

Package ‘ReplicationSuccess’

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Type Package

Title Design and Analysis of Replication Studies

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Description Provides utilities for the design and analysis of replication studies. Features both traditional methods based on statistical significance and more recent methods such as the sceptical p-value; Held L. (2020) <[doi:10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)>. Also provides related methods including the harmonic mean chi-squared test; Held, L. (2020), <[doi:10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)>, and intrinsic credibility; Held, L. (2019) <[doi:10.1098/rsos.181534](https://doi.org/10.1098/rsos.181534)>. Contains datasets from four large-scale replication projects.

URL <https://SamCH93.github.io/ReplicationSuccess/>

BugReports <https://github.com/SamCH93/ReplicationSuccess/issues/>

Depends

VignetteBuilder knitr

Suggests knitr, roxygen2, testthat

License GPL (>= 2)

LazyData true

NeedsCompilation no

RoxygenNote 7.1.1

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ci2se	<i>Convert between estimates, z-values, p-values, and confidence intervals</i>
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Description

Convert between estimates, z-values, p-values, and confidence intervals

Usage

```
ci2se(lower, upper, conf.level = 0.95, ratio = FALSE)
```

```
ci2estimate(lower, upper, ratio = FALSE, antilog = FALSE)
```

```
ci2z(lower, upper, conf.level = 0.95, ratio = FALSE)
```

```
ci2p(
  lower,
```

```

    upper,
    conf.level = 0.95,
    ratio = FALSE,
    alternative = c("two.sided", "one.sided", "less", "greater")
)

z2p(z, alternative = c("two.sided", "one.sided", "less", "greater"))

p2z(p, alternative = c("two.sided", "one.sided", "less", "greater"))

```

Arguments

lower	Numeric vector of lower confidence interval bounds.
upper	Numeric vector of upper confidence interval bounds.
conf.level	The confidence level of the confidence intervals. Default is 0.95.
ratio	Indicates whether the confidence interval is for a ratio, e.g. an odds ratio, relative risk or hazard ratio. If TRUE, the standard error of the log ratio is computed. Defaults to FALSE.
antilog	Indicates whether the estimate is reported on the ratio scale. Only applies if ratio = TRUE. Defaults to FALSE.
alternative	Direction of the alternative of the p-value. Either "two.sided" (default), "one.sided", "less", or "greater". If "one.sided" or "two.sided" is specified, the z-value is assumed to be positive.
z	Numeric vector of z-values.
p	Numeric vector of p-values.

Details

z2p is the vectorized version of .z2p_. [Vectorize](#) is used to vectorize the function.

p2z is the vectorized version of .p2z_. [Vectorize](#) is used to vectorize the function.

Value

ci2se returns a numeric vector of standard errors.

ci2estimate returns a numeric vector of parameter estimates.

ci2z returns a numeric vector of z-values.

ci2p returns a numeric vector of p-values.

z2p returns a numeric vector of p-values.

p2z returns a numeric vector of z-values.

Examples

```

ci2se(lower = 1, upper = 3)
ci2se(lower = 1, upper = 3, ratio = TRUE)
ci2se(lower = 1, upper = 3, conf.level = 0.9)

```

```

ci2estimate(lower = 1, upper = 3)
ci2estimate(lower = 1, upper = 3, ratio = TRUE)
ci2estimate(lower = 1, upper = 3, ratio = TRUE, antilog = TRUE)

ci2z(lower = 1, upper = 3)
ci2z(lower = 1, upper = 3, ratio = TRUE)
ci2z(lower = 1, upper = 3, conf.level = 0.9)

ci2p(lower = 1, upper = 3)
ci2p(lower = 1, upper = 3, alternative = "one.sided")

z2p(z = c(1, 2, 5))
z2p(z = c(1, 2, 5), alternative = "less")
z2p(z = c(1, 2, 5), alternative = "greater")
z <- seq(-3, 3, by = 0.01)
plot(z, z2p(z), type = "l", xlab = "z", ylab = "p", ylim = c(0, 1))
lines(z, z2p(z, alternative = "greater"), lty = 2)
legend("topright", c("two-sided", "greater"), lty = c(1, 2), bty = "n")

p2z(p = c(0.005, 0.01, 0.05))
p2z(p = c(0.005, 0.01, 0.05), alternative = "greater")
p2z(p = c(0.005, 0.01, 0.05), alternative = "less")
p <- seq(0.001, 0.05, 0.0001)
plot(p, p2z(p), type = "l", ylim = c(0, 3.5), ylab = "z")
lines(p, p2z(p, alternative = "greater"), lty = 2)
legend("bottomleft", c("two-sided", "greater"), lty = c(1, 2), bty = "n")

```

effectSizeReplicationSuccess

Computes the minimum relative effect size to achieve replication success

Description

The minimum relative effect size (replication to original) to achieve replication success is computed based on the result of the original study and the corresponding variance ratio.

Usage

```

effectSizeReplicationSuccess(
  zo,
  c = 1,
  level = 0.025,
  alternative = c("one.sided", "two.sided"),
  type = c("golden", "nominal", "liberal", "controlled")
)

```

Arguments

zo	Numeric vector of z-values from original studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
level	Replication success level. Default is 0.025.
alternative	Specifies if the replication success level is "one.sided" (default) or "two.sided". If the replication success level is one-sided, then effect size calculations are based on a one-sided assessment of replication success in the direction of the original effect estimate.
type	Type of recalibration. Can be either "golden" (default), "nominal" (no recalibration), "liberal", or "controlled". "golden" ensures that for an original study just significant at the specified level, replication success is only possible for replication effect estimates larger than the original one. See levelSceptical for details about recalibration types.

Details

effectSizeReplicationSuccess is the vectorized version of .effectSizeReplicationSuccess_. [Vectorize](#) is used to vectorize the function.

Value

The minimum relative effect size to achieve replication success.

Author(s)

Leonhard Held, Charlotte Micheloud, Samuel Pawel, Florian Gerber

References

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[sampleSizeReplicationSuccess](#), [levelSceptical](#)

Examples

```
po <- c(0.001, 0.002, 0.01, 0.02, 0.025)
zo <- p2z(po, alternative = "one.sided")

effectSizeReplicationSuccess(zo = zo, c = 1, level = 0.025,
                             alternative = "one.sided", type = "golden")

effectSizeReplicationSuccess(zo = zo, c = 10, level = 0.025,
                             alternative = "one.sided", type = "golden")
```

```
effectSizeReplicationSuccess(zo = zo, c = 2, level = 0.025,
                             alternative = "one.sided", type = "nominal")

effectSizeReplicationSuccess(zo = zo, c = 2, level = 0.05,
                             alternative = "two.sided", type = "nominal")
```

effectSizeSignificance

Computes the minimum relative effect size to achieve significance of the replication study

Description

The minimum relative effect size (replication to original) to achieve significance of the replication study is computed based on the result of the original study and the corresponding variance ratio.

Usage

```
effectSizeSignificance(
  zo,
  c = 1,
  level = 0.025,
  alternative = c("one.sided", "two.sided")
)
```

Arguments

zo	Numeric vector of z-values from original studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
level	Significance level. Default is 0.025.
alternative	Specifies if the significance level is "one.sided" (default) or "two.sided".

Details

effectSizeSignificance is the vectorized version of .effectSizeSignificance_. [Vectorize](#) is used to vectorize the function.

Value

The minimum relative effect size to achieve significance in the replication study.

Author(s)

Charlotte Micheloud, Samuel Pawel, Florian Gerber

References

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[effectSizeReplicationSuccess](#)

Examples

```
po <- c(0.001, 0.002, 0.01, 0.02, 0.025)
zo <- p2z(po, alternative = "one.sided")

effectSizeSignificance(zo = zo, c = 1, level = 0.025,
                      alternative = "one.sided")

effectSizeSignificance(zo = zo, c = 1, level = 0.05,
                      alternative = "two.sided")

effectSizeSignificance(zo = zo, c = 50, level = 0.025,
                      alternative = "one.sided")
```

hMeanChiSq

harmonic mean chi-squared test

Description

p-values and confidence intervals from the harmonic mean chi-squared test.

