

Package: DelayedEffect.Design (via r-universe)

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Title Sample Size and Power Calculations using the APPLE, SEPPLE, APPLE+ and SEPPLE+ Methods

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Description Provides sample size and power calculations when the treatment time-lag effect is present and the lag duration is either homogeneous across the individual subject, or varies heterogeneously from individual to individual within a certain domain and following a specific pattern. The methods used are described in Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017) <doi:10.1002/sim.7157>.

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data	<i>Data for examples</i>
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Description

Data for examples.

Details

A data frame used in the examples.

Examples

```
data(data, package="DelayedEffect.Design")

# Display some of the data
data[1:5, ]
```

DelayedEffect.Design	<i>Sample size and power calculations using the APPLE, SEPPLE, APPLE+ and SEPPLE+ methods</i>
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Description

An R package for sample size and power calculation when the treatment time-lag effect is present. The package incorporates two specific lag assumptions:

1. the lag duration is homogeneous across the individual subject;
2. the lag duration varies heterogeneously from individual to individual within a certain domain and following a specific pattern.

Details

The four new methods in this package for performing the sample size and power calculations are:

1. Analytic Power calculation method based on Piecewise weighted Log-rank tEst (APPLE),
2. Simulation-based Empirical Power calculation method based on Piecewise weighted Log-rank tEst (SEPPLE),
3. Analytic Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect (APPLE+),
4. Simulation-based Empirical Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect (SEPPLE+).

See the reference for details of these methods. Specifically, APPLE and SEPPLE assume that the lag duration is homogeneous across the individual subject, whereas APPLE and SEPPLE assume that the lag duration varies heterogeneously from individual to individual or from study to study within a certain domain and following a specific pattern. The functions for computing power corresponding to the above methods are `pow.APPLE`, `pow.SEPPLE`, `pow.APPLE.plus`, `pow.SEPPLE.plus` and `pow.SEPPLE.random.DE`. These can be compared to `pow.sim.logrk` and `pow.sim.logrk.rankdom.DE` which compute the power from a simulation-based algorithm using the regular log-rank test which ignores the existence of lag effects. The package also includes the function `N.APPLE`, `N.APPLE.plus` to back calculate the sample size given the power and hazard ratio, and the functions `HR.APPLE` and `HR.APPLE.plus` to back calculate the hazard ratio given the power and sample size, respectively, using the close-from APPLE and APPLE+ methods.

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen <Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

- Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.
- Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

GPW.logrank

Generalized Piecewise Weighted Logrank Test

Description

Compute the p-value based on the generalized piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a uniform pattern.

Usage

```
GPW.logrank(data, obs.time, time.to.event, event.status, trt.group, tl, tu)
```

Arguments

<code>data</code>	Data frame
<code>obs.time</code>	Column name in data for the observational time.
<code>time.to.event</code>	Column name in data for the event time.
<code>event.status</code>	Column name in data for the event status, where 0 denotes being censored, and 1 denotes events.
<code>trt.group</code>	Column name in data for the treatment group, where 0 denotes controls, and 1 denotes treated subjects.
<code>t1</code>	Lower bound of delayed duration domain
<code>tu</code>	Upper bound of delayed duration domain

Value

The p-value of the test.

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov> , Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.SEPPLE.plus](#)

Examples

```
data(data, package="DelayedEffect.Design")
GPW.logrank(data, "X", "time_to_event", "event_status", "Z", 30, 30*11)
```

HR.APPLE

APPLE hazard ratio computation

Description

Perform the post-delay hazard ratio calculation given power and sample size using the close-form APPLE method based on the piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration is homogeneous across the individual subject

Usage

```
HR.APPLE(lambda1, t1, p, N, tao, A, beta, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard or NULL (see details)
t1	Delayed duration or NULL (see details)
p	Proportion of subjects who survive beyond the delayed period or NULL (see details)
N	Sample size
tao	Total study duration
A	Total enrollment duration
beta	Type II error rate; Power=1-beta
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE is an acronym for:

Analytic Power calculation method based on Piecewise weighted Log-rank tEst. See the reference for details of this method.

Out of the three input parameters lambda1, t1 and p, only two need to be specified, the remaining one will be computed internally from the formula $\lambda_1 = -\log(p)/t_1$. If all three are not NULL, then lambda1 will be set to $-\log(p)/t_1$ regardless of the user input value.

Value

The hazard ratio

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE](#), [N.APPLE](#)

Examples

