theta)]^2 \psi}\right)
$$
,

where z_a is the 100a-th percentile of the standard normal distribution,

$$
\delta = \frac{1}{p_{00}} + \frac{1}{p_{01}} + \frac{1}{p_{10}} + \frac{1}{p_{11}},
$$

 ψ is the proportion of subjects died of the disease of interest, and $p_{00} = Pr(X_1 = 0, \text{and}, X_2 = 1)$ 0), $p_{01} = Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10} = Pr(X_1 = 1, \text{and}, X_2 = 0), p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 0)$ 1, and, $X_2 = 1$).

 $p_{00}, p_{01}, p_{10}, p_{11}$, and ψ will be estimated from the pilot data.

Value

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

powerEpiInt.default0 31

See Also

[powerEpiInt.default0](#page-30-1), [powerEpiInt2](#page-34-1)

Examples

```
# generate a toy pilot data set
X1 \leftarrow c(rep(1, 39), rep(0, 61))set.seed(123456)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.25, 0.75), replace = TRUE)
powerEpilnt(X1 = X1,X2 = X2,
     failureFlag = failureFlag,
     n = 184,
     theta = 3,
     alpha = 0.05)
```
powerEpiInt.default0 *Power Calculation Testing Interaction Effect for Cox Proportional Hazards Regression*

Description

Power calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
powerEpiInt.default0(n,
       theta,
       p,
       psi,
       G,
       rho2,
       alpha = 0.05
```


,

Details

This is an implementation of the power calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),
$$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\gamma) = \theta$ is

$$
power = \Phi\left(-z_{1-\alpha/2} + \sqrt{\frac{n}{G}[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)}\right)
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1), and

$$
G = \frac{[(1-q)(1-p_0)p_0 + q(1-p_1)p_1]^2}{(1-q)q(1-p_0)p_0(1-p_1)p_1}.
$$

If X_1 and X_2 are uncorrelated, we have $p_0 = p_1 = p$ leading to $1/[(1 - q)q]$. For $q = 0.5$, we have $G=4.$

Value

The power of the test.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[powerEpiInt.default1](#page-32-1), [powerEpiInt2](#page-34-1)

powerEpiInt.default1 33

Examples

```
# Example at the end of Section 4 of Schmoor et al. (2000).
powerEpiInt.default0(n = 184,
      theta = 3,
      p = 0.61,
      psi = 139 / 184,
                    G = 4.79177,rho2 = 0.015^2,
       alpha = 0.05)
```
powerEpiInt.default1 *Power Calculation Testing Interaction Effect for Cox Proportional Hazards Regression*

Description

Power calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
powerEpiInt.default1(n,
       theta,
       psi,
       p00,
       p01,
       p10,
       p11,
       alpha = 0.05)
```


Details

This is an implementation of the power calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemoilogical studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),
$$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\gamma) = \theta$ is:

power =
$$
\Phi\left(-z_{1-\alpha/2} + \sqrt{\frac{n}{\delta}[\log(\theta)]^2 \psi}\right)
$$
,

where z_a is the 100a-th percentile of the standard normal distribution,

$$
\delta = \frac{1}{p_{00}} + \frac{1}{p_{01}} + \frac{1}{p_{10}} + \frac{1}{p_{11}},
$$

 ψ is the proportion of subjects died of the disease of interest, and $p_{00} = Pr(X_1 = 0, \text{and}, X_2 = 1)$ 0), $p_{01} = Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10} = Pr(X_1 = 1, \text{and}, X_2 = 0), p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 0)$ 1, and, $X_2 = 1$).

Value

The power of the test.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[powerEpiInt.default0](#page-30-1), [powerEpiInt2](#page-34-1)

```
# Example at the end of Section 4 of Schmoor et al. (2000).
# p00, p01, p10, and p11 are calculated based on Table III on page 448
# of Schmoor et al. (2000).
powerEpiInt.default1(n = 184,
       theta = 3.
       psi = 139 / 184,
                    p00 = 50 / 184,
      p01 = 21 / 184,
       p10 = 78 / 184,
       p11 = 35 / 184,
                     alpha = 0.05
```


