## Package 'PF'

June 6, 2024

```
Type Package
Title Prevented fraction
Version 9.6.8
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Description Functions related to PF (prevented fraction). Calculate incidence
      density ratio, confidence interval, and Rao-Scott weights of PF by multiple
      methods. See http://goo.gl/eJ6Rxi for definition of PF.
      No endorsement, claim, or warranty is implied for this package. It is made
      available for investigational or pedagogical use only.
License file LICENSE
URL https://www.aphis.usda.gov/animal_health/vet_biologics/publications/
      STATWI0007.pdf, https:
      //github.com/ABS-dev/PF/blob/master/doc/PFPackageManual.pdf
BugReports https://github.com/ABS-dev/PF/issues
LazyLoad true
LazyData true
Depends R (>= 4.1)
Imports methods,
      plyr,
      dplyr,
      tidyr
Suggests magrittr,
      testthat,
      knitr,
      R.rsp,
      rmarkdown
Collate 'aaa.r'
      'aab.r'
      'aac.r'
      'class.r'
      'generics.r'
      'PF.r'
      'PF-package.r'
      'IDRlsi.r'
      'IDRsc.r'
```

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'RR1si.r'
'RRmh.r'
'RRmpWald.r'
'RRor.r'
'RRotsst.r'
'RRsc.r'
'RRStr.r'
'RRtosst.r'
'rsbWt.r'
'tauWt.r'
Encoding UTF-8
<b>Roxygen</b> list(markdown = TRUE)
RoxygenNote 7.2.3
Config/build/clean-inst-doc FALSE
VignetteBuilder R.rsp

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### **Description**

Internal function.

#### Usage

```
.rr.score.asymp(y, alpha = 0.05, iter.max = 18, converge = 1e-04, mn = FALSE)
```

### **Arguments**

y data alpha alpha

iter.max maximum number of iterations

converge convergence criterion

mn boolean whether to calculate MN or use default value of 1.0

### **Examples**

# none

bird bird dataset

### Description

bird dataset

#### **Format**

a data.frame with 6 observations of the following 4 variables, no NAs

y number positive

**n** total number in group tx x all

tx treatment 'vac' or 'con'

all all?

### References

we need some references

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birdm

birdm dataset

### Description

birdm dataset

#### **Format**

```
a data.frame with 6 observations of the following 4 variables, no NAs
y number positive
n total number in group tx x all
tx treatment 'vac' or 'con'
all all?
```

#### References

we need some references

IDRlsi

IDR likelihood support interval.

### Description

Estimates likelihood support interval for the incidence density ratio or prevented fraction based on it.

### Usage

```
IDRlsi(
  y = NULL,
  formula = NULL,
  data = NULL,
  alpha = 0.05,
  k = 8,
  use.alpha = FALSE,
  pf = TRUE,
  converge = 1e-08,
  rnd = 3,
  start = NULL,
  trace.it = FALSE,
  iter.max = 24,
  compare = c("con", "vac")
)
```

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#### **Arguments**

Data vector  $c(y_1, n_1, y_2, n_2)$  where y are the positives, n are the total, and group

1 is compared to group 2 (control or reference).

formula Formula of the form  $cbind(y, n) \sim x$ , where y is the number positive, n is the

group size, x is a factor with two levels of treatment.#'

data data.frame containing variables of the formula.

alpha Complement of the confidence level.

k Likelihood ratio criterion.

use.alpha Base choice of k on its relationship to alpha?

pf Estimate *IDR* or its complement *PF*?

converge Convergence criterion

rnd Number of digits for rounding. Affects display only, not estimates.

start describe here.

trace.it Verbose tracking of the iterations?
iter.max Maximum number of iterations

compare Text vector stating the factor levels: compare[1] is the vaccinate group to which

compare[2] (control or reference) is compared.

#### **Details**

Estimates likelihood support interval for the incidence density ratio based on orthogonal factoring of reparameterized likelihood. The incidence density is the number of cases per subject-time; its distribution is assumed Poisson.