```

lambda1 <- NULL
t1      <- 183
p       <- 0.7
N       <- 200
tao     <- 365*3
A       <- 365
beta    <- 0.2
HR.APPLE(lambda1, t1, p, N, tao, A, beta)

```

HR.APPLE.plus

APPLE+ hazard ratio computation

Description

Perform the post-delay hazard ratio calculation given power and sample size using the close-form APPLE+ method based on the generalized piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern.

Usage

```
HR.APPLE.plus(lambda1, t1, tu, N, tao, A, beta, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
N	Sample size
tao	Total study duration
A	Total enrollment duration
beta	Type II error rate; Power=1-beta
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE+ is an acronym for:

Analytic Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect. See the reference for details of this method.

Value

The hazard ratio

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE.plus](#), [N.APPLE.plus](#)

Examples

```
lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
N       <- 200
tao     <- 365*3
A       <- 365
beta    <- 0.2
HR.APPLE.plus(lambda1, t1, tu, N, tao, A, beta)
```

N.APPLE

APPLE sample size computation

Description

Perform the sample size calculation given the power and post-delay hazard ratio using the closeform APPLE method based on the piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration is homogeneous across the individual subject

Usage

```
N.APPLE(lambda1, t1, p, HR, tao, A, beta, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard or NULL (see details)
t1	Delayed duration or NULL (see details)
p	Proportion of subjects who survive beyond the delayed period or NULL (see details)
HR	Post-delay hazard ratio, defined as the post-delay hazard rate of the treatment group compared to that of the control group

tao	Total study duration
A	Total enrollment duration
beta	Type II error rate; Power=1-beta
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE is an acronym for:

Analytic Power calculation method based on Piecewise weighted Log-rank tEst. See the reference for details of this method.

Out of the three input parameters lambda1, t1 and p, only two need to be specified, the remaining one will be computed internally from the formula $\lambda_1 = -\log(p)/t_1$. If all three are not NULL, then lambda1 will be set to $-\log(p)/t_1$ regardless of the user input value.

Value

The sample size

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE](#), [HR.APPLE](#)

Examples

```
lambda1 <- NULL
t1      <- 183
p       <- 0.7
HR      <- 0.55
tao     <- 365*3
A       <- 365
beta    <- 0.2
N.APPLE(lambda1, t1, p, HR, tao, A, beta)
```

N.APPLE.plus

APPLE+ sample size computation

Description

Perform the sample size calculation given the power and post-delay hazard ratio using the close-form APPLE+ method based on the generalized piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern.

Usage

```
N.APPLE.plus(lambda1, t1, tu, HR, tao, A, beta, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
HR	Post-delay hazard ratio after tu, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
beta	Type II error rate; Power=1-beta
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE+ is an acronym for:

Analytic Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect. See the reference for details of this method.

Value

The sample size

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE.plus](#), [HR.APPLE.plus](#)

Examples

```
lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
HR      <- 1.3
tao     <- 365*3
A       <- 365
beta    <- 0.2
N.APPLE.plus(lambda1, t1, tu, HR, tao, A, beta)
```

pow.APPLE

APPLE power computation

Description

Perform the power calculation using the close-form APPLE method based on the piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration is homogeneous across the individual subject

Usage

```
pow.APPLE(lambda1, t1, p, N, HR, tao, A, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard or NULL (see details)
t1	Delayed duration or NULL (see details)
p	Proportion of subjects who survive beyond the delayed period or NULL (see details)
N	Sample size
HR	Post-delay hazard ratio, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE is an acronym for:

Analytic Power calculation method based on Piecewise weighted Log-rank tEst. See the reference for details of this method.

Out of the three input parameters λ_1 , t_1 and p , only two need to be specified, the remaining one will be computed internally from the formula $\lambda_1 = -\log(p)/t_1$. If all three are not NULL, then λ_1 will be set to $-\log(p)/t_1$ regardless of the user input value.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[N.APPLE](#), [HR.APPLE](#), [pow.SEPPLE](#), [pow.sim.logrk](#)

Examples