Description

Power calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
powerEpiInt2(n,
      theta,
      psi,
      mya,
      myb,
      myc,
      myd,
      alpha = 0.05
```
Arguments

Details

This is an implementation of the power calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemiological studies:

 $h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the power required to detect a hazard ratio as small as $\exp(\gamma) = \theta$ is

power =
$$
\Phi\left(-z_{1-\alpha/2} + \sqrt{\frac{n}{G}[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)}\right)
$$
,

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1), and

$$
G = \frac{[(1 - q)(1 - p_0)p_0 + q(1 - p_1)p_1]^2}{(1 - q)q(1 - p_0)p_0(1 - p_1)p_1}
$$

,

and $p0 = Pr(X_1 = 1 | X_2 = 0) = myc/(mya + myc)$, $p1 = Pr(X_1 = 1 | X_2 = 1) =$ $myd/(myb + myd), p = Pr(X_1 = 1) = (myc + myd)/n_{obs}, q = Pr(X_2 = 1) = (myb +$ $myd)/n_{obs}, n_{obs} = mya + myb + myc + myd.$

 p_{00} = $Pr(X_1 = 0, \text{and}, X_2 = 0), p_{01}$ = $Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10}$ = $Pr(X_1 = 0, \text{and}, X_2 = 0)$ 1, and, $X_2 = 0$, $p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 1)$.

Value

The power of the test.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[powerEpiInt.default0](#page-30-1), [powerEpiInt.default1](#page-32-1)

```
# Example at the end of Section 4 of Schmoor et al. (2000).
# mya, myb, myc, and myd are obtained from Table III on page 448
# of Schmoor et al. (2000).
powerEpiInt2(n = 184,
      theta = 3,
      psi = 139 / 184,
             mya = 50,
      myb = 21,
      myc = 78,
      myd = 35,
      alpha = 0.05)
```
ssize.stratify *Sample size calculation for Survival Analysis with Binary Predictor and Exponential Survival Function*

Description

Sample size calculation for survival analysis with binary predictor and exponential survival function.

Usage

```
ssize.stratify(
    power,
    timeUnit,
    gVec,
    PVec,
    HR,
    lambda0Vec,
    alpha = 0.05,
    verbose = TRUE)
```
Arguments

Details

We assume (1) there is only one predictor and no covariates in the survival model (exponential survival function); (2) there are m strata; (3) the predictor x is a binary variable indicating treatment group 1 ($x = 1$) or treatment group 0 ($x = 0$); (3) the treatment effect is constant over time (proportional hazards); (4) the hazard ratio is the same in all strata, and (5) the data will be analyzed by the stratified log rank test.

The sample size formula is Formula (1) on page 801 of Palta M and Amini SB (1985):

$$
n = (Z_{\alpha} + Z_{\beta})^2 / \mu^2
$$

where α is the Type I error rate, β is the Type II error rate (power= $1 - \beta$), Z_{α} is the $100(1 - \alpha)$ -th percentile of standard normal distribution, and

$$
\mu = \log(\delta) \sqrt{\sum_{s=1}^{m} g_s P_s (1 - P_s) V_s}
$$

and

$$
V_s = P_s \left[1 - \frac{1}{\lambda_{1s}} \left\{ \exp\left[-\lambda_{1s}(T-1)\right] - \exp(-\lambda_{1s}T) \right\} \right] + (1 - P_s) \left[1 - \frac{1}{\lambda_{0s}} \left\{ \exp\left[-\lambda_{0s}(T-1)\right] - \exp(-\lambda_{0s}T) \right\} \right]
$$

In the above formulas, m is the number of strata, T is the total study length, δ is the hazard ratio, g_s is the proportion of the total sample size in stratum s , P_s is the proportion of stratum s , which is in treatment group 1, and λ_{is} is the hazard for the *i*-th treatment group in stratum *s*.

Value

The sample size.

References

Palta M and Amini SB. (1985). Consideration of covariates and stratification in sample size determination for survival time studies. *Journal of Chronic Diseases*. 38(9):801-809.

See Also

[power.stratify](#page-7-1)

```
# example on page 803 of Palta M and Amini SB. (1985).
n <- ssize.stratify(
  power = 0.9,
  timeUnit = 1.25,
  gVec = c(0.5, 0.5),
  PVec = c(0.5, 0.5),
  HR = 1 / 1.91,
  lambda0Vec = c(2.303, 1.139),
  alpha = 0.05,
  verbose = TRUE
)
```


Description

Sample size calculation for the Comparison of Survival Curves Between Two Groups under the Cox Proportional-Hazards Model for clinical trials. Some parameters will be estimated based on a pilot data set.

Usage