Usage

```
hMeanChiSq(
  z,
  w = rep(1, length(z)),
  alternative = c("greater", "less", "two.sided", "none"),
  bound = FALSE
)

hMeanChiSqMu(
  thetahat,
  se,
  w = rep(1, length(thetahat)),
  mu = 0,
  alternative = c("greater", "less", "two.sided", "none"),
  bound = FALSE
)

hMeanChiSqCI(
```

```

  thetahat,
  se,
  w = rep(1, length(thetahat)),
  alternative = c("two.sided", "greater", "less", "none"),
  level = 0.95,
  wGamma = rep(1, length(unique(thetahat)) - 1)
)

```

Arguments

<code>z</code>	Numeric vector of z-values.
<code>w</code>	Numeric vector of weights.
<code>alternative</code>	Either "greater" (default), "less", "two.sided", or "none". Specifies the alternative to be considered in the computation of the p-value.
<code>bound</code>	If FALSE (default), p-values that cannot be computed are reported as NaN. If TRUE, they are reported as "> bound".
<code>thetahat</code>	Numeric vector of parameter estimates.
<code>se</code>	Numeric vector of standard errors.
<code>mu</code>	The null hypothesis value. Defaults to 0.
<code>level</code>	Numeric vector specifying the level of the confidence interval. Defaults to 0.95.
<code>wGamma</code>	Numeric vector of length <code>unique(thetahat) - 1</code> specifying weights used to summarize the gamma values, i.e., the local minima of the p-value function between the thetahats. Defaults is a vector of 1s.

Value

`hMeanChiSq`: returns the p-values from the harmonic mean chi-squared test based on the study-specific z-values.

`hMeanChiSqMu`: returns the p-value from the harmonic mean chi-squared test based on study-specific estimates and standard errors.

`hMeanChiSqCI`: returns a list containing confidence interval(s) obtained by inverting the harmonic mean chi-squared test based on study-specific estimates and standard errors. The list contains:

`CI` Confidence interval(s).

If the `alternative` is "none", the list also contains:

`gamma` Local minima of the p-value function between the thetahats.

`gammaMean` Mean of all gammas weighted by `wGamma`.

`gammaHMean` Harmonic mean of all gammas weighted by `wGamma`.

Author(s)

Leonhard Held, Florian Gerber

References

Held, L. (2020). The harmonic mean chi-squared test to substantiate scientific findings. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **69**, 697-708. doi: [10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)

Examples

```
## Example from Fisher (1999) as discussed in Held (2020)
pvalues <- c(0.0245, 0.1305, 0.00025, 0.2575, 0.128)
lower <- c(0.04, 0.21, 0.12, 0.07, 0.41)
upper <- c(1.14, 1.54, 0.60, 3.75, 1.27)
se <- ci2se(lower = lower, upper = upper, ratio = TRUE)
thetahat <- ci2estimate(lower = lower, upper = upper, ratio = TRUE)

## hMeanChiSq() -----
hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           alternative = "less")
hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           alternative = "two.sided")
hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           alternative = "none")

hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           w = 1 / se^2, alternative = "less")
hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           w = 1 / se^2, alternative = "two.sided")
hMeanChiSq(z = p2z(p = pvalues, alternative = "less"),
           w = 1 / se^2, alternative = "none")

## hMeanChiSqMu() -----
hMeanChiSqMu(thetahat = thetahat, se = se, alternative = "two.sided")
hMeanChiSqMu(thetahat = thetahat, se = se, w = 1 / se^2,
             alternative = "two.sided")
hMeanChiSqMu(thetahat = thetahat, se = se, alternative = "two.sided",
             mu = -0.1)

## hMeanChiSqCI() -----
## two-sided
CI1 <- hMeanChiSqCI(thetahat = thetahat, se = se, w = 1 / se^2,
                  alternative = "two.sided")
CI2 <- hMeanChiSqCI(thetahat = thetahat, se = se, w = 1 / se^2,
                  alternative = "two.sided", level = 0.99875)

## one-sided
CI1b <- hMeanChiSqCI(thetahat = thetahat, se = se, w = 1 / se^2,
                   alternative = "less", level = 0.975)
CI2b <- hMeanChiSqCI(thetahat = thetahat, se = se, w = 1 / se^2,
                   alternative = "less", level = 1 - 0.025^2)

## confidence intervals on hazard ratio scale
print(exp(CI1$CI), digits = 2)
print(exp(CI2$CI), digits = 2)
print(exp(CI1b$CI), digits = 2)
```

```

print(exp(CI2b$CI), digits = 2)

## example with confidence region consisting of disjunct intervals
thetahat2 <- c(-3.7, 2.1, 2.5)
se2 <- c(1.5, 2.2, 3.1)
level <- 0.95; alpha <- 1 - level
muSeq <- seq(-7, 6, length.out = 1000)
pValueSeq <- hMeanChiSqMu(thetahat = thetahat2, se = se2,
                          alternative = "none", mu = muSeq)
(hm <- hMeanChiSqCI(thetahat = thetahat2, se = se2, alternative = "none"))

plot(x = muSeq, y = pValueSeq, type = "l", panel.first = grid(lty = 1),
     xlab = expression(mu), ylab = "p-value")
abline(v = thetahat2, h = alpha, lty = 2)
arrows(x0 = hm$CI[, 1], x1 = hm$CI[, 2], y0 = alpha,
       y1 = alpha, col = "darkgreen", lwd = 3, angle = 90, code = 3)
points(hm$gamma, col = "red", pch = 19, cex = 2)

```

levelEquivalent

Computes the level for the sceptical p-value

Description

The equivalent level for the sceptical p-value is computed based on the specified limiting relative effect size, the replication success level, and the alternative.

Usage

```
levelEquivalent(dinf, level = 0.025, alternative = c("one.sided", "two.sided"))
```

Arguments

dinf	The limiting relative effect size bound below which replication success is deemed to be impossible for borderline significant original studies with p-values equal to level.
level	Numeric vector of required replication success levels. Defaults to 0.025.
alternative	Either "one.sided" (default) or "two.sided". Specifies if the replication success level is one-sided or two-sided. If the replication success level is one-sided, then a one-sided level for the sceptical p-value is computed.

Value

Equivalent levels for the sceptical p-value corresponding to $d_{min}=1$. Thus, at the equivalent level, replication success is impossible for borderline significant original studies and shrinkage of the replication effect estimate.

Author(s)

Leonhard Held

References

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[pSceptical](#), [levelSceptical](#)

Examples

```
levelEquivalent(dinf = 0.8, level = 0.025)
levelEquivalent(dinf = 0.8, level = 0.05, alternative="two.sided")
```

levelSceptical	<i>Computes the level for the sceptical p-value</i>
----------------	---

Description

The level for the sceptical p-value is computed based on the specified alternative and calibration type.

Usage

```
levelSceptical(
  level,
  alternative = c("one.sided", "two.sided"),
  type = c("golden", "nominal", "liberal", "controlled")
)
```

Arguments

level	Numeric vector of required replication success levels.
alternative	Either "one.sided" (default) or "two.sided". Specifies if the replication success level is one-sided or two-sided. If the replication success level is one-sided, then a one-sided level for the sceptical p-value is computed.
type	The calibration type can be either "golden" (default), "nominal", "liberal", or "controlled". type = "golden" ensures that for an original study just significant at the specified level, replication success is only possible if the replication effect estimate is larger than the original one. If type = "controlled", the type-I

error rate is equal to level^2 (for `alternative = "two.sided"`) or $2 \times \text{level}^2$ (for `alternative = "one.sided"`) if the variance ratio is equal to 1. The type `"nominal"` ensures that the type-I error rate is always smaller than level^2 . Significance of both the original and replication study at `level` is then a necessary but not sufficient requirement for replication success. If type is `"liberal"` then significance of both studies is a sufficient requirement for replication success if the variance ratio is equal to 1.

Value

Levels for the sceptical p-value.

Author(s)

Leonhard Held

References

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Held, L. (2020). The harmonic mean chi-squared test to substantiate scientific findings. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **69**, 697-708. doi: [10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

Examples

```
levelSceptical(level = 0.025, alternative = "one.sided", type = "nominal")
levelSceptical(level = 0.025, alternative = "one.sided", type = "liberal")
levelSceptical(level = 0.025, alternative = "one.sided", type = "controlled")
levelSceptical(level = 0.025, alternative = "one.sided", type = "golden")
```

pBox

Computes Box's tail probability

Description

pBox computes Box's tail probabilities based on the z-values of the original and the replication study, the corresponding variance ratio, and the significance level.

Usage

```
pBox(zo, zr, c, level = 0.05, alternative = c("two.sided", "one.sided"))
zBox(zo, zr, c, level = 0.05, alternative = c("two.sided", "one.sided"))
```

Arguments

zo	Numeric vector of z-values from the original studies.
zr	Numeric vector of z-values from replication studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
level	Numeric vector of significance levels. Default is 0.05.
alternative	Either "two.sided" (default) or "one.sided". Specifies whether two-sided or one-sided Box's tail probabilities are computed.

Value

pBox returns Box's tail probabilities.

zBox returns the z-values used in pBox.

Author(s)

Leonhard Held

References

Box, G.E.P. (1980). Sampling and Bayes' inference in scientific modelling and robustness (with discussion). *Journal of the Royal Statistical Society, Series A*, **143**, 383-430.