```
lambda1 <- NULL
t1      <- 183
p       <- 0.7
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
pow.APPLE(lambda1, t1, p, N, HR, tao, A)
```

pow.APPLE.plus

APPLE+ power computation

Description

Perform the power calculation using the close-form APPLE+ method based on the generalized piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern.

Usage

```
pow.APPLE.plus(lambda1, t1, tu, N, HR, tao, A, ap=0.5, alpha=0.05)
```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
N	Sample size
HR	Post-delay hazard ratio after tu, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.

Details

APPLE+ is an acronym for:

Analytic Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect. See the reference for details of this method.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[N.APPLE.plus](#), [HR.APPLE.plus](#)

Examples

```

lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
N       <- 200
HR      <- 1.3
tao     <- 365*3
A       <- 365
pow.APPLE.plus(lambda1, t1, tu, N, HR, tao, A)

```

pow.SEPPLE

SEPPLE power computation

Description

Perform the power calculation using the numeric SEPPLE method based on the piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration is homogeneous across the individual subject

Usage

```
pow.SEPPLE(lambda1, t1, p, N, HR, tao, A, ap=0.5, alpha=0.05, nsim=10000)
```

Arguments

lambda1	Baseline hazard or NULL (see details)
t1	Delayed duration or NULL (see details)
p	Proportion of subjects who survive beyond the delayed period or NULL (see details)
N	Sample size
HR	Post-delay hazard ratio, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.
nsim	Number of simulations. The default is 10000.

Details

SEPPLE is an acronym for:
Simulation-based Empirical Power calculation method based on Piecewise weighted Log-rank tEst.
See the reference for details of this method.

Out of the three input parameters lambda1, t1 and p, only two need to be specified, the remaining one will be computed internally from the formula $\lambda = -\log(p)/t1$. If all three are not NULL, then lambda1 will be set to $-\log(p)/t1$ regardless of the user input value.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov> , Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE](#), [pow.sim.logrk](#)

Examples

```
lambda1 <- NULL
t1      <- 183
p       <- 0.7
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
pow.SEPPLE(lambda1, t1, p, N, HR, tao, A, nsim=1000)
```

pow.SEPPLE.plus

SEPPLE+ power computation

Description

Perform the power calculation using the numeric SEPPLE+ method based on the generalized piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern.

Usage

```
pow.SEPPLE.plus(lambda1, t1, tu, N, HR, tao, A, dist="uniform",
                 shape1=NULL, shape2=NULL, ap=0.5, alpha=0.05, nsim=10000)
```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
N	Sample size
HR	Post-delay hazard ratio after tu, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
dist	One of "uniform", "beta" or "gamma", for the lag distribution
shape1	NULL or a positive parameter value for the beta or gamma distribution.
shape2	NULL or a positive parameter value for the beta or gamma distribution.
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.
nsim	Number of simulations. The default is 10000.

Details

SEPPLE+ is an acronym for:
Simulation-based Empirical Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect. See the reference for details of this method.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

- Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.
- Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.SEPPLE.random.DE](#), [pow.sim.logrk.random.DE](#)

Examples

```

lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
shape1  <- 5
shape2  <- 5
pow.SEPPLE.plus(lambda1, t1, tu, N, HR, tao, A, dist="beta",
                 shape1=shape1, shape2=shape2, nsim=1000)

```

pow.SEPPLE.random.DE *SEPPLE+ power computation*

Description

Perform the power calculation using the numeric SEPPL method based on the piecewise weighted log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern. The purpose of this function is to evaluate the property of SEPPL which assumes the lag duration is homogeneous across the individual subject, when applied under the random scenario where the lag duration, in fact, varies heterogeneously.

Usage

```

pow.SEPPLE.random.DE(lambda1, t1, tu, N, HR, tao, A, t.fixed, dist="uniform",
                     shape1=NULL, shape2=NULL, ap=0.5, alpha=0.05, nsim=10000)

```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
N	Sample size
HR	Post-delay hazard ratio after tu, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
t.fixed	Fixed duration in SEPPL
dist	One of "uniform", "beta" or "gamma", for the lag distribution
shape1	NULL or a positive parameter value for the beta or gamma distribution.
shape2	NULL or a positive parameter value for the beta or gamma distribution.
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.
nsim	Number of simulations. The default is 10000.

Details

SEPPLE+ is an acronym for:
Simulation-based Empirical Power calculation method based on generalized Piecewise weighted Log-rank tEst with random treatment time-lag effect. See the reference for details of this method.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov> , Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.SEPPLE.plus](#), [pow.sim.logrk.random.DE](#)

Examples