```
ssizeCT(formula,
dat,
power,
k,
RR,
alpha = 0.05
```
Arguments

Details

This is an implementation of the sample size calculation method described in Section 14.12 (page 807) of Rosner (2006). The method was proposed by Freedman (1982).

The movitation of this function is that some times we do not have information about m or p_E and p_C available, but we have a pilot data set that can be used to estimate p_E and p_C hence m, where $m = n_{E}p_{E} + n_{C}p_{C}$ is the expected total number of events over both groups, n_{E} and n_{C} are numbers of participants in group E (experimental group) and group C (control group), respectively. p_E is the probability of failure in group E (experimental group) over the maximum time period of the study (t years). p_C is the probability of failure in group C (control group) over the maximum time period of the study (t years).

Suppose we want to compare the survival curves between an experimental group (E) and a control group (C) in a clinical trial with a maximum follow-up of t years. The Cox proportional hazards regression model is assumed to have the form:

$$
h(t|X_1) = h_0(t) \exp(\beta_1 X_1).
$$

Let n_E be the number of participants in the E group and n_C be the number of participants in the C group. We wish to test the hypothesis $H0:RR = 1$ versus $H1:RR$ not equal to 1, where $RR = \exp(\beta_1)$ =underlying hazard ratio for the E group versus the C group. Let RR be the postulated hazard ratio, α be the significance level. Assume that the test is a two-sided test. If the ratio of participants in group E compared to group $C = n_E/n_C = k$, then the number of participants needed in each group to achieve a power of $1 - \beta$ is

$$
n_E = \frac{mk}{k p_E + p_C}, n_C = \frac{m}{k p_E + p_C}
$$

where

$$
m = \frac{1}{k} \left(\frac{kRR + 1}{RR - 1} \right)^2 (z_{1 - \alpha/2} + z_{1 - \beta})^2,
$$

and $z_{1-\alpha/2}$ is the 100(1 − $\alpha/2$)-th percentile of the standard normal distribution $N(0, 1)$. p_C and p_E can be calculated from the following formulaes:

$$
p_C = \sum_{i=1}^{t} D_i, p_E = \sum_{i=1}^{t} E_i,
$$

where $D_i = \lambda_i A_i C_i$, $E_i = RR\lambda_i B_i C_i$, $A_i = \prod_{j=0}^{i-1} (1 - \lambda_j)$, $B_i = \prod_{j=0}^{i-1} (1 - RR\lambda_j)$, $C_i =$ $\prod_{j=0}^{i-1}(1-\delta_j)$. And λ_i is the probability of failure at time i among participants in the control group, given that a participant has survived to time $i - 1$ and is not censored at time $i - 1$, i.e., the approximate hazard time i in the control group, $i = 1, ..., t$; $RRlambda_i$ is the probability of failure at time i among participants in the experimental group, given that a participant has survived to time $i-1$ and is not censored at time $i-1$, i.e., the approximate hazard time i in the experimental group, $i = 1, ..., t$; delta is the probability that a participant is censored at time i given that he was followed up to time i and has not failed, $i = 0, 1, ..., t$, which is assumed the same in each group.

Value

- mat.lambda a matrix with 9 columns and nTimes+1 rows, where nTimes is the number of observed time points for the control group in the data set. The 9 columns are (1) time - observed time point for the control group; (2) lambda; (3) RR1ambda; (4) delta; (5) A; (6) B; (7) C; (8) D; (9) E. Please refer to the Details section for the definitions of elements of these quantities. See also Table 14.24 on page 809 of Rosner (2006).
- mat.event a matrix with 5 columns and nTimes+1 rows, where nTimes is the number of observed time points for control group in the data set. The 5 columns are (1) time - observed time point for the control group; (2) nEvent.C - number of events in

Note

(1) The estimates of $RRlambda_i = RR * \lambda_i$. That is, RR1 ambda is not directly estimated based on data from the experimental group; (2) The sample size formula assumes that the central-limit theorem is valid and hence is appropriate for large samples. (3) n_E and n_C will be rounded up to integers.

References

Freedman, L.S. (1982). Tables of the number of patients required in clinical trials using the log-rank test. *Statistics in Medicine*. 1: 121-129

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

See Also

[ssizeCT.default](#page-41-1)

```
# Example 14.42 in Rosner B. Fundamentals of Biostatistics.
# (6-th edition). (2006) page 809
library(survival)
data(Oph)
res <- ssizeCT(formula = Surv(times, status) ~ group,
dat = 0ph,power = 0.8,
 k = 1,
 RR = 0.7,
 alpha = 0.05)
# Table 14.24 on page 809 of Rosner (2006)
print(round(res$mat.lambda, 4))
# Table 14.12 on page 787 of Rosner (2006)
print(round(res$mat.event, 4))
# the sample size
```
print(res\$ssize)

ssizeCT.default *Sample Size Calculation in the Analysis of Survival Data for Clinical Trials*

Description

Sample size calculation for the Comparison of Survival Curves Between Two Groups under the Cox Proportional-Hazards Model for clinical trials.

Usage