Likelihood support intervals are usually formed based on the desired likelihood ratio, often 1/8 or 1/32. Under some conditions the log likelihood ratio may follow the chi square distribution. If so, then  $\alpha=1-F(2log(k),1)$ , where F is a chi-square CDF. RRsc() will make the conversion from  $\alpha$  to k if use.alpha = TRUE.

The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1 - y1, y2, n2 - y2), 2, 2, byrow = TRUE).

#### Value

A rrsi object with the following elements.

estimate vector with point and interval estimate

estimator either PF or IDR

y data.frame with "y1", "n1", "y2", "n2" values.

k Likelihood ratio criterion

rnd how many digits to round the display alpha complement of confidence level

### Author(s)

PF-package

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#### References

Royall R. *Statistical Evidence: A Likelihood Paradigm*. Chapman & Hall, Boca Raton, 1997. Section 7.2.

#### See Also

**IDRsc** 

```
# Both examples represent the same observation, with data entry by vector
# and matrix notation.
y_{\text{vector}} <- c(26, 204, 10, 205)
IDRlsi(y_vector, pf = FALSE)
# 1/8 likelihood support interval for IDR
# corresponds to 95.858% confidence
# (under certain assumptions)
# IDR LL UL
# 2.61 1.26 5.88
y_{matrix} \leftarrow matrix(c(26, 178, 10, 195), 2, 2, byrow = TRUE)
y_matrix
      [, 1] [, 2]
# [1, ] 26 178
# [2, ] 10 195
IDRlsi(y_matrix, pf = FALSE)
# 1/8 likelihood support interval for IDR
# corresponds to 95.858% confidence
# (under certain assumptions)
# IDR
# IDR LL UL
# 2.61 1.26 5.88
data1 <- data.frame(group = rep(c("treated", "control"), each = 5),</pre>
             n = c(rep(41, 4), 40, rep(41, 5)),
             y = c(4, 5, 7, 6, 4, 1, 3, 3, 2, 1),
             cage = rep(paste('cage', 1:5), 2))
IDRlsi(data = data1, formula = cbind(y, n) ~ group,
              compare = c("treated", "control"), pf = FALSE)
\# 1/8 likelihood support interval for IDR
# corresponds to 95.858% confidence
   (under certain assumptions)
# IDR
# IDR LL UL
```

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IDRsc

IDR confidence interval.

### Description

Estimates confidence interval for the incidence density ratio or prevented fraction based on it.

### Usage

```
IDRsc(
  y = NULL,
  data = NULL,
  formula = NULL,
  compare = c("con", "vac"),
  alpha = 0.05,
  pf = TRUE,
  rnd = 3
)
```

### **Arguments**

у	Data vector c(y1, n1, y2, n2) where y are the positives, n are the total, and group 1 is compared to group 2 (control or reference).
data	data.frame containing variables of formula.
formula	Formula of the form $cbind(y, n) \sim x$ , where y is the number positive, n is the group size, x is a factor with two levels of treatment.
compare	Text vector stating the factor levels: compare[1] is the vaccinate group to which compare[2] (control or reference) is compared.
alpha	Complement of the confidence level.
pf	Estimate <i>IDR</i> , or its complement <i>PF</i> ?
rnd	Number of digits for rounding. Affects display only, not estimates.

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#### **Details**

The incidence density is the number of cases per subject-time; its distribution is assumed Poisson. IDRsc estimates a confidence interval for the incidence density ratio using Siev's formula based on

the Poisson score statistic. 
$$IDR = \widehat{IDR} \left\{ 1 + \left( \frac{1}{y_1} + \frac{1}{y_2} \right) \frac{z_{\alpha/2}^2}{2} \pm \frac{z_{\alpha/2}^2}{2y_1y_2} \sqrt{y_{\bullet} \left( y_{\bullet} z_{\alpha/2}^2 + 4y_1y_2 \right)} \right\}$$

The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1 - y1, y2, n2 - y2), 2, 2, byrow = TRUE).

#### Value

A rr1 object with the following elements.

estimate vector with point and interval estimate

estimator either *PF* or *IDR* 

y data vector

rnd how many digits to round the display alpha complement of confidence level

#### Author(s)

PF-package

#### References

Siev D, 1994. Estimating vaccine efficacy in prospective studies. *Preventive Veterinary Medicine* 20:279-296, Appendix 1.