```
lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
t.fixed <- (t1+tu)/2
shape1  <- 5
shape2  <- 5
pow.SEPPLE.random.DE(lambda1, t1, tu, N, HR, tao, A, t.fixed, dist="beta",
                      shape1=shape1, shape2=shape2, nsim=1000)
```

pow.sim.logrk

Simulated log-rank power computation

Description

Perform the power calculation using a simulation-based method based on the regular log-rank test when the treatment time-lag effect is present and the lag duration is homogeneous across the individual subject

Usage

```
pow.sim.logrk(lambda1, t1, p, N, HR, tao, A, ap=0.5, alpha=0.05, nsim=10000)
```

Arguments

lambda1	Baseline hazard or NULL (see details)
t1	Delayed duration or NULL (see details)
p	Proportion of subjects who survive beyond the delayed period or NULL (see details)
N	Sample size
HR	Post-delay hazard ratio, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.
nsim	Number of simulations. The default is 10000.

Details

Out of the three input parameters lambda1, t1 and p, only two need to be specified, the remaining one will be computed internally from the formula $\lambda_1 = -\log(p)/t_1$. If all three are not NULL, then lambda1 will be set to $-\log(p)/t_1$ regardless of the user input value.

Value

The power

Author(s)

Zhenzhen Xu <Zhenzhen.Xu@fda.hhs.gov>, Boguang Zhen<Boguang.Zhen@fda.hhs.gov>, Yongsoek Park <yongpark@pitt.edu> and Bin Zhu <bin.zhu@nih.gov>

References

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.APPLE](#), [pow.SEPPLE](#)

Examples

```

lambda1 <- NULL
t1      <- 183
p       <- 0.7
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
pow.sim.logrk(lambda1, t1, p, N, HR, tao, A, nsim=1000)

```

```
pow.sim.logrk.random.DE
```

Simulated log-rank power computation

Description

Perform the power calculation using a simulation-based method based on the regular log-rank test when the treatment time-lag effect is present and the lag duration varies heterogeneously from individual to individual or from study to study, within a certain domain and following a specific pattern.

Usage

```
pow.sim.logrk.random.DE(lambda1, t1, tu, N, HR, tao, A, dist="uniform",
  shape1=NULL, shape2=NULL, ap=0.5, alpha=0.05, nsim=10000)
```

Arguments

lambda1	Baseline hazard
t1	Lower bound of delayed duration domain
tu	Upper bound of delayed duration domain
N	Sample size
HR	Post-delay hazard ratio after tu, defined as the post-delay hazard rate of the treatment group compared to that of the control group
tao	Total study duration
A	Total enrollment duration
dist	One of "uniform", "beta" or "gamma", for the lag distribution
shape1	NULL or a positive parameter value for the beta or gamma distribution.
shape2	NULL or a positive parameter value for the beta or gamma distribution.
ap	Experimental-control allocation ratio. The default is 0.5.
alpha	Type I error rate (two-sided). The default is 0.05.
nsim	Number of simulations. The default is 10000.

Details

The regular log-rank test is used here

Value

The power

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References

Xu, Z., Park, Y., Zhen, B. & Zhu, B. (2017). Achieving optimal power of logrank test with random treatment time-lag effect. *Biometrika*. Under review.

Xu, Z., Zhen, B., Park, Y., & Zhu, B. (2017). Designing therapeutic cancer vaccine trials with delayed treatment effect. *Statistics in medicine*, 36(4), 592-605.

See Also

[pow.SEPPLE.plus](#), [pow.SEPPLE.random.DE](#)

Examples

```
lambda1 <- 0.001982
t1      <- 30
tu      <- 30*11
N       <- 200
HR      <- 0.55
tao     <- 365*3
A       <- 365
shape1  <- 5
shape2  <- 5
pow.sim.logrk.random.DE(lambda1, t1, tu, N, HR, tao, A, dist="beta",
                        shape1=shape1, shape2=shape2, nsim=1000)
```

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