```
ssizeCT.default(power,
 k,
 pE,
  pC,
  RR,
  alpha = 0.05
```
Arguments

Details

This is an implementation of the sample size calculation method described in Section 14.12 (page 807) of Rosner (2006). The method was proposed by Freedman (1982).

Suppose we want to compare the survival curves between an experimental group (E) and a control group (C) in a clinical trial with a maximum follow-up of t years. The Cox proportional hazards regression model is assumed to have the form:

$$
h(t|X_1) = h_0(t) \exp(\beta_1 X_1).
$$

Let n_E be the number of participants in the E group and n_C be the number of participants in the C group. We wish to test the hypothesis $H0:RR = 1$ versus $H1:RR$ not equal to 1, where

ssizeCT.default 43

 $RR = \exp(\beta_1)$ =underlying hazard ratio for the E group versus the C group. Let RR be the postulated hazard ratio, α be the significance level. Assume that the test is a two-sided test. If the ratio of participants in group E compared to group $C = n_E/n_C = k$, then the number of participants needed in each group to achieve a power of $1 - \beta$ is

$$
n_E = \frac{mk}{kp_E + pc}, n_C = \frac{m}{kp_E + pc}
$$

where

$$
m = \frac{1}{k} \left(\frac{kRR + 1}{RR - 1} \right)^2 (z_{1 - \alpha/2} + z_{1 - \beta})^2,
$$

and $z_{1-\alpha/2}$ is the 100(1 – $\alpha/2$)-th percentile of the standard normal distribution $N(0, 1)$.

Value

A two-element vector. The first element is n_E and the second element is n_C .

Note

(1) The sample size formula assumes that the central-limit theorem is valid and hence is appropriate for large samples. (2) n_E and n_C will be rounded up to integers.

References

Freedman, L.S. (1982). Tables of the number of patients required in clinical trials using the log-rank test. *Statistics in Medicine*. 1: 121-129

Rosner B. (2006). *Fundamentals of Biostatistics*. (6-th edition). Thomson Brooks/Cole.

See Also

[ssizeCT](#page-38-1)

```
# Example 14.42 in Rosner B. Fundamentals of Biostatistics.
# (6-th edition). (2006) page 809
ssizeCT.default(power = 0.8,
 k = 1,
 pE = 0.3707,
 pC = 0.4890,RR = 0.7,
  alpha = 0.05)
```


Description

Sample size calculation for Cox proportional hazards regression with two covariates for Epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates.

Usage

```
ssizeEpi(X1,
  X2,
  failureFlag,
  power,
  theta,
  alpha = 0.05
```
Arguments

Details

This is an implementation of the sample size formula derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 has to be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal

ssizeEpi 45

to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a power of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2}{[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1 - q)}{p(1 - p)}}
$$

,

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

 p, ρ^2 , and ψ will be estimated from a pilot study.

Value

Note

(1) The calculated sample size will be round up to an integer.

(2) The formula can be used to calculate sample size required for a randomized trial study by setting rho2=0.

(3) When rho2=0, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio $\exp(\beta_1) = \theta$ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

See Also

[ssizeEpi.default](#page-45-1)

Examples

```
# generate a toy pilot data set
X1 <- c(rep(1, 39), rep(0, 61))
set.seed(123456)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.5, 0.5), replace = TRUE)
ssizeEpi(X1 = X1,
 X2 = X2,
  failureFlag = failureFlag,
        power = 0.80,
  theta = 2,
  alpha = 0.05
```
ssizeEpi.default *Sample Size Calculation for Cox Proportional Hazards Regression*

Description

Sample size calculation for Cox proportional hazards regression with two covariates for Epidemiological Studies. The covariate of interest should be a binary variable. The other covariate can be either binary or non-binary. The formula takes into account competing risks and the correlation between the two covariates.

Usage

```
ssizeEpi.default(power,
  theta,
  p,
  psi,
  rho2,
  alpha = 0.05
```


Details

This is an implementation of the sample size formula derived by Latouche et al. (2004) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2),
$$

where the covariate X_1 is of our interest. The covariate X_1 has to be a binary variable taking two possible values: zero and one, while the covariate X_2 can be binary or continuous.

Suppose we want to check if the hazard of $X_1 = 1$ is equal to the hazard of $X_1 = 0$ or not. Equivalently, we want to check if the hazard ratio of $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a power of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2}{[\log(\theta)]^2 p(1-p)\psi(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}}
$$

,

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1).

Value

The required sample size to achieve the specified power with the given type I error rate.

Note

(1) The calculated sample size will be round up to an integer.

(2) The formula can be used to calculate sample size required for a randomized trial study by setting rho2=0.

(3) When rho2=0, the formula derived by Latouche et al. (2004) looks the same as that derived by Schoenfeld (1983). Latouche et al. (2004) pointed out that in this situation, the interpretations are different hence the two formulae are actually different. In Latouched et al. (2004), the hazard ratio $\exp(\beta_1) = \theta$ measures the difference of effect of a covariate at two different levels on the subdistribution hazard for a particular failure, while in Schoenfeld (1983), the hazard ratio θ measures the difference of effect on the cause-specific hazard.

References

Schoenfeld DA. (1983). Sample-size formula for the proportional-hazards regression model. *Biometrics*. 39:499-503.

Latouche A., Porcher R. and Chevret S. (2004). Sample size formula for proportional hazards modelling of competing risks. *Statistics in Medicine*. 23:3263-3274.

See Also

[ssizeEpi](#page-43-1)

Examples