Graham PL, Mengersen K, Morton AP, 2003. Confidence limits for the ratio of two rates based on likelihood scores:non-iterative method *Statistics in Medicine* 22:2071-2083.

Siev D, 2004. Letter to the editor. *Statistics in Medicine* 23:693. (Typographical error in formula: replace the two final minus signs with subscript dots.)

#### See Also

**IDRlsi** 

```
# All examples represent the same observation, with data entry by vector,
# matrix, and formula+data notation.

y_vector <- c(26, 204, 10, 205)
IDRsc(y_vector, pf = FALSE)

# IDR
# 95% interval estimates

# IDR LL UL
# 2.61 1.28 5.34

y_matrix <- matrix(c(26, 178, 10, 195), 2, 2, byrow = TRUE)
y_matrix
# [, 1] [, 2]
# [1, ] 26 178</pre>
```

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```
# [2, ] 10 195
IDRsc(y_matrix, pf = FALSE)
# IDR
# 95% interval estimates
# IDR LL UL
# 2.61 1.28 5.34
require(dplyr)
data1 <- data.frame(group = rep(c("treated", "control"), each = 5),</pre>
           n = c(rep(41, 4), 40, rep(41, 5)),
           y = c(4, 5, 7, 6, 4, 1, 3, 3, 2, 1),
           cage = rep(paste('cage', 1:5), 2))
data2 <- data1 |>
 group_by(group) |>
  summarize(sum_y = sum(y),
  sum_n = sum(n)
IDRsc(data = data2, formula = cbind(sum_y, sum_n) ~ group,
   compare = c("treated", "control"), pf = FALSE)
# IDR
# 95% interval estimates
# IDR LL UL
# 2.61 1.28 5.34
```

New

New dataset

### **Description**

New dataset

### **Format**

```
a data frame with 52 observations of the following 3 variables, no NAs
```

```
cage cage ID. 1 - 26
```

tx treatment. one of 'con' or 'vac'

**pos** numeric indicator of positive response. 0 = FALSE or 1 = TRUE

### References

We need some references

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pf-class

Data class pf

### Description

data class pf

### **Fields**

- [estimator] either "PF" or "IDR"
- [rnd] how many digits to round display
- [alpha] complement of c.i.

#### Author(s)

PF-package

### See Also

```
rr1, rrsi, rrsc, rrstr
```

phiWt

Binomial dispersion parameter.

### **Description**

MME estimate of dispersion parameter phi.

### Usage

```
phiWt(fit, subset.factor = NULL, fit.only = TRUE, show.warns = FALSE)
```

### **Arguments**

fit A glm object.

subset.factor Factor for estimating phi by subset.

fit.only Return only the new fit? If FALSE, also returns the weights and phi estimates.

show.warns Show warnings

#### **Details**

Estimates binomial dispersion parameter  $\phi$  by the method of moments. Refits the model, weighting the observations by  $1/\phi$ . Uses quasibinomial family in glm().

#### Value

A list with the following elements.

fit the new model fit, updated by the estimated weights

weights vector of weights
phi vector of phi estimates

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### Author(s)

PF-package

### References

Wedderburn RWM, 1974. Quasi-likelihood functions, generalized linear models, and the Gauss-Newton method. *Biometrika* 61:439-447.

#### See Also

```
tauWt, RRor.
```

### **Examples**

```
birdm.fit <- glm(cbind(y, n - y)~tx-1, binomial, birdm)
RRor(phiWt(birdm.fit))
#
# 95% t intervals on 4 df
#
# PF
# PF LL UL
# 0.479 -0.537 0.823
#
# mu.hat LL UL
# txcon 0.768 0.95 0.367
# txvac 0.400 0.78 0.111
#</pre>
```

print.rr1

Print values for PF data obhects.

### Description

Print values for PF data obhects.

### Usage