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Examples

```
pBox(zo = p2z(0.01), zr = p2z(0.02), c = 2)
pBox(zo = p2z(0.02), zr = p2z(0.01), c = 1/2)
pBox(zo = p2z(0.02, alternative = "one.sided"),
      zr = p2z(0.01, alternative = "one.sided"),
      c = 1/2, alternative = "one.sided")
```

pIntrinsic

Computes the p-value for intrinsic credibility

Description

Computes the p-value for intrinsic credibility

Usage

```
pIntrinsic(  
  p = z2p(z, alternative = alternative),  
  z = NULL,  
  alternative = c("two.sided", "one.sided"),  
  type = c("Held", "Matthews")  
)
```

Arguments

<code>p</code>	numeric vector of p-values.
<code>z</code>	numeric vector of z-values. Default is NULL.
<code>alternative</code>	Either "two.sided" (default) or "one.sided". Specifies if the p-value is two-sided or one-sided. If the p-value is one-sided, then a one-sided p-value for intrinsic credibility is computed.
<code>type</code>	Type of intrinsic p-value. Default is "Held" as in Held (2019). The other option is "Matthews" as in Matthews (2018).

Value

p-values for intrinsic credibility.

Author(s)

Leonhard Held

References

Matthews, R. A. J. (2018). Beyond 'significance': principles and practice of the analysis of credibility. *Royal Society Open Science*, **5**, 171047. doi: [10.1098/rsos.171047](https://doi.org/10.1098/rsos.171047)

Held, L. (2019). The assessment of intrinsic credibility and a new argument for $p < 0.005$. *Royal Society Open Science*, **6**, 181534. doi: [10.1098/rsos.181534](https://doi.org/10.1098/rsos.181534)

Examples

```
p <- c(0.005, 0.01, 0.05)  
pIntrinsic(p = p)  
pIntrinsic(p = p, type = "Matthews")  
pIntrinsic(p = p, alternative = "one.sided")  
pIntrinsic(p = p, alternative = "one.sided", type = "Matthews")  
  
pIntrinsic(z = 2)
```

 powerReplicationSuccess

Computes the power for replication success

Description

Computes the power for replication success based on the result of the original study, the corresponding variance ratio, and the design prior.

Usage

```
powerReplicationSuccess(
  zo,
  c = 1,
  level = 0.025,
  designPrior = c("conditional", "predictive", "EB"),
  alternative = c("one.sided", "two.sided"),
  type = c("golden", "nominal", "liberal", "controlled"),
  shrinkage = 0,
  h = 0,
  strict = FALSE
)
```

Arguments

zo	Numeric vector of z-values from original studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
level	Numeric vector of replication success levels. The default is 0.025.
designPrior	Either "conditional" (default), "predictive", or "EB". If "EB", the power is computed under a predictive distribution, where the contribution of the original study is shrunken towards zero based on the evidence in the original study (with an empirical Bayes shrinkage estimator).
alternative	Either "one.sided" (default) or "two.sided". Specifies if the replication success level is one-sided or two-sided. If the replication success level is one-sided, then power calculations are based on a one-sided assessment of replication success in the direction of the original effect estimates.
type	Recalibration type can be either "golden" (default), "nominal" (no recalibration), "liberal", or "controlled". <code>type = "golden"</code> ensures that for an original study just significant at the specified <code>level</code> , replication success is only possible if the replication effect estimate is larger than the original one. See levelSceptical for details about recalibration types.

shrinkage	Numeric vector with values in [0,1). Defaults to 0. Specifies the shrinkage of the original effect estimate towards zero, e.g., the effect is shrunken by a factor of 25% for shrinkage = 0.25. Is only taken into account if the designPrior is "conditional" or "predictive".
h	Numeric vector of relative heterogeneity variances i.e., the ratio of the heterogeneity variance to the variance of the original effect estimate. Default is 0 (no heterogeneity). Is only taken into account when designPrior = "predictive" or designPrior = "EB".
strict	Logical vector indicating whether the probability for replication success in the opposite direction of the original effect estimate should also be taken into account. Default is FALSE. Only taken into account when alternative = "two.sided".

Details

powerReplicationSuccess is the vectorized version of .powerReplicationSuccess_. [Vectorize](#) is used to vectorize the function.

Value

The power for replication success.

Author(s)

Leonhard Held, Charlotte Micheloud, Samuel Pawel

References

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[sampleSizeReplicationSuccess](#), [pSceptical](#), [levelSceptical](#)

Examples

```
## larger sample size in replication (c > 1)
powerReplicationSuccess(z0 = p2z(0.005), c = 2, level = 0.025, designPrior = "conditional")
powerReplicationSuccess(z0 = p2z(0.005), c = 2, level = 0.025, designPrior = "predictive")

## smaller sample size in replication (c < 1)
powerReplicationSuccess(z0 = p2z(0.005), c = 1/2, level = 0.025, designPrior = "conditional")
powerReplicationSuccess(z0 = p2z(0.005), c = 1/2, level = 0.025, designPrior = "predictive")

powerReplicationSuccess(z0 = p2z(0.00005), c = 2, level = 0.05,
                        alternative = "two.sided", strict = TRUE, shrinkage = 0.9)
powerReplicationSuccess(z0 = p2z(0.00005), c = 2, level = 0.05,
```



```
alternative = "two.sided", strict = FALSE, shrinkage = 0.9)
```

powerSignificance *Computes the power for significance*

Description

The power for significance is computed based on the result of the original study, the corresponding variance ratio, and the design prior.

Usage

```
powerSignificance(
  zo,
  c = 1,
  level = 0.025,
  designPrior = c("conditional", "predictive", "EB"),
  alternative = c("one.sided", "two.sided"),
  h = 0,
  shrinkage = 0,
  strict = FALSE
)
```

Arguments

zo	Numeric vector of z-values from original studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
level	Numeric vector of significance levels. The default is 0.025.
designPrior	Either "conditional" (default), "predictive", or "EB". If "EB", the power is computed under a predictive distribution, where the contribution of the original study is shrunken towards zero based on the evidence in the original study (with an empirical Bayes shrinkage estimator).
alternative	Either "one.sided" (default) or "two.sided". Specifies if the significance level is one-sided or two-sided. If the significance level is one-sided, then power calculations are based on a one-sided assessment of significance in the direction of the original effect estimates.
h	The relative between-study heterogeneity, i.e., the ratio of the heterogeneity variance to the variance of the original effect estimate. Default is 0 (no heterogeneity). Is only taken into account when designPrior = "predictive" or designPrior = "EB".
shrinkage	Numeric vector with values in [0,1). Defaults to 0. Specifies the shrinkage of the original effect estimate towards zero, e.g., the effect is shrunken by a factor of 25% for shrinkage = 0.25. Is only taken into account if the designPrior is "conditional" or "predictive".

strict Logical vector indicating whether the probability for significance in the opposite direction of the original effect estimate should also be taken into account. Default is FALSE. Only taken into account when `alternative = "two.sided"`.

Details

`powerSignificance` is the vectorized version of `.powerSignificance_`. [Vectorize](#) is used to vectorize the function.

Value

The probability that a replication study yields a significant effect estimate in the specified direction.

Author(s)

Leonhard Held, Samuel Pawel, Charlotte Micheloud, Florian Gerber

References

- Goodman, S. N. (1992). A comment on replication, p-values and evidence, *Statistics in Medicine*, **11**, 875–879. doi: [10.1002/sim.4780110705](https://doi.org/10.1002/sim.4780110705)
- Senn, S. (2002). Letter to the Editor, *Statistics in Medicine*, **21**, 2437–2444.
- Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)
- Pawel, S., Held, L. (2020). Probabilistic forecasting of replication studies. *PLoS ONE*. **15**, e0231416. doi: [10.1371/journal.pone.0231416](https://doi.org/10.1371/journal.pone.0231416)
- Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[sampleSizeSignificance](#), [powerSignificanceInterim](#)

Examples

```
powerSignificance(zo = p2z(0.005), c = 2)
powerSignificance(zo = p2z(0.005), c = 2, designPrior = "predictive")
powerSignificance(zo = p2z(0.005), c = 2, alternative = "two.sided")
powerSignificance(zo = -3, c = 2, designPrior = "predictive",
                  alternative = "one.sided")
powerSignificance(zo = p2z(0.005), c = 1/2)
powerSignificance(zo = p2z(0.005), c = 1/2, designPrior = "predictive")
powerSignificance(zo = p2z(0.005), c = 1/2, alternative = "two.sided")
powerSignificance(zo = p2z(0.005), c = 1/2, designPrior = "predictive",
                  alternative = "two.sided")
powerSignificance(zo = p2z(0.005), c = 1/2, designPrior = "predictive",
                  alternative = "one.sided", h = 0.5, shrinkage = 0.5)
powerSignificance(zo = p2z(0.005), c = 1/2, designPrior = "EB",
                  alternative = "two.sided", h = 0.5)
```

```
# power as function of original p-value
po <- seq(0.0001, 0.06, 0.0001)
plot(po, powerSignificance(zo = p2z(po), designPrior = "conditional"),
      type = "l", ylim = c(0, 1), lwd = 1.5, las = 1, ylab = "Power",
      xlab = expression(italic(p)[o]))
lines(po, powerSignificance(zo = p2z(po), designPrior = "predictive"),
      lwd = 2, lty = 2)
lines(po, powerSignificance(zo = p2z(po), designPrior = "EB"),
      lwd = 1.5, lty = 3)
legend("topright", legend = c("conditional", "predictive", "EB"),
      title = "Design prior", lty = c(1, 2, 3), lwd = 1.5, bty = "n")
```

powerSignificanceInterim

Interim power of a replication study

Description

Computes the power of a replication study taking into account data from an interim analysis.