```
# Examples at the end of Section 5.2 of Latouche et al. (2004)
# for a cohort study.
ssizeEpi.default(power = 0.80,
  theta = 2,
   p = 0.39,
  psi = 0.505,
                 rho2 = 0.132^2,
   alpha = 0.05)
```
ssizeEpiCont *Sample Size Calculation for Cox Proportional Hazards Regression with Nonbinary Covariates for Epidemiological Studies*

Description

Sample size calculation for Cox proportional hazards regression with nonbinary covariates for Epidemiological Studies.

Usage

```
ssizeEpiCont(formula,
     dat,
      var.X1,
      var.failureFlag,
     power,
      theta,
      alpha = 0.05)
```


Details

This is an implementation of the sample size calculation formula derived by Hsieh and Lavori (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2,
$$

where the covariate X_1 is a nonbinary variable and X_2 is a vector of other covariates.

Suppose we want to check if the hazard ratio of the main effect $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a sample size of $1 - \beta$ is

$$
n = \frac{\left(z_{1-\alpha/2} + z_{1-\beta}\right)^2}{[\log(\theta)]^2 \sigma^2 \psi (1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution, $\sigma^2 = Var(X_1)$, ψ is the proportion of subjects died of the disease of interest, and ρ is the multiple correlation coefficient of the following linear regression:

$$
x_1=b_0+\boldsymbol{b}^T\boldsymbol{x}_2.
$$

That is, $\rho^2 = R^2$, where R^2 is the proportion of variance explained by the regression of X_1 on the vector of covriates X_2 .

 rho^2 , σ^2 , and ψ will be estimated from a pilot study.

Value

Note

(1) Hsieh and Lavori (2000) assumed one-sided test, while this implementation assumed two-sided test. (2) The formula can be used to calculate ssize for a randomized trial study by setting rho2=0.

References

Hsieh F.Y. and Lavori P.W. (2000). Sample-size calculation for the Cox proportional hazards regression model with nonbinary covariates. *Controlled Clinical Trials*. 21:552-560.

See Also

[ssizeEpiCont.default](#page-49-1)

Examples

```
# generate a toy pilot data set
set.seed(123456)
X1 <- rnorm(100, mean = 0, sd = 0.3126)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.25, 0.75), replace = TRUE)
dat \leq data.frame(X1 = X1, X2 = X2, failureFlag = failureFlag)
ssizeEpiCont(formula = X1 ~ X2,
     dat = dat,var.X1 = "X1",
     var.failureFlag = "failureFlag",
            power = 0.806,theta = exp(1),
      alpha = 0.05)
```
ssizeEpiCont.default *Sample Size Calculation for Cox Proportional Hazards Regression with Nonbinary Covariates for Epidemiological Studies*

Description

Sample size calculation for Cox proportional hazards regression with nonbinary covariates for Epidemiological Studies.

Usage

```
ssizeEpiCont.default(power,
       theta,
       sigma2,
      psi,
       rho2,
      alpha = 0.05
```


Details

This is an implementation of the sample size calculation formula derived by Hsieh and Lavori (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2,
$$

where the covariate X_1 is a nonbinary variable and X_2 is a vector of other covariates.

Suppose we want to check if the hazard ratio of the main effect $X_1 = 1$ to $X_1 = 0$ is equal to 1 or is equal to $\exp(\beta_1) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a sample size of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2}{[\log(\theta)]^2 \sigma^2 \psi(1-\rho^2)},
$$

 α

where z_a is the 100a-th percentile of the standard normal distribution, $\sigma^2 = Var(X_1)$, ψ is the proportion of subjects died of the disease of interest, and ρ is the multiple correlation coefficient of the following linear regression:

$$
x_1=b_0+\boldsymbol{b}^T\boldsymbol{x}_2.
$$

That is, $\rho^2 = R^2$, where R^2 is the proportion of variance explained by the regression of X_1 on the vector of covriates X_2 .

Value

The total number of subjects required.

Note

(1) Hsieh and Lavori (2000) assumed one-sided test, while this implementation assumed two-sided test. (2) The formula can be used to calculate ssize for a randomized trial study by setting rho2=0.

References

Hsieh F.Y. and Lavori P.W. (2000). Sample-size calculation for the Cox proportional hazards regression model with nonbinary covariates. *Controlled Clinical Trials*. 21:552-560.

See Also

[ssizeEpiCont](#page-47-1)