```
## S3 method for class 'rr1'
print(x, ...)
## S3 method for class 'rror'
print(x, ...)
## S3 method for class 'rrsi'
print(x, ...)
## S3 method for class 'rrmp'
print(x, ...)
## S3 method for class 'rrsc'
print(x, ...)
## S3 method for class 'rrsc'
print(x, ...)
```

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### **Arguments**

x object of class rr1, rror, rrsi, rrmp, rrstr, rrsc other arguments not used by this method

rat ataset

### Description

rat dataset

### **Format**

a data.frame with 32 observations of the following 3 variables, no NAs

y number positive

n total number

group treatment group: 'control' or 'treated'

### References

Weil's rat data (Table 1 of Rao and Scott)

rr1-class

Data class rr1

### Description

Data class rr1

### **Fields**

- [estimate] vector with point and interval estimate
- [estimator] either "PF" or "IDR"
- [y]data.frame with restructured input
- [rnd]how many digits to round display
- [alpha]complement of c.i.

### Author(s)

PF-package

### See Also

IDRsc, RRotsst, RRtosst

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RRlsi RR likelihood support interval.

### Description

likelihood support interval for the risk ratio or prevented fraction by the likelihood profile.

### Usage

```
RRlsi(
  y = NULL,
  formula = NULL,
  data = NULL,
  compare = c("vac", "con"),
  alpha = 0.05,
  k = 8,
  use.alpha = FALSE,
  pf = TRUE,
  iter.max = 50,
  converge = 1e-06,
  rnd = 3,
  start = NULL,
  track = FALSE,
  full.track = FALSE
)
```

### Arguments

full.track

у	Data vector c(y1, n1, y2, n2) where y are the positives, n are the total, and group 1 is compared to group 2 (control or reference group).
formula	Formula of the form $cbind(y, n) \sim x$ , where y is the number positive, n is the group size, x is a factor with two levels of treatment.
data	data.frame containing variables of formula.
compare	Text vector stating the factor levels: compare[1] is the vaccinate group to which compare[2] (control or reference) is compared.
alpha	Complement of the confidence level (see details).
k	Likelihood ratio criterion.
use.alpha	Base choice of k on its relationship to alpha?
pf	Estimate <i>RR</i> or its complement <i>PF</i> ?
iter.max	Maximum number of iterations
converge	Convergence criterion
rnd	Number of digits for rounding. Affects display onlyRR, not estimates.
start	Optional starting value.
track	Verbose tracking of the iterations?

Verbose tracking of the iterations?

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#### **Details**

Estimates a likelihood support interval for RR or PF by the profile likelihood method using the DUD algorithm.

Likelihood support intervals are usually formed based on the desired likelihood ratio, often 1/8 or 1/32. Under some conditions the log likelihood ratio may follow the chi square distribution. If so, then  $\alpha = 1 - F(2log(k), 1)$ , where F is a chi-square CDF. RRlsi() will make the conversion from  $\alpha$  to k if use alpha = TRUE.

The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1-y1, y2, n2-y2), 2, 2, byrow = TRUE).

#### Value

An object of class rrsi with the following fields:

```
estimate matrix of point and interval estimates - see details estimator either "PF" or "RR"

y data.frame with "y1", "n1", "y2", "n2" values.

rnd how many digits to round the display

k likelihood ratio criterion

alpha complement of confidence level
```

#### Author(s)

PF-package

### References

Royall R. Statistical Evidence: A Likelihood Paradigm. Chapman & Hall, Boca Raton, 1997. Section 7.6

Ralston ML, Jennrich RI, 1978. DUD, A Derivative-Free Algorithm for Nonlinear Least Squares. *Technometrics* 20:7-14.