Usage

```
powerSignificanceInterim(
  zo,
  zi,
  c = 1,
  f = 1/2,
  level = 0.025,
  designPrior = c("conditional", "informed predictive", "predictive"),
  analysisPrior = c("flat", "original"),
  alternative = c("one.sided", "two.sided"),
  shrinkage = 0
)
```

Arguments

zo	Numeric vector of z-values from original studies.
zi	Numeric vector of z-values from interim analyses of replication studies.
c	Ratio of the sample size of the replication study to the sample size of the original study. Default is 1.
f	Fraction of the replication study already completed. Default is 0.5.
level	Significance level. Default is 0.025.
designPrior	Either "conditional" (default), "informed predictive", or "predictive". "informed predictive" refers to an informative normal prior coming from the original study. "predictive" refers to a flat prior.

analysisPrior	Either "flat" (default) or "original".
alternative	Either "one.sided" (default) or "two.sided". Specifies if the significance level is one-sided or two-sided.
shrinkage	Numeric vector with values in [0,1). Defaults to 0. Specifies the shrinkage of the original effect estimate towards zero, e.g., the effect is shrunken by a factor of 25% for shrinkage=0.25.

Details

This is an extension of `powerSignificance()` and adapts the ‘interim power’ from section 6.6.3 of Spiegelhalter et al. (2004) to the setting of replication studies.

`powerSignificanceInterim` is the vectorized version of `.powerSignificanceInterim_`. [Vectorize](#) is used to vectorize the function.

Value

The probability of statistical significance in the specified direction at the end of the replication study given the data collected so far in the replication study.

Author(s)

Charlotte Micheloud

References

Spiegelhalter, D. J., Abrams, K. R., and Myles, J. P. (2004). Bayesian Approaches to Clinical Trials and Health-Care Evaluation, volume 13. John Wiley & Sons

Micheloud, C., Held, L. (2021). Power Calculations for Replication Studies. <https://arxiv.org/abs/2004.10814>

See Also

[sampleSizeSignificance](#), [powerSignificance](#)

Examples

```
powerSignificanceInterim(zo = 2, zi = 2, c = 1, f = 1/2,
                        designPrior = "conditional",
                        analysisPrior = "flat")
```

```
powerSignificanceInterim(zo = 2, zi = 2, c = 1, f = 1/2,
                        designPrior = "informed predictive",
                        analysisPrior = "flat")
```

```
powerSignificanceInterim(zo = 2, zi = 2, c = 1, f = 1/2,
                        designPrior = "predictive",
                        analysisPrior = "flat")
```

```
powerSignificanceInterim(zo = 2, zi = -2, c = 1, f = 1/2,
                        designPrior = "conditional",
```

```

analysisPrior = "flat")

powerSignificanceInterim(zo = 2, zi = 2, c = 1, f = 1/2,
  designPrior = "conditional",
  analysisPrior = "flat",
  shrinkage = 0.25)

```

PPpSceptical

Compute project power of the sceptical p-value

Description

The project power of the sceptical p-value is computed for a specified level of replication success, the relative variance, significance level and power for a standard significance test of the original study, and the alternative hypothesis.

Usage

```

PPpSceptical(
  level,
  c,
  alpha,
  power,
  alternative = c("one.sided", "two.sided", "greater", "less"),
  type = c("golden", "nominal", "liberal", "controlled")
)

```

Arguments

level	Numeric vector of levels of replication success.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
alpha	Significance level for a standard significance test in the original study.
power	Power to detect the assumed effect with a standard significance test in the original study.
alternative	Either "one.sided" (default), "two.sided", "greater", or "less". If "one.sided", the type-I error rate is computed based on a one-sided assessment of replication success in the direction of the original effect estimate. If "two.sided", the type-I error rate is computed based on a two-sided assessment of replication success regardless of the direction of the original and replication effect estimate. If "greater" or "less", the type-I error rate is computed based on a one-sided assessment of replication success in the pre-specified direction of the original and replication effect estimate.

type Type of recalibration. Can be either "golden" (default), "nominal" (no recalibration), "liberal", or "controlled". "golden" ensures that for an original study just significant at the specified level, replication success is only possible if the replication effect estimate is at least as large as the original one. See [levelSceptical](#) for details about recalibration types.

Details

PPpSceptical is the vectorized version of `.PPpSceptical_`. [Vectorize](#) is used to vectorize the function.

Value

The project power.

Author(s)

Samuel Pawel, Leonhard Held

References

Held, L. (2020). The harmonic mean chi-squared test to substantiate scientific findings. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **69**, 697-708. doi: [10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

Maca, J., Gallo, P., Branson, M., and Maurer, W. (2002). Reconsidering some aspects of the two-trials paradigm. *Journal of Biopharmaceutical Statistics*, **12**, 107-119. doi: [10.1081/bip120006450](https://doi.org/10.1081/bip120006450)

See Also

[pSceptical](#), [levelSceptical](#), [T1EpSceptical](#)

Examples

```
## compare project power for different levels of replication success
levels <- c("nominal" = levelSceptical(level = 0.025, type = "nominal"),
           "liberal" = levelSceptical(level = 0.025, type = "liberal"),
           "controlled" = levelSceptical(level = 0.025, type = "controlled"),
           "golden" = levelSceptical(level = 0.025, type = "golden"))
c <- seq(0.4, 5, by = 0.01)
alpha <- 0.025
power <- 0.9
pp <- sapply(X = levels, FUN = function(l) {
  PPpSceptical(level = l, c = c, alpha, power, alternative = "one.sided",
               type = "nominal")
})

## compute project power of 2 trials rule
za <- qnorm(p = 1 - alpha)
mu <- za + qnorm(p = power)
pp2TR <- power*pnorm(q = za, mean = sqrt(c)*mu, lower.tail = FALSE)
```

```

matplot(x = c, y = pp*100, type = "l", lty = 1, lwd = 2, las = 1, log = "x",
        xlab = bquote(italic(c)), ylab = "Project power (%)", xlim = c(0.4, 5),
        ylim = c(0, 100))
lines(x = c, y = pp2TR*100, col = length(levels) + 1, lwd = 2)
abline(v = 1, lty = 2)
abline(h = 90, lty = 2, col = "lightgrey")
legend("bottomright", legend = c(names(levels), "2TR"), lty = 1, lwd = 2,
       col = seq(1, length(levels) + 1))

```

predictionInterval *Prediction interval for effect estimate of replication study*

Description

Computes a prediction interval for the effect estimate of the replication study.

Usage

```

predictionInterval(
  thetao,
  seo,
  ser,
  tau = 0,
  conf.level = 0.95,
  designPrior = "predictive"
)

```

Arguments

thetao	Numeric vector of effect estimates from original studies.
seo	Numeric vector of standard errors of the original effect estimates.
ser	Numeric vector of standard errors of the replication effect estimates.
tau	Between-study heterogeneity standard error. Default is 0 (no heterogeneity). Is only taken into account when designPrior is "predictive" or "EB".
conf.level	The confidence level of the prediction intervals. Default is 0.95.
designPrior	Either "predictive" (default), "conditional", or "EB". If "EB", the contribution of the original study to the predictive distribution is shrunken towards zero based on the evidence in the original study (with empirical Bayes).

Details

This function computes a prediction interval and a mean estimate under a specified predictive distribution of the replication effect estimate. Setting designPrior = "conditional" is not recommended since this ignores the uncertainty of the original effect estimate. See Patil, Peng, and Leek (2016) and Pawel and Held (2020) for details.

predictionInterval is the vectorized version of .predictionInterval_. [Vectorize](#) is used to vectorize the function.

Value

A data frame with the following columns

lower	Lower limit of prediction interval,
mean	Mean of predictive distribution,
upper	Upper limit of prediction interval.