```
# example in the EXAMPLE section (page 557) of Hsieh and Lavori (2000).
# Hsieh and Lavori (2000) assumed one-sided test,
# while this implementation assumed two-sided test.
# Hence alpha=0.1 here (two-sided test) will correspond
# to alpha=0.05 of one-sided test in Hsieh and Lavori's (2000) example.
ssizeEpiCont.default(power = 0.806,
       theta = exp(1),
       sigma2 = 0.3126^2,
                    psi = 0.738,
```
 $rho2 = 0.1837,$ alpha = 0.1)

ssizeEpiInt *Sample Size Calculation Testing Interaction Effect for Cox Proportional Hazards Regression*

Description

Sample size calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
ssizeEpiInt(X1,
     X2,
     failureFlag,
     power,
     theta,
     alpha = 0.05)
```
Arguments

Details

This is an implementation of the sample size calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemoilogical studies:

 $h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),$

where both covariates X_1 and X_2 are binary variables.

ssizeEpiInt 53

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve the desired power $1 - \beta$ is:

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2 G}{[\log(\theta)]^2 \psi(1-p)p(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1 - q)}{p(1 - p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1), and

$$
G = \frac{[(1-q)(1-p_0)p_0 + q(1-p_1)p_1]^2}{(1-q)q(1-p_0)p_0(1-p_1)p_1},
$$

and $p0 = Pr(X_1 = 1 | X_2 = 0) = myc/(mya + myc)$, $p1 = Pr(X_1 = 1 | X_2 = 1) =$ $myd/(myb+myd), p = Pr(X₁ = 1) = (myc+myd)/n, q = Pr(X₂ = 1) = (myb+myd)/n,$ $n = mya + myb + myc + myd.$

 p_{00} = $Pr(X_1 = 0, \text{and}, X_2 = 0), p_{01} = Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10} = Pr(X_1 = 0, \text{and}, X_2 = 1)$ 1, and, $X_2 = 0$, $p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 1)$.

 $p_{00}, p_{01}, p_{10}, p_{11}$, and ψ will be estimated from the pilot data.

Value

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[ssizeEpiInt.default0](#page-53-1), [ssizeEpiInt2](#page-57-1)

Examples

```
# generate a toy pilot data set
X1 \leftarrow c(rep(1, 39), rep(0, 61))set.seed(123456)
X2 \leq - sample(c(0, 1), 100, replace = TRUE)
failureFlag <- sample(c(0, 1), 100, prob = c(0.25, 0.75), replace = TRUE)
ssizeEpiInt(X1 = X1,
     X2 = X2,
     failureFlag = failureFlag,
     power = 0.88,
     theta = 3,
     alpha = 0.05)
```
ssizeEpiInt.default0 Sample Size Calculation Testing Interaction Effect for Cox Propor*tional Hazards Regression*

Description

Sample size calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
ssizeEpiInt.default0(power,
       theta,
       p,
       psi,
       G,
       rho2,
       alpha = 0.05
```


Details

This is an implementation of the sample size calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemiological studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),
$$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a power of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2 G}{[\log(\theta)]^2 \psi(1-p)p(1-\rho^2)},
$$

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1), and

$$
G = \frac{[(1 - q)(1 - p_0)p_0 + q(1 - p_1)p_1]^2}{(1 - q)q(1 - p_0)p_0(1 - p_1)p_1}
$$

If X_1 and X_2 are uncorrelated, we have $p_0 = p_1 = p$ leading to $1/[(1 - q)q]$. For $q = 0.5$, we have $G=4.$

Value

.

The total number of subjects required.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[ssizeEpiInt.default1](#page-55-1), [ssizeEpiInt2](#page-57-1)

Examples

```
# Example at the end of Section 4 of Schmoor et al. (2000).
ssizeEpiInt.default0(power = 0.8227,
      theta = 3,
      p = 0.61,
      psi = 139 / 184,
                    G = 4.79177,rho2 = 0.015^2,
       alpha = 0.05)
```
ssizeEpiInt.default1 *Sample Size Calculation Testing Interaction Effect for Cox Proportional Hazards Regression*

Description

Sample size calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

ssizeEpiInt.default1(power, theta, psi, p00,

```
p01,
p10,
p11,
alpha = 0.05
```


Details

This is an implementation of the sample size calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemoilogical studies:

$$
h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),
$$

where both covariates X_1 and X_2 are binary variables.

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a power of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2 \delta}{\lceil \log(\theta) \rceil^2 \psi},
$$

 Ω

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest,

$$
\delta = \frac{1}{p_{00}} + \frac{1}{p_{01}} + \frac{1}{p_{10}} + \frac{1}{p_{11}},
$$

and $p_{00} = Pr(X_1 = 0, \text{and}, X_2 = 0), p_{01} = Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10} = Pr(X_1 = 0, \text{and}, X_2 = 0)$ 1, and, $X_2 = 0$, $p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 1)$.

Value

The ssize of the test.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[ssizeEpiInt.default0](#page-53-1), [ssizeEpiInt2](#page-57-1)