```
# All examples represent the same observation, with data entry by vector,
# matrix, and formula+data notation.

y_vector <- c(4, 24, 12, 28)
RRlsi(y_vector)

# 1/8 likelihood support interval for PF

# corresponds to 95.858% confidence
# (under certain assumptions)

# PF

# PF LL UL
# 0.6111 0.0168 0.8859

y_matrix <- matrix(c(4, 20, 12, 16), 2, 2, byrow = TRUE)</pre>
```

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```
y_matrix
# [, 1] [, 2]
#[1,] 4 20
# [2, ] 12 16
RRlsi(y_matrix)
# 1/8 likelihood support interval for PF
# corresponds to 95.858% confidence
   (under certain assumptions)
# PF
     PF
            LL
                   UL
#
# 0.6111 0.0168 0.8859
require(dplyr)
data1 <- data.frame(group = rep(c("treated", "control"), each = 2),</pre>
 y = c(1, 3, 7, 5),
 n = c(12, 12, 14, 14),
 cage = rep(paste('cage', 1:2), 2))
data2 <- data1 |>
  group_by(group) |>
  summarize(sum_y = sum(y),
   sum_n = sum(n)
RRlsi(data = data2, formula = cbind(sum_y, sum_n) ~ group,
   compare = c("treated", "control"))
# 1/8 likelihood support interval for PF
# corresponds to 95.858% confidence
# (under certain assumptions)
# PF
# PF
        LL
               UL
# 0.6111 0.0168 0.8859
```

RRmh

Mantel-Haenszel method, CI for common RR over strata or clusters with sparse data.

### Description

Estimates confidence intervals for the risk ratio or prevented fraction from clustered or stratified data, using a Mantel-Haenszel estimator for sparse data.

### Usage

```
RRmh(
  formula = NULL,
  data = NULL,
  compare = c("vac", "con"),
  Y,
```

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```
alpha = 0.05,
pf = TRUE,
rnd = 3
)
```

### **Arguments**

formula Formula of the form  $cbind(y, n) \sim x + cluster(w)$ , where y is the number

positive, n is the group size, x is a factor with two levels of treatment, and w is a

factor indicating the clusters.

data data. frame containing variables for formula

compare Text vector stating the factor levels: compare[1] is the vaccinate group to which

compare[2] (control or reference) is compared.

Y Matrix of data,  $K \times 4$ . Each row is a stratum or cluster. The columns are

y1, n1, y2, n2, where the y's are the number of positive in each group, and the n is the total in each group. Group 1 corresponds to vaccinates and group 2 are controls or reference. If data entered by formula and dataframe, Y is generated

automatically.

alpha Complement of the confidence level.

pf Estimate *RR* or its complement *PF*?

rnd Number of digits for rounding. Affects display only, not estimates.

#### **Details**

Based on the Mantel-Haenszel (1959) procedure for sparse data developed by Greenland and Robins (1985). The confidence limits are based on asymptotic normality of the log(risk ratio). Agresti and Hartzel (2000) favor this procedure for small, sparse data sets, but they warn that it is less efficient than maximum likelihood for large data sets.

#### Value

An object of class rr1 with the following fields.

estimate vector of point and interval estimates: point estimate, lower confidence limit,

upper confidence limit

estimator either "PF" or "RR"

y data.frame of restructured input
rnd how many digits to round the display
alpha complement of confidence level

#### Note

If either all y1's or all y2's are zero, a division by zero may occur, and a NaN returned for some values.

Vignette Examples for Stratified Designs forthcoming with more examples.

Call to this function may be one of two formats: (1) specify data and formula or (2) as a matrix Y

```
RRmh(formula, data, compare = c('b', 'a'), pf = TRUE, alpha = 0.05, rnd = 3)
RRmh(Y, pf = TRUE, alpha = 0.05, rnd = 3)
```

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#### Author(s)

PF-package

#### References

Mantel N, Haenszel W, 1959. Statistical aspects of the analysis of data from retrospective studies of disease. *Journal of the National Cancer Institute* 22: 719-748.

Greenland S, Robins JM, 1985. Estimation of a common effect parameter from sparse follow-up data. *Biometrics* 41: 55-68. Errata, 45: 1323-1324.

Agresti A, Hartzel J, 2000. Strategies for comparing treatments on a binary response with multicentre data. *Statistics in Medicine* 19: 1115-1139.

Lachin JM, 2000. *Biostatistical Methods: The Assessment of Relative Risks* (Wiley, New York), Sec. 4.3.1.

#### See Also

rr1

#### **Examples**