Author(s)

Samuel Pawel

References

Patil, P., Peng, R. D., Leek, J. T. (2016). What should researchers expect when they replicate studies? A statistical view of replicability in psychological science. *Perspectives on Psychological Science*, **11**, 539-544. doi: [10.1177/1745691616646366](https://doi.org/10.1177/1745691616646366)

Pawel, S., Held, L. (2020). Probabilistic forecasting of replication studies. *PLoS ONE*. **15**, e0231416. doi: [10.1371/journal.pone.0231416](https://doi.org/10.1371/journal.pone.0231416)

Examples

```
predictionInterval(thetao = c(1.5, 2, 5), seo = 1, ser = 0.5, designPrior = "EB")

# compute prediction intervals for replication projects
data("RProjects", package = "ReplicationSuccess")
parOld <- par(mfrow = c(2, 2))
for (p in unique(RProjects$project)) {
  data_project <- subset(RProjects, project == p)
  PI <- predictionInterval(thetao = data_project$fiso, seo = data_project$se_fiso,
                          ser = data_project$se_fisr)
  PI <- tanh(PI) # transforming back to correlation scale
  within <- (data_project$rr < PI$upper) & (data_project$rr > PI$lower)
  coverage <- mean(within)
  color <- ifelse(within == TRUE, "#333333B3", "#8B0000B3")
  study <- seq(1, nrow(data_project))
  plot(data_project$rr, study, col = color, pch = 20,
        xlim = c(-0.5, 1), xlab = expression(italic(r)[r]),
        main = paste0(p, ": ", round(coverage*100, 1), "% coverage"))
  arrows(PI$lower, study, PI$upper, study, length = 0.02, angle = 90,
        code = 3, col = color)
  abline(v = 0, lty = 3)
}
par(parOld)
```

pReplicate	<i>Probability of replicating an effect of Killeen (2005)</i>
------------	---

Description

Computes the probability that a replication study yields an effect estimate in the same direction as in the original study.

Usage

```
pReplicate(  
  po = NULL,  
  zo = p2z(p = po, alternative = alternative),  
  c,  
  alternative = "two.sided"  
)
```

Arguments

po	Numeric vector of p-values from the original study, default is NULL.
zo	Numeric vector of z-values from the original study. Is calculated from po, if necessary.
c	The ratio of the variances of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
alternative	Either "two.sided" (default) or "one.sided". Specifies whether the p-value is two-sided or one-sided.

Details

This extends the statistic `p_rep` ("the probability of replicating an effect") by Killeen (2005) to the case of possibly unequal sample sizes, see also Senn (2002).

Value

The probability that a replication study yields an effect estimate in the same direction as in the original study.

Author(s)

Leonhard Held

References

- Killeen, P. R. (2005). An alternative to null-hypothesis significance tests. *Psychological Science*, **16**, 345–353. doi: [10.1111/j.09567976.2005.01538.x](https://doi.org/10.1111/j.09567976.2005.01538.x)
- Senn, S. (2002). Letter to the Editor, *Statistics in Medicine*, **21**, 2437–2444.
- Held, L. (2019). The assessment of intrinsic credibility and a new argument for $p < 0.005$. *Royal Society Open Science*, **6**, 181534. doi: [10.1098/rsos.181534](https://doi.org/10.1098/rsos.181534)

Examples

```
pReplicate(po = c(0.05, 0.01, 0.001), c = 1)
pReplicate(po = c(0.05, 0.01, 0.001), c = 2)
pReplicate(po = c(0.05, 0.01, 0.001), c = 2, alternative = "one.sided")
pReplicate(zo = c(2, 3, 4), c = 1)
```

protzko2020

Data from Protzko et al. (2020)

Description

Data from "High Replicability of Newly-Discovered Social-behavioral Findings is Achievable" by Protzko et al. (2020). The variables are as follows:

experiment Experiment name
 type Type of study, either "original", "self-replication", or "external-replication"
 lab The lab which conducted the study, either 1, 2, 3, or 4.
 smd Standardized mean difference effect estimate
 se Standard error of standardized mean difference effect estimate
 n Total sample size of the study

Usage

```
data("protzko2020")
```

Format

A data frame with 80 rows and 6 variables

Details

This data set originates from a prospective replication project involving four laboratories. Each of them conducted four original studies and for each original study a replication study was carried out within the same lab (self-replication) and by the other three labs (external-replication). Most studies used simple between-subject designs with two groups and a continuous outcome so that for each study, an estimate of the standardized mean difference (SMD) could be computed from the group means, group standard deviations, and group sample sizes. For studies with covariate adjustment and/or binary outcomes, effect size transformations as described in the supplementary material of

Protzko (2020) were used to obtain effect estimates and standard errors on SMD scale. The data set is licensed under a CC-BY Attribution 4.0 International license, see <https://creativecommons.org/licenses/by/4.0/> for the terms of reuse.

Source

The relevant files were downloaded from <https://osf.io/42ef9/> on January 24, 2022. The R markdown script "Decline effects main analysis.Rmd" was executed and the relevant variables from the objects "ES_experiments" and "decline_effects" were saved.

References

Protzko, J., Krosnick, J., Nelson, L. D., Nosek, B. A., Axt, J., Berent, M., ... Schooler, J. (2020, September 10). High Replicability of Newly-Discovered Social-behavioral Findings is Achievable. doi: [10.31234/osf.io/n2a9x](https://doi.org/10.31234/osf.io/n2a9x)

Protzko, J., Berent, M., Buttrick, N., DeBell, M., Roeder, S. S., Walleczek, J., ... Nosek, B. A. (2021, January 5). Results & Data. Retrieved from <https://osf.io/42ef9/>

Examples

```
data("protzko2020", package = "ReplicationSuccess")

## forestplots of effect estimates
graphics.off()
parOld <- par(mar = c(5, 8, 4, 2), mfrow = c(4, 4))
experiments <- unique(protzko2020$experiment)
for (ex in experiments) {
  ## compute CIs
  dat <- subset(protzko2020, experiment == ex)
  za <- qnorm(p = 0.975)
  plotDF <- data.frame(lower = dat$smd - za*dat$se,
                      est = dat$smd,
                      upper = dat$smd + za*dat$se)
  colpalette <- c("#000000", "#1B9E77", "#D95F02")
  cols <- colpalette[dat$type]
  yseq <- seq(1, nrow(dat))

  ## forestplot
  plot(x = plotDF$est, y = yseq, xlim = c(-0.15, 0.8),
       ylim = c(0.8*min(yseq), 1.05*max(yseq)), type = "n",
       yaxt = "n", xlab = "Effect estimate (SMD)", ylab = "")
  abline(v = 0, col = "#0000004D")
  arrows(x0 = plotDF$lower, x1 = plotDF$upper, y0 = yseq, angle = 90,
        code = 3, length = 0.05, col = cols)
  points(y = yseq, x = plotDF$est, pch = 20, lwd = 2, col = cols)
  axis(side = 2, at = yseq, las = 1, labels = dat$type, cex.axis = 0.85)
  title(main = ex)
}
par(parOld)
```

pSceptical

Computes the sceptical p-value and z-value

Description

Computes sceptical p-values and z-values based on the z-values of the original and the replication study and the corresponding variance ratio. If specified, the p-values are recalibrated.

Usage

```
pSceptical(
  zo,
  zr,
  c,
  alternative = c("one.sided", "two.sided"),
  type = c("golden", "nominal", "liberal", "controlled")
)

zSceptical(zo, zr, c)
```

Arguments

zo	Numeric vector of z-values from original studies.
zr	Numeric vector of z-values from replication studies.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.
alternative	Either "one.sided" (default) or "two.sided". If "one.sided", the sceptical p-value is based on a one-sided assessment of replication success in the direction of the original effect estimate. If "two.sided", the sceptical p-value is based on a two-sided assessment of replication success regardless of the direction of the original and replication effect estimate.
type	Type of recalibration. Can be either "golden" (default), "nominal", "liberal", or "controlled". Setting type to "nominal" corresponds to no recalibration as in Held et al. (2020). A recalibration is applied if type is "liberal", "controlled", or "golden", and the sceptical p-value can then be interpreted on the same scale as an ordinary p-value (e.g., a one-sided sceptical p-value can be thresholded at the conventional 0.025 level). See levelSceptical for details about recalibration types.

Details

pSceptical is the vectorized version of .pSceptical_. [Vectorize](#) is used to vectorize the function.

Value

pSceptical returns the sceptical p-value.

zSceptical returns the z-value of the sceptical p-value.