```
# Example at the end of Section 4 of Schmoor et al. (2000).
# p00, p01, p10, and p11 are calculated based on Table III on page 448
# of Schmoor et al. (2000).
ssizeEpiInt.default1(power = 0.8227,
       theta = 3,
       psi = 139 / 184,
                     p00 = 50/184,
       p01 = 21 / 184,
       p10 = 78 / 184,
       p11 = 35 / 184,
                     alpha = 0.05)
```
ssizeEpiInt2 *Sample Size Calculation Testing Interaction Effect for Cox Proportional Hazards Regression*

Description

Sample size calculation testing interaction effect for Cox proportional hazards regression with two covariates for Epidemiological Studies. Both covariates should be binary variables. The formula takes into account the correlation between the two covariates.

Usage

```
ssizeEpiInt2(power,
      theta,
      psi,
      mya,
      myb,
      myc,
      myd,
      alpha = 0.05
```
Arguments

Details

This is an implementation of the sample size calculation formula derived by Schmoor et al. (2000) for the following Cox proportional hazards regression in the epidemiological studies:

 $h(t|x_1, x_2) = h_0(t) \exp(\beta_1 x_1 + \beta_2 x_2 + \gamma(x_1 x_2)),$

where both covariates X_1 and X_2 are binary variables.

ssizeEpiInt2 59

Suppose we want to check if the hazard ratio of the interaction effect $X_1X_2 = 1$ to $X_1X_2 = 0$ is equal to 1 or is equal to $\exp(\gamma) = \theta$. Given the type I error rate α for a two-sided test, the total number of subjects required to achieve a power of $1 - \beta$ is

$$
n = \frac{(z_{1-\alpha/2} + z_{1-\beta})^2 G}{[\log(\theta)]^2 \psi(1-p)p(1-\rho^2)}
$$

,

,

where z_a is the 100a-th percentile of the standard normal distribution, ψ is the proportion of subjects died of the disease of interest, and

$$
\rho = corr(X_1, X_2) = (p_1 - p_0) \times \sqrt{\frac{q(1-q)}{p(1-p)}},
$$

and $p = Pr(X_1 = 1), q = Pr(X_2 = 1), p_0 = Pr(X_1 = 1 | X_2 = 0),$ and $p_1 = Pr(X_1 = 1 | X_2 = 0)$ 1), and

$$
G = \frac{[(1 - q)(1 - p_0)p_0 + q(1 - p_1)p_1]^2}{(1 - q)q(1 - p_0)p_0(1 - p_1)p_1}
$$

and $p0 = Pr(X_1 = 1 | X_2 = 0) = myc/(mya + myc)$, $p1 = Pr(X_1 = 1 | X_2 = 1) =$ $myd/(myb+myd), p = Pr(X₁ = 1) = (myc+myd)/n, q = Pr(X₂ = 1) = (myb+myd)/n,$ $n = mya + myb + myc + myd.$

 p_{00} = $Pr(X_1 = 0, \text{and}, X_2 = 0), p_{01} = Pr(X_1 = 0, \text{and}, X_2 = 1), p_{10} = Pr(X_1 = 0, \text{and}, X_2 = 1)$ 1, and, $X_2 = 0$, $p_{11} = Pr(X_1 = 1, \text{and}, X_2 = 1)$.

Value

The total number of subjects required.

References

Schmoor C., Sauerbrei W., and Schumacher M. (2000). Sample size considerations for the evaluation of prognostic factors in survival analysis. *Statistics in Medicine*. 19:441-452.

See Also

[ssizeEpiInt.default0](#page-53-1), [ssizeEpiInt.default1](#page-55-1)