```
## Table 1 from Gart (1985)
## as data frame
# tx group "b" is control
RRmh(cbind(y, n) \sim tx + cluster(clus),
    Table6,
    compare = c('a', 'b'), pf = FALSE)
# RR
# 95% interval estimates
   RR LL UL
# 2.67 1.37 5.23
## or as matrix
RRmh(Y = table6, pf = FALSE)
# RR
# 95% interval estimates
  RR LL UL
# 2.67 1.37 5.23
```

rrmp-class

Data class rrmp

### **Description**

data class rrmp

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#### **Fields**

- [estimate] vector with point and interval estimate
- [estimator] either "PF" or "IDR"
- [y] data vector
- [rnd] how many digits to round display
- [alpha] complement of c.i.
- [compare] text vector, same as input
- [multvec] data.frame showing the multinomial representation of the data

### Author(s)

PF-package

### See Also

RRmpWald

RRmpWald

Wald confidence intervals for RR from matched pairs

### **Description**

Estimates confidence intervals for the risk ratio or prevented fraction from matched pairs.

### Usage

```
RRmpWald(
  formula = NULL,
  data = NULL,
  compare = c("vac", "con"),
  affected = 1,
  x,
  alpha = 0.05,
  pf = TRUE,
  tdist = TRUE,
  df = NULL,
  rnd = 3
)
```

### **Arguments**

affected

formula	Formula of the form $y \sim x + cluster(w)$ , where y is the indicator for an individual's positive response, x is a factor with two levels of treatment, and w identifies the pairs.
data	data.frame containing variables in formula
compare	Text vector stating the factor levels: compare[1] is the vaccinate group to which compare[2] (control or reference) is compared.

Indicator for positive response

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Χ	Alternative	data i	nput.	Instead	of	formul	a and	data	frame,	data may	be input as
	C		~			•					

frequency vector. See example for how to order this vector.

alpha Complement of the confidence level pf Estimate *RR* or its complement *PF*?

tdist Use t distribution?

df Degrees of freedom. When NULL, the function will default to df = N - 2, where

N is the total number of pairs.

rnd Number of digits for rounding. Affects display only, not estimates.

#### **Details**

Estimates confidence intervals for the risk ratio or prevented fraction from matched pairs. The response is the tetranomial vector c(11, 12, 21, 22), where the first index is the row and the the second index is the column when displayed as a 2x2 table. Wald type confidence intervals are found by applying the delta method to the multinomial variance. This method fails when there are no responders in one of the treatment groups.

```
Alternative forms of data entry are illustrated by the output, say Y, where c(Y$xtable) = Y$frequec = Y$multvec$Freq.
```

If RR = 0 (PF = 1), the function will return degenerate interval.

#### Value

A rrmp object with the following fields:

estimate vector of point and interval estimates - see details

estimator either "PF" or "RR"

compare text vector, same as input

alpha complement of confidence level

rnd how many digits to round the display

multvec data frame showing the multinomial representation of the data

#### Note

Experimental functions for estimating profile likelihood intervals are in the CVBmisc package.

Call to this function may be one of two formats: (1) specify data and formula or (2) as a vector x

```
RRmpWald(formula, data, compare = c('vac', 'con'), affected = 1, alpha = 0.05,
pf = TRUE, tdist = TRUE, df = NULL, rnd = 3)

RRmpWald(x, compare = c('vac', 'con'), affected = 1, alpha = 0, 05,
pf = TRUE, tdist = TRUE, df = NULL, rnd = 3)
```

#### Author(s)

PF-package

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#### **Examples**

```
RRmpWald(pos ~ tx + cluster(cage), New, compare = c('vac', 'con'))
# PF
# 95% interval estimates
#
#
    PF
         LL
               UL
# 0.550 0.183 0.752
require(magrittr)
thistable <- New |>
 tidyr::spread(tx, pos) |>
 dplyr::mutate(vac = factor(vac, levels = 1:0),
   con = factor(con, levels = 1:0)) |>
 with(table(vac, con))
thistable
    con
# vac 1 0
  1 7 2
# 0 13 4
as.vector(thistable)
#[1] 7 13 2 4
RRmpWald(x = as.vector(thistable))
# PF
# 95% interval estimates
    PF
        LL
# 0.550 0.183 0.752
```

RRor

RR estimate from logistic regression.

### Description

Model based interval estimate of the risk ratio or prevented fraction from a logistic regression model.

### Usage