Author(s)

Leonhard Held

References

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[sampleSizeReplicationSuccess](#), [powerReplicationSuccess](#), [levelSceptical](#)

Examples

```
## no recalibration (type = "nominal") as in Held (2020)
pSceptical(zo = p2z(0.01), zr = p2z(0.02), c = 2, alternative = "one.sided",
           type = "nominal")

## recalibration with golden level as in Held, Micheloud, Pawel (2020)
pSceptical(zo = p2z(0.01), zr = p2z(0.02), c = 2, alternative = "one.sided",
           type = "golden")

## two-sided p-values 0.01 and 0.02, relative sample size 2
pSceptical(zo = p2z(0.01), zr = p2z(0.02), c = 2, alternative = "one.sided")
## reverse the studies
pSceptical(zo = p2z(0.02), zr = p2z(0.01), c = 1/2, alternative = "one.sided")
## both p-values 0.01, relative sample size 2
pSceptical(zo = p2z(0.01), zr = p2z(0.01), c = 2, alternative = "two.sided")

zSceptical(zo = 2, zr = 3, c = 2)
zSceptical(zo = 3, zr = 2, c = 2)
```

pvalueBound

Bound for the p-values entering the harmonic mean chi-squared test

Description

Necessary or sufficient bounds for significance of the harmonic mean chi-squared test are computed for n one-sided p-values.

Usage

```
pvalueBound(alpha, n, type = c("necessary", "sufficient"))
```

Arguments

alpha	Numeric vector specifying the significance level.
n	The number of p-values.
type	Either "necessary" (default) or "sufficient". If "necessary", the necessary bounds are computed. If "sufficient", the sufficient bounds are computed.

Value

The bound for the p-values.

Author(s)

Leonhard Held

References

Held, L. (2020). The harmonic mean chi-squared test to substantiate scientific findings. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **69**, 697-708. doi: [10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)

See Also

[hMeanChiSq](#)

Examples

```
pvalueBound(alpha = 0.025^2, n = 2, type = "necessary")
pvalueBound(alpha = 0.025^2, n = 2, type = "sufficient")
```

Qtest

Q-test to assess compatibility between original and replication effect estimate

Description

Computes p-value from meta-analytic Q-test to assess compatibility between original and replication effect estimate.

Usage

```
Qtest(thetao, thetar, seo, ser)
```

Arguments

thetao	Numeric vector of effect estimates from original studies.
thetar	Numeric vector of effect estimates from replication studies.
seo	Numeric vector of standard errors of the original effect estimates.
ser	Numeric vector of standard errors of the replication effect estimates.

Details

This function computes the p-value from a meta-analytic Q-test assessing compatibility between original and replication effect estimate. Rejecting compatibility when the p-value is smaller than alpha is equivalent with rejecting compatibility based on a $(1 - \alpha)$ prediction interval.

Value

p-value from Q-test.

Author(s)

Samuel Pawel

References

Hedges, L. V., Schauer, J. M. (2019). More Than One Replication Study Is Needed for Unambiguous Tests of Replication. *Journal of Educational and Behavioral Statistics*, **44**, 543-570. doi: [10.3102/1076998619852953](https://doi.org/10.3102/1076998619852953)

See Also

[predictionInterval](#)

Examples

```
Qtest(thetao = 2, thetar = 0.5, seo = 1, ser = 0.5)
```

RProjects

Data from four large-scale replication projects

Description

Data from *Reproducibility Project Psychology (RPP)*, *Experimental Economics Replication Project (EERP)*, *Social Sciences Replication Project (SSRP)*, *Experimental Philosophy Replicability Project (EPRP)*. The variables are as follows:

study Study identifier, usually names of authors from original study

project Name of replication project

ro Effect estimate of original study on correlation scale

rr Effect estimate of replication study on correlation scale
 fiso Effect estimate of original study transformed to Fisher-z scale
 fisr Effect estimate of replication study transformed to Fisher-z scale
 se_fiso Standard error of Fisher-z transformed effect estimate of original study
 se_fisr Standard error of Fisher-z transformed effect estimate of replication study
 po Two-sided p-value from significance test of effect estimate from original study
 pr Two-sided p-value from significance test of effect estimate from replication study
 pm_belief Peer belief about whether replication effect estimate will achieve statistical significance elicited through prediction market (only available for EERP and SSRP)
 no Sample size in original study
 nr Sample size in replication study

Usage

```
data(RProjects)
```

Format

A data frame with 143 rows and 13 variables

Details

Two-sided p-values were calculated assuming normality of Fisher-z transformed effect estimates. From the RPP only the *meta-analytic subset* is included, which consists of 73 out of 100 study pairs for which the standard error of the z-transformed correlation coefficient can be computed. For the RPP sample sizes were recalculated from the reported standard errors of Fisher z-transformed correlation coefficients. From the EERP only 31 out of 40 study pairs are included where effective sample size for original and replication study are available simultaneously. For more details about how the the data was preprocessed see source below and supplement S1 of Pawel and Held (2020).

Source

RPP: The source files were downloaded from <https://github.com/CenterForOpenScience/rpp/>. The "masterscript.R" file was executed and the relevant variables were extracted from the generated "final" object (standard errors of Fisher-z transformed correlations) and "MASTER" object (everything else). The data set is licensed under a CC0 1.0 Universal license, see <https://creativecommons.org/publicdomain/zero/1.0/> for the terms of reuse.

EERP: The source files were downloaded from <https://osf.io/pnwuz/>. The required data were then manually extracted from the code in the files "effectdata.py" (sample sizes) and "create_studydetails.do" (everything else). Data regarding the prediction market and survey beliefs were manually extracted from table S3 of the supplementary materials of the EERP. The authors of this R package have been granted permission to share this data set by the coordinators of the EERP.

SSRP: The relevant variables were extracted from the file "D3 - ReplicationResults.csv" downloaded from <https://osf.io/abu7k>. For replications which underwent only the first stage, the data from the first stage were taken as the data for the replication study. For the replications which

reached the second stage, the pooled data from both stages were taken as the data for the replication study. Data regarding survey and prediction market beliefs were extracted from the "D6 - MeanPeerBeliefs.csv" file, which was downloaded from <https://osf.io/vr6p8/>. The data set is licensed under a CC0 1.0 Universal license, see <https://creativecommons.org/publicdomain/zero/1.0/> for the terms of reuse.

EPRP: Data were taken from the "XPhiReplicability_CompleteData.csv" file, which was downloaded from <https://osf.io/4ewkh/>. The authors of this R package have been granted permission to share this data set by the coordinators of the EPRP.

References

- Camerer, C. F., Dreber, A., Forsell, E., Ho, T.-H., Huber, J., Johannesson, M., ... Hang, W. (2016). Evaluating replicability of laboratory experiments in economics. *Science*, **351**, 1433-1436. doi: [10.1126/science.aaf0918](https://doi.org/10.1126/science.aaf0918)
- Camerer, C. F., Dreber, A., Holzmeister, F., Ho, T.-H., Huber, J., Johannesson, M., ... Wu, H. (2018). Evaluating the replicability of social science experiments in Nature and Science between 2010 and 2015. *Nature Human Behaviour*, **2**, 637-644. doi: [10.1038/s415620180399z](https://doi.org/10.1038/s415620180399z)
- Cova, F., Strickland, B., Abatista, A., Allard, A., Andow, J., Attie, M., ... Zhou, X. (2018). Estimating the reproducibility of experimental philosophy. *Review of Philosophy and Psychology*. doi: [10.1007/s1316401804009](https://doi.org/10.1007/s1316401804009)
- Open Science Collaboration. (2015). Estimating the reproducibility of psychological science. *Science*, **349**, aac4716. doi: [10.1126/science.aac4716](https://doi.org/10.1126/science.aac4716)
- Pawel, S., Held, L. (2020). Probabilistic forecasting of replication studies. *PLoS ONE*. **15**, e0231416. doi: [10.1371/journal.pone.0231416](https://doi.org/10.1371/journal.pone.0231416)

See Also

[SSRP](#)

Examples

```
data("RProjects", package = "ReplicationSuccess")

## Computing key quantities
RProjects$zo <- RProjects$fiso/RProjects$se_fiso
RProjects$zr <- RProjects$fisr/RProjects$se_fisr
RProjects$c <- RProjects$se_fiso^2/RProjects$se_fisr^2

## Computing one-sided p-values for alternative = "greater"
RProjects$po1 <- z2p(z = RProjects$zo, alternative = "greater")
RProjects$pr1 <- z2p(z = RProjects$zr, alternative = "greater")

## Plots of effect estimates
parOld <- par(mfrow = c(2, 2))
for (p in unique(RProjects$project)) {
  data_project <- subset(RProjects, project == p)
  plot(rr ~ ro, data = data_project, ylim = c(-0.5, 1),
       xlim = c(-0.5, 1), main = p, xlab = expression(italic(r)[o]),
       ylab = expression(italic(r)[r]))
}
```

```

    abline(h = 0, lty = 2)
    abline(a = 0, b = 1, col = "grey")
  }
  par(parOld)

  ## Plots of peer beliefs
  RProjects$significant <- factor(RProjects$pr < 0.05,
                                levels = c(FALSE, TRUE),
                                labels = c("no", "yes"))

  parOld <- par(mfrow = c(1, 2))
  for (p in c("Experimental Economics", "Social Sciences")) {
    data_project <- subset(RProjects, project == p)
    boxplot(pm_belief ~ significant, data = data_project, ylim = c(0, 1),
            main = p, xlab = "Replication effect significant", ylab = "Peer belief")
    stripchart(pm_belief ~ significant, data = data_project, vertical = TRUE,
              add = TRUE, pch = 1, method = "jitter")
  }
  par(parOld)

  ## Computing the sceptical p-value
  ps <- with(RProjects, pSceptical(zo = fiso/se_fiso,
                                  zr = fisr/se_fisr,
                                  c = se_fiso^2/se_fisr^2))

```

sampleSizeReplicationSuccess

Computes the required relative sample size to achieve replication success based on power or on the minimum relative effect size

Description

The relative sample size to achieve replication success is computed based on the z-value of the original study, the replication success level, the type of recalibration and either the power or the minimum relative effect size. When the approach based on power is used, the design prior also has to be specified.