```
# Example at the end of Section 4 of Schmoor et al. (2000).
# mya, myb, myc, and myd are obtained from Table III on page 448
# of Schmoor et al. (2000).
ssizeEpiInt2(power = 0.8227,
      theta = 3,
      psi = 139 / 184,
             mya = 50,
      myb = 21,
      myc = 78.
      myd = 35,
      alpha = 0.05)
```
Index

∗ datasets Oph , [6](#page-5-0) ∗ design numDEpi , [2](#page-1-0) numDEpi.default , [4](#page-3-0) powerCT , [15](#page-14-0) powerCT.default , [18](#page-17-0) powerCT.default0 , [20](#page-19-0) powerEpi , [21](#page-20-0) powerEpi.default , [23](#page-22-0) powerEpiCont, [25](#page-24-0) powerEpiCont.default, [27](#page-26-0) powerEpiInt , [29](#page-28-0) powerEpiInt.default0 , [31](#page-30-0) powerEpiInt.default1, [33](#page-32-0) powerEpiInt2, [35](#page-34-0) ssizeCT, [39](#page-38-0) ssizeCT.default, [42](#page-41-0) ssizeEpi , [44](#page-43-0) ssizeEpi.default , [46](#page-45-0) ssizeEpiCont, [48](#page-47-0) ssizeEpiCont.default, [50](#page-49-0) ssizeEpiInt , [52](#page-51-0) ssizeEpiInt.default0 , [54](#page-53-0) ssizeEpiInt.default1, [56](#page-55-0) ssizeEpiInt2, [58](#page-57-0) ∗ method powerConLogistic.bin , [10](#page-9-0) powerConLogistic.con , [12](#page-11-0) ∗ survival numDEpi , [2](#page-1-0) numDEpi.default , [4](#page-3-0) power.stratify , [8](#page-7-0) powerCT , [15](#page-14-0) powerCT.default , [18](#page-17-0) powerCT.default0 , [20](#page-19-0) powerEpi , [21](#page-20-0) powerEpi.default , [23](#page-22-0) powerEpiCont , [25](#page-24-0)

powerEpiCont.default , [27](#page-26-0) powerEpiInt , [29](#page-28-0) powerEpiInt.default0 , [31](#page-30-0) powerEpiInt.default1 , [33](#page-32-0) powerEpiInt2 , [35](#page-34-0) ssize.stratify, [37](#page-36-0) ssizeCT, [39](#page-38-0) ssizeCT.default, [42](#page-41-0) ssizeEpi , [44](#page-43-0) ssizeEpi.default, [46](#page-45-0) ssizeEpiCont, [48](#page-47-0) ssizeEpiCont.default, [50](#page-49-0) ssizeEpiInt , [52](#page-51-0) ssizeEpiInt.default0 , [54](#page-53-0) ssizeEpiInt.default1 , [56](#page-55-0) ssizeEpiInt2 , [58](#page-57-0) numDEpi , [2](#page-1-0) , *[6](#page-5-0)* numDEpi.default , *[4](#page-3-0)* , [4](#page-3-0) Oph , [6](#page-5-0) power.stratify , [8](#page-7-0) , *[38](#page-37-0)*

powerConLogistic.bin, [10](#page-9-0) powerConLogistic.con , [12](#page-11-0) powerCT , [15](#page-14-0) , *[19](#page-18-0)* , *[21](#page-20-0)* powerCT.default , *[17](#page-16-0)* , [18](#page-17-0) , *[21](#page-20-0)* powerCT.default0 , *[17](#page-16-0)* , *[19](#page-18-0)* , [20](#page-19-0) powerEpi , [21](#page-20-0) , *[25](#page-24-0)* powerEpi.default , *[23](#page-22-0)* , [23](#page-22-0) powerEpiCont , [25](#page-24-0) , *[28](#page-27-0)* powerEpiCont.default , *[26](#page-25-0)* , [27](#page-26-0) powerEpiInt , [29](#page-28-0) powerEpiInt.default0 , *[31](#page-30-0)* , [31](#page-30-0) , *[34](#page-33-0)* , *[36](#page-35-0)* powerEpiInt.default1 , *[32](#page-31-0)* , [33](#page-32-0) , *[36](#page-35-0)* powerEpiInt2 , *[31](#page-30-0) , [32](#page-31-0)* , *[34](#page-33-0)* , [35](#page-34-0)

ssize.stratify, [9](#page-8-0), [37](#page-36-0) ssizeCT , [39](#page-38-0) , *[43](#page-42-0)* ssizeCT.default , *[41](#page-40-0)* , [42](#page-41-0)

I NDEX 61

ssizeEpi , [44](#page-43-0) , *[48](#page-47-0)* ssizeEpi.default, [45](#page-44-0), [46](#page-45-0) ssizeEpiCont , [48](#page-47-0) , *[51](#page-50-0)* ssizeEpiCont.default , *[49](#page-48-0)* , [50](#page-49-0) ssizeEpiInt , [52](#page-51-0) ssizeEpiInt.default0 , *[54](#page-53-0)* , [54](#page-53-0) , *[57](#page-56-0)* , *[59](#page-58-0)* ssizeEpiInt.default1 , *[55](#page-54-0)* , [56](#page-55-0) , *[59](#page-58-0)* ssizeEpiInt2 , *[54](#page-53-0) , [55](#page-54-0)* , *[57](#page-56-0)* , [58](#page-57-0)