```
RRor(
  fit = NULL,
  beta.hat = NULL,
  var.beta.hat = NULL,
  degf = NULL,
  which = c(1, 2),
  pf = TRUE,
  norm = FALSE,
  alpha = 0.05,
  rnd = 3
)
```

RRor 21

#### **Arguments**

fit A glm object.

beta.hat Parameters estimates from a logistic regression with no intercept.

var.beta.hat Variance-covariance matrix from a logistic regression with no intercept.

degf Degrees of freedom.

which Numeric vector indicating which parameters to compare, so that RR = compare[2]/compare[1]

pf Estimate *RR* or its complement *PF*?

norm Estimate confidence interval using quantiles of Guassian rather

than t distribution quantiles?

alpha Complement of the confidence level.

rnd Number of digits for rounding. Affects display only, not estimates.

#### **Details**

Estimates confidence intervals using the delta method on parameters from a generalized linear model with logit link.

 $RR = \mu_2/\mu_1$ , where  $\mu_i$  are the estimated probabilities from the model.

#### Value

A rror object with the following fields.

estimate vector with point and interval estimate

estimator either *PF* or *RR* 

mu matrix with rows giving probability estimates for each of the groups

rnd how many digits to round the display

alpha complement of confidence level

norm logical indicating Gaussian or t interval

degf degrees of freedom

#### Note

```
Call to this function may be one of two formats: (1) specify fit or (2) beta.hat, var.beta.hat, degf RRor(fit, degf = NULL, pf = TRUE, alpha = 0.05, which = c(1, 2), norm = TRUE, rnd = 3) RRor(beta.hat, var.beta.hat, degf, pf = TRUE, alpha = 0.05, which = c(1, 2), norm = TRUE,
```

### Author(s)

PF-package

rnd = 3)

### See Also

rror, phiWt, tauWt StatWI007 for more examples

22 rror-class

#### **Examples**

```
bird.fit <- glm(cbind(y, n - y) \sim tx - 1, binomial, bird)
RRor(tauWt(bird.fit))
# 95% t intervals on 4 df
#
# PF
     PF
#
         LL
                   UL
#
 0.500 -0.583 0.842
       mu.hat LL
# txcon 0.733 0.943 0.3121
# txvac 0.367 0.752 0.0997
RRor(phiWt(bird.fit))
# 95% t intervals on 4 df
#
# PF
     PF
#
           LL
                   UL
# 0.500 -0.583 0.842
       mu.hat LL
#
# txcon 0.733 0.943 0.3121
# txvac 0.367 0.752 0.0997
```

rror-class

Data class rror

### Description

data class rror

### **Fields**

- [estimate] vector with point and interval estimate
- [estimator] either "PF" or "IDR"
- [y] data vector
- [rnd] how many digits to round display
- [alpha] complement of c.i.
- [norm] logical indicating Gaussian or t interval
- [degf] degrees of freedom
- [mu] matrix with rows giving probability estimates for each of the groups

### Author(s)

PF-package

### See Also

RRor

RRotsst 23

RRotsst

RR exact CI, OTSST method.

### Description

Estimates confidence interval for the risk ratio or prevented fraction, exact method based on the score statistic (inverts one two-sided test).

### Usage