Usage

```

sampleSizeReplicationSuccess(
  zo,
  power = NA,
  d = NA,
  level = 0.025,
  alternative = c("one.sided", "two.sided"),
  type = c("golden", "nominal", "liberal", "controlled"),
  designPrior = c("conditional", "predictive", "EB"),
  shrinkage = 0,
  h = 0
)

```

Arguments

zo	Numeric vector of z-values from original studies.
power	The power to achieve replication success.
d	The minimum relative effect size (ratio of the effect estimate from the replication study to the effect estimate from the original study) to achieve replication success.
level	Numeric vector of replication success levels. The default is 0.025.
alternative	Either "one.sided" (default) or "two.sided". Specifies if the replication success level is one-sided or two-sided.
type	Type of recalibration. Can be either "golden" (default), "nominal" (no recalibration), "liberal", "controlled". "golden" ensures that for an original study just significant at the specified level, replication success is only possible if the replication effect estimate is at least as large as the original one. See levelSceptical for details about recalibration types.
designPrior	Is only taken into account when power is specified. Either "conditional" (default), "predictive", or "EB". If "EB", the power is computed under a predictive distribution where the contribution of the original study is shrunken towards zero based on the evidence in the original study (with an empirical Bayes shrinkage estimator).
shrinkage	Is only taken into account when power is specified. A number in [0,1) with default 0. Specifies the shrinkage of the original effect estimate towards zero (e.g., the effect is shrunken by a factor of 25% for shrinkage = 0.25). Is only taken into account when the designPrior is "conditional" or "predictive".
h	Is only taken into account when power is specified and designPrior is "predictive" or "EB". The relative between-study heterogeneity, i.e., the ratio of the heterogeneity variance to the variance of the original effect estimate. Default is 0 (no heterogeneity).

Details

sampleSizeReplicationSuccess is the vectorized version of `.sampleSizeReplicationSuccess_`. [Vectorize](#) is used to vectorize the function.

Value

The relative sample size for replication success. If impossible to achieve the desired power for specified inputs NaN is returned.

Author(s)

Leonhard Held, Charlotte Micheloud, Samuel Pawel, Florian Gerber

References

Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[pSceptical](#), [powerReplicationSuccess](#), [levelSceptical](#)

Examples

```
## based on power
sampleSizeReplicationSuccess(zo = p2z(0.0025), power = 0.8, level = 0.025,
                             type = "golden")
sampleSizeReplicationSuccess(zo = p2z(0.0025), power = 0.8, level = 0.025,
                             type = "golden", designPrior = "predictive")

## based on minimum relative effect size
sampleSizeReplicationSuccess(zo = p2z(0.0025), d = 0.9, level = 0.025,
                             type = "nominal")
sampleSizeReplicationSuccess(zo = p2z(0.0025), d = 0.9, level = 0.025,
                             type = "golden")
```

sampleSizeSignificance

Computes the required relative sample size to achieve significance based on power or on the minimum relative effect size

Description

The relative sample size to achieve significance of the replication study is computed based on the z-value of the original study, the significance level and either the power or the minimum relative effect size. When the approach based on power is used, the arguments design prior, shrinkage, and relative heterogeneity also have to be specified.

Usage

```
sampleSizeSignificance(
  zo,
  power = NA,
  d = NA,
  level = 0.025,
  alternative = c("one.sided", "two.sided"),
  designPrior = c("conditional", "predictive", "EB"),
  h = 0,
  shrinkage = 0
)
```

Arguments

zo	A vector of z-values from original studies.
power	The power to achieve replication success.
d	The minimum relative effect size (ratio of the effect estimate from the replication study to the effect estimate from the original study).
level	Significance level. Default is 0.025.
alternative	Either "one.sided" (default) or "two.sided". Specifies direction of the alternative. "one.sided" assumes an effect in the same direction as the original estimate.
designPrior	Is only taken into account when power is specified. Either "conditional" (default), "predictive", or "EB". If "EB", the power is computed under a predictive distribution where the contribution of the original study is shrunken towards zero based on the evidence in the original study (with an empirical Bayes shrinkage estimator).
h	Is only taken into account when power is specified and designPrior is "predictive" or "EB". The relative between-study heterogeneity, i.e., the ratio of the heterogeneity variance to the variance of the original effect estimate. Default is 0 (no heterogeneity).
shrinkage	Is only taken into account when power is specified. A number in [0,1) with default 0. Specifies the shrinkage of the original effect towards zero (e.g., shrinkage = 0.25 implies shrinkage by a factor of 25%). Is only taken into account when designPrior is "conditional" or "predictive".

Details

sampleSizeSignificance is the vectorized version of .sampleSizeSignificance_. [Vectorize](#) is used to vectorize the function.

Value

The relative sample size to achieve significance in the specified direction. If impossible to achieve the desired power for specified inputs NaN is returned.

Author(s)

Leonhard Held, Samuel Pawel, Charlotte Micheloud, Florian Gerber

References

- Held, L. (2020). A new standard for the analysis and design of replication studies (with discussion). *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, **183**, 431-448. doi: [10.1111/rssa.12493](https://doi.org/10.1111/rssa.12493)
- Pawel, S., Held, L. (2020). Probabilistic forecasting of replication studies. *PLoS ONE*. **15**, e0231416. doi: [10.1371/journal.pone.0231416](https://doi.org/10.1371/journal.pone.0231416)
- Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[powerSignificance](#)

Examples

```

sampleSizeSignificance(zo = p2z(0.005), power = 0.8)
sampleSizeSignificance(zo = p2z(0.005, alternative = "two.sided"), power = 0.8)
sampleSizeSignificance(zo = p2z(0.005), power = 0.8, designPrior = "predictive")

sampleSizeSignificance(zo = 3, power = 0.8, designPrior = "predictive",
                       shrinkage = 0.5, h = 0.25)
sampleSizeSignificance(zo = 3, power = 0.8, designPrior = "EB", h = 0.5)

# sample size to achieve 0.8 power as function of original p-value
zo <- p2z(seq(0.0001, 0.05, 0.0001))
oldPar <- par(mfrow = c(1,2))
plot(z2p(zo), sampleSizeSignificance(zo = zo, designPrior = "conditional", power = 0.8),
     type = "l", ylim = c(0.5, 10), log = "y", lwd = 1.5, ylab = "Relative sample size",
     xlab = expression(italic(p)[o]), las = 1)
lines(z2p(zo), sampleSizeSignificance(zo = zo, designPrior = "predictive", power = 0.8),
     lwd = 2, lty = 2)
lines(z2p(zo), sampleSizeSignificance(zo = zo, designPrior = "EB", power = 0.8),
     lwd = 1.5, lty = 3)
legend("topleft", legend = c("conditional", "predictive", "EB"),
     title = "Design prior", lty = c(1, 2, 3), lwd = 1.5, bty = "n")

sampleSizeSignificance(zo = p2z(0.005), d = 1)
sampleSizeSignificance(zo = p2z(0.005), d = 0.5)
# sample size based on minimum relative effect size of 0.8
zo <- p2z(seq(0.0001, 0.05, 0.0001))
plot(z2p(zo), sampleSizeSignificance(zo = zo, d = 0.8, level = 0.025),
     type = "l", ylim = c(0.5, 10), log = "y", lwd = 1.5, ylab = "Relative sample size",
     xlab = expression(italic(p)[o]), las = 1)
par(oldPar)

```

Description

Data from the *Social Sciences Replication Project* (SSRP) including the details of the interim analysis. The variables are as follows:

