Package ‘LogrankPower’

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Type Package

Title Log-Rank Test Power Calculation

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**LRPower**  

*Log-rank Test Power Calculation*

**Description**

This function is used to perform power calculation of the Log-rank test based on simulation datasets, with user specified total sample size (in one simulation), type I error, effect size, total number of simulation datasets, sample size ratio between comparison groups, the death rate in the reference group, and the distribution of follow-up time (negative binomial).

**Usage**

```r
LRPower(total.sample.size, type.I.error = 0.05, effect.size=0.6,  
simulation.n=1000, group.sample.size.ratio=1, reference.group.incidence=0.5,  
time.distribution.para=c(100, 0.8))
```

**Arguments**

- `total.sample.size`: Total sample size in this simulation dataset.
- `type.I.error`: The significance level for performing the Log-rank test. `type.I.error=0.05` by default.
- `effect.size`: A scalar parameter (between 0 and 1). `effect.size=0.2` (small effect size); `effect.size=0.5` (median effect size); `effect.size=0.8` (large effect size). `effect.size=0.6` by default.
- `simulation.n`: Total number of simulation datasets used for power calculation. `simulation.n=1000` by default.
- `group.sample.size.ratio`: Ratio between samples of contrast group and the reference or control group. `group.sample.size.ratio=1` by default.
- `reference.group.incidence`: Incidence of death or event of interest in the reference patient group. `reference.group.incidence=0.5` by default.
- `time.distribution.para`: A numeric vector of length 2. This vector is used to specify the negative binomial distribution that will be used to simulate follow-up time. The 1st element in this vector is the target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer. Then 2nd parameter is the probability of success in each trial. `time.distribution.para=c(100, 0.8)` by default.

**Details**

By default, this function will generate `simulation.n=1000` simulation datasets to calculate the power of the log-rank test at significance level of `type.I.error = 0.05`. 
The simulation datasets are generated following the steps below:

1) determine sample size of the reference patient group (group 1) by calculating
   \[
   \text{round}\left(\frac{\text{total.sample.size}}{\left(\text{group.sample.size.ratio}+1\right)}\right);
   \]
2) determine sample size of the contrast group (group 2) \(\text{total.sample.size} - \text{reference.group.n}\);
3) generate follow-up time for both the reference and the contrast group using negative bionomial
distribution with user specified parameters \(\text{time.to.event.distribution.para}\);
4) generate vital status indicator for the reference group by sampling from the bionomial distribution
with the number of trials = 1 and the probability of success/death = \(\text{reference.group.incidence}\);
5) generate vital status indicator for the contrast group by sampling from the bionomial distribution
with the number of trials = 1 and
   the probability of success/death = \(\text{reference.group.incidence} \times (1 - \text{effect.size})\);

The log-rank test p-values are calculated based on the results of \texttt{survdiff()} with \(\text{rho}=0\).

**Value**

\texttt{LRPower()} returns a point estimate of the power.

**Examples**

```r
set.seed(1234)

LRPower(total.sample.size=160, type.I.error = 0.05, effect.size=0.2,
       simulation.n=1000, group.sample.size.ratio=1, reference.group.incidence=0.5)
LRPower(total.sample.size=160, effect.size=0.5)
LRPower(total.sample.size=160, effect.size=0.8)

total.sample.size <- 160
effect.size <- c(1:20)/20
power <- sapply(effect.size, function(x){
  LRPower(total.sample.size, effect.size=x, simulation.n=100)
});
plot(effect.size, power,
    main=paste("Log-rank Test Power Calculation (n=" total.sample.size,"")
    , sep=""),
    type="b")
```

**Description**

This function is used to generate 1 simulation dataset for power calculation of the Log-rank test,
with user specified total sample size, effect size, sample size ratio between comparison groups,
the death rate in the reference group, and the distribution of follow-up time (negative binomial).
LRPowerSimulation

Usage

LRPowerSimulation(total.sample.size, effect.size=0.6, group.sample.size.ratio=1, reference.group.incidence=0.5, time.distribution.para=c(100, 0.8))

Arguments

total.sample.size
Total sample size in this simulation dataset.

effect.size
A scalar parameter (between 0 and 1). effect.size=0.2 (small effect size); effect.size=0.5 (median effect size); effect.size=0.8 (large effect size). effect.size=0.6 by default.

group.sample.size.ratio
Ratio between samples of contrast group and the reference or control group. group.sample.size.ratio=1 by default.

reference.group.incidence
The incidence of death or event of interest in the reference patient group. reference.group.incidence=0.5 by default.

time.distribution.para
A numeric vector of length 2. This vector is used to specify the negative binomial distribution that will be used to simulate follow-up time. The 1st element in this vector is the target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer. Then 2nd parameter is the probability of success in each trial. time.distribution.para=c(100, 0.8) by default.

Details

This function is used to generate one simulation dataset following the steps below:
1) determine sample size of the reference patient group (group 1) by calculating round(total.sample.size/(group.sample.size.ratio+1));
2) determine sample size of the contrast group (group 2) total.sample.size -reference.group.n;
3) generate follow-up time for both the referencee and the contrast group using negative binomial distribution with user specified parameters time.to.event.distribution.para;
4) generate vital status indicator for the reference group by sampling from the binomial distribution with the number of trials = 1 and the probability of success/death = reference.group.incidence;
5) generate vital status indicator for the contrast group by sampling from the binomial distribution with the number of trials = 1 and the probability of success/death = reference.group.incidence*(1-effect.size).

Value

LRPowerSimulation() returns a data frame of size total.sample.size by 4. The 1st column is the follow-up time. The 2nd column is the vital status (death=1). The 3rd column indicates the comparison groups. The 4th column is an object of class Surv.
Examples

```r
set.seed(1234)
surv.data <- LRPowerSimulation(total.sample.size=160, effect.size=0.5,
group.sample.size.ratio=1, reference.group.incidence=0.5)
ggsurvplot(survfit(SurvObj ~ group, surv.data), surv.median.line = "hv",
           conf.int = TRUE, pval=TRUE)
```
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