`study` Study identifier, usually names of authors from original study
`ro` Effect estimate of original study on correlation scale
`ri` Effect estimate of replication study at the interim analysis on correlation scale
`rr` Effect estimate of replication study at the final analysis on correlation scale
`fiso` Effect estimate of original study transformed to Fisher-z scale

fisi Effect estimate of replication study at the interim analysis transformed to Fisher-z scale
fifr Effect estimate of replication study at the final analysis transformed to Fisher-z scale
se_fiso Standard error of Fisher-z transformed effect estimate of original study
se_fisi Standard error of Fisher-z transformed effect estimate of replication study at the interim analysis
se_fifr Standard error of Fisher-z transformed effect estimate of replication study at the final analysis
no Sample size in original study
ni Sample size in replication study at the interim analysis
nr Sample size in replication study at the final analysis
po Two-sided p-value from significance test of effect estimate from original study
pi Two-sided p-value from significance test of effect estimate from replication study at the interim analysis
pr Two-sided p-value from significance test of effect estimate from replication study at the final analysis
n75 Sample size calculated to have 90% power in replication study to detect 75% of the original effect size (expressed as the correlation coefficient r)
n50 Sample size calculated to have 90% power in replication study to detect 50% of the original effect size (expressed as the correlation coefficient r)

Usage

```
data(SSRP)
```

Format

A data frame with 21 rows and 18 variables

Details

Two-sided p-values were calculated assuming normality of Fisher-z transformed effect estimates. A two-stage procedure was used for the replications. In stage 1, the authors had 90% power to detect 75% of the original effect size at the 5% significance level in a two-sided test. If the original result replicated in stage 1 (two-sided P-value < 0.05 and effect in the same direction as in the original study), the data collection was stopped. If not, a second data collection was carried out in stage 2 to have 90% power to detect 50% of the original effect size for the first and the second data collections pooled. n75 and n50 are the planned sample sizes calculated to reach 90% power in stage 1 and 2, respectively. They sometimes differ from the sample sizes that were actually collected (ni and nr, respectively). See supplementary information of Camerer et al. (2018) for details.

Source

<https://osf.io/abu7k>

References

Camerer, C. F., Dreber, A., Holzmeister, F., Ho, T.-H., Huber, J., Johannesson, M., ... Wu, H. (2018). Evaluating the replicability of social science experiments in Nature and Science between 2010 and 2015. *Nature Human Behaviour*, **2**, 637-644. doi: [10.1038/s415620180399z](https://doi.org/10.1038/s415620180399z)

See Also

[RProjects](#)

Examples

```
# plot of the sample sizes
plot(ni ~ no, data = SSRP, ylim = c(0, 2500), xlim = c(0, 400),
     xlab = expression(n[o]), ylab = expression(n[i]))
abline(a = 0, b = 1, col = "grey")

plot(nr ~ no, data = SSRP, ylim = c(0, 2500), xlim = c(0, 400),
     xlab = expression(n[o]), ylab = expression(n[r]))
abline(a = 0, b = 1, col = "grey")
```

T1EpSceptical

Compute type-I error rate of the sceptical p-value

Description

The type-I error rate of the sceptical p-value is computed for a specified level of replication success, the relative variance, and the alternative hypothesis.

Usage

```
T1EpSceptical(
  level,
  c,
  alternative = c("one.sided", "two.sided", "greater", "less"),
  type = c("golden", "nominal", "liberal", "controlled")
)
```

Arguments

level	Numeric vector of levels of replication success.
c	Numeric vector of variance ratios of the original and replication effect estimates. This is usually the ratio of the sample size of the replication study to the sample size of the original study.

alternative	Either "one.sided" (one.sided), "two.sided", "greater", or "less". If "one.sided", the type-I error rate is computed based on a one-sided assessment of replication success in the direction of the original effect estimate. If "two.sided", the type-I error rate is computed based on a two-sided assessment of replication success regardless of the direction of the original and replication effect estimate. If "greater" or "less", the type-I error rate is computed based on a one-sided assessment of replication success in the pre-specified direction of the original and replication effect estimate.
type	Type of recalibration. Can be either "golden" (default), "nominal" (no recalibration), "liberal", or "controlled". "golden" ensures that for an original study just significant at the specified level, replication success is only possible if the replication effect estimate is at least as large as the original one. See levelSceptical for details about recalibration types.

Details

T1EpSceptical is the vectorized version of .T1EpSceptical_. [Vectorize](#) is used to vectorize the function.

Value

The type-I error rate.

Author(s)

Samuel Pawel, Leonhard Held

References

Held, L. (2020). The harmonic mean chi-squared test to substantiate scientific findings. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, **69**, 697-708. doi: [10.1111/rssc.12410](https://doi.org/10.1111/rssc.12410)

Held, L., Micheloud, C., Pawel, S. (2021). The assessment of replication success based on relative effect size. <https://arxiv.org/abs/2009.07782>

See Also

[pSceptical](#), [levelSceptical](#), [PPpSceptical](#)

Examples

```
## compare type-I error rate for different levels of replication success
levels <- c("nominal" = levelSceptical(level = 0.025, type = "nominal"),
           "liberal" = levelSceptical(level = 0.025, type = "liberal"),
           "controlled" = levelSceptical(level = 0.025, type = "controlled"),
           "golden" = levelSceptical(level = 0.025, type = "golden"))
c <- seq(0.2, 5, by = 0.05)
t1 <- sapply(X = levels, FUN = function(l) {
  T1EpSceptical(level = 1, c = c, alternative = "one.sided", type = "nominal")
})
matplot(x = c, y = t1*100, type = "l", lty = 1, lwd = 2, las = 1, log = "x",
```

```

      xlab = bquote(italic(c)), ylab = "Type-I error (%)", xlim = c(0.2, 5))
legend("topright", legend = names(levels), lty = 1, lwd = 2, col = seq_along(levels))

## check that one.sided controlled level controls type-I error rate for c = 1
## at alpha = 0.05*0.025 = 0.00125
T1EpSceptical(level = levelSceptical(level = 0.025, alternative = "one.sided",
                                     type = "controlled"),
              c = 1, alternative = "one.sided", type = "nominal")

```

thresholdIntrinsic *Computes the p-value threshold for intrinsic credibility*

Description

Computes the p-value threshold for intrinsic credibility

Usage

```

thresholdIntrinsic(
  alpha,
  alternative = c("two.sided", "one.sided"),
  type = c("Held", "Matthews")
)

```

Arguments

alpha	Numeric vector of intrinsic credibility levels.
alternative	Either "two.sided" (default) or "one.sided". Specifies if the threshold is for one-sided or two-sided p-values.
type	Either "Held" (default) or "Matthews". Type of intrinsic p-value threshold, see Held (2019) and Matthews (2018) for more information.

Value

The threshold for intrinsic credibility.

Author(s)

Leonhard Held

References

Matthews, R. A. J. (2018). Beyond 'significance': principles and practice of the analysis of credibility. *Royal Society Open Science*, **5**, 171047. doi: [10.1098/rsos.171047](https://doi.org/10.1098/rsos.171047)

Held, L. (2019). The assessment of intrinsic credibility and a new argument for $p < 0.005$. *Royal Society Open Science*, **6**, 181534. doi: [10.1098/rsos.181534](https://doi.org/10.1098/rsos.181534)

Examples

```
thresholdIntrinsic(alpha = c(0.005, 0.01, 0.05))
thresholdIntrinsic(alpha = c(0.005, 0.01, 0.05), alternative = "one.sided")
```

```
unirootAll           Find multiple roots in interval
```

Description

Searches the interval from lower to upper for several roots (i.e., zero's) of a univariate function f .

Usage

```
unirootAll(
  f,
  interval,
  lower = min(interval),
  upper = max(interval),
  n = 1000,
  tol = .Machine$double.eps^0.2,
  maxiter = 1000,
  trace = 0,
  ...
)
```

Arguments

<code>f</code>	the function for which the root is sought. <code>f</code> should be vectorized in the first argument.
<code>interval</code>	A vector containing the end-points of the interval to be searched for the root.
<code>lower</code>	The lower end point of the interval to be searched.
<code>upper</code>	The upper end point of the interval to be searched.
<code>n</code>	Number of subintervals on which <code>link[stats]{uniroot}</code> is called. Default is 1000.
<code>tol</code>	See help of <code>link[stats]{uniroot}</code> .
<code>maxiter</code>	See help of <code>link[stats]{uniroot}</code> .
<code>trace</code>	See help of <code>link[stats]{uniroot}</code> .
<code>...</code>	Additional named or unnamed arguments to be passed to <code>f</code> .

Value

A numeric vector of the roots found in the interval.

Author(s)

Florian Gerber

References

This function is inspired by `rootSolve::uniroot.all()`, package version 1.8.2.2.

See Also

[Vectorize](#)

Examples

```
f <- function (x) cos(2*x)^3
(roots <- unirootAll(f = f, interval = c(0, 10)))
f(roots)
```

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