```
RRotsst(
  y = NULL,
  data = NULL,
  formula = NULL,
  compare = c("vac", "con"),
  alpha = 0.05,
  pf = TRUE,
  stepstart = 0.1,
  iter.max = 36,
  converge = 1e-06,
  rnd = 3,
  trace.it = FALSE,
  nuisance.points = 120,
  gamma = 1e-06
)
```

evaluation)

### Arguments

У	Data vector c(y1, n1, y2, n2) where y are the positives, n are the total, and group 1 is compared to group 2 (control or reference).			
data	data.frame containing variables of the formula.			
formula	Formula of the form $cbind(y, n) \sim x$ , where y is the number positive, n is the group size, x is a factor with two levels of treatment.			
compare	Text vector stating the factor levels: compare[1] is the vaccinate group to which compare[2] (control or reference) is compared.			
alpha	Complement of the confidence level.			
pf	Estimate <i>RR</i> or its complement <i>PF</i> ?			
stepstart	starting interval for step search			
iter.max	Maximum number of iterations			
converge	Convergence criterion			
rnd	Number of digits for rounding. Affects display only, not estimates.			
trace.it	Verbose tracking of the iterations?			
nuisance.points				
	number of points over which to evaluate nuisance parameter			
gamma	parameter for Berger-Boos correction (restricts range of nuisance parameter			

RRotsst RRotsst

#### **Details**

Estimates confidence intervals based on the score statistic that are 'exact' in the sense of accounting for discreteness. The score statistic is used to select tail area tables, and the binomial probability is estimated over the tail area by taking the maximum over the nuisance parameter. Algorithm is a simple step search.

```
The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1 - y1, y2, n2 - y2), 2, 2, byrow = TRUE).
```

#### Value

An object of class rr1 with the following fields:

```
estimate vector with point and interval estimate
estimator either "PF" or "RR"

y data.frame with "y1", "n1", "y2", "n2" values.
rnd how many digits to round the display
alpha complement of confidence level
```

#### Author(s)

PF-package

#### References

Koopman PAR, 1984. Confidence intervals for the ratio of two binomial proportions. *Biometrics* 40:513-517.

Agresti A, Min Y, 2001. On small-sample confidence intervals for parameters in discrete distribution. *Biometrics* 57: 963-971.

Berger RL, Boos DD, 1994. P values maximized over a confidence set for the nuisance parameter. *Journal of the American Statistical Association* 89:214-220.

### See Also

```
RRtosst, rr1.
```

```
# All examples represent the same observation, with data entry by multiple
# options.

y_vector <- c(4, 24, 12, 28)
RRotsst(y_vector, rnd = 3)

# PF
# 95% interval estimates

# PF LL UL
# 0.6111 0.0148 0.8519

y_matrix <- matrix(c(4, 20, 12, 16), 2, 2, byrow = TRUE)
RRotsst(y_matrix, rnd = 3)</pre>
```

RRsc 25

```
# PF
# 95% interval estimates
    PF
           LL
                   UL
# 0.6111 0.0148 0.8519
require(dplyr)
data1 <- data.frame(group = rep(c("treated", "control"), each = 2),</pre>
 y = c(1, 3, 7, 5),
 n = c(12, 12, 14, 14),
 cage = rep(paste('cage', 1:2), 2))
data2 <- data1 |>
  group_by(group) |>
  summarize(sum_y = sum(y),
   sum_n = sum(n)
RRotsst(data = data2, formula = cbind(sum_y, sum_n) ~ group,
  compare = c("treated", "control"))
# 95% interval estimates
# PF
        LL
# 0.6111 0.0148 0.8519
```

RRsc

RR score based asymptotic CI.

#### **Description**

Estimates confidence intervals for the risk ratio or prevented fraction based on the score statistic.

### Usage

```
RRsc(
  y = NULL,
  data = NULL,
  formula = NULL,
  compare = c("vac", "con"),
  alpha = 0.05,
  pf = TRUE,
  trace.it = FALSE,
  iter.max = 18,
  converge = 1e-06,
  rnd = 3
)
```

### Arguments

Data vector c(y1, n1, y2, n2) where y are the positives, n are the total, and group 1 is compared to group 2 (control or reference group).

data data.frame containing variables of formula.

26 RRsc

formula Formula of the form  $cbind(y, n) \sim x$ , where y is the number positive, n is the

group size, x is a factor with two levels of treatment.

compare Text vector stating the factor levels: compare[1] is the vaccinate group to which

compare[2] (control or reference) is compared.

alpha Complement of the confidence level. pf Estimate RR or its complement PF? trace.it Verbose tracking of the iterations? iter.max Maximum number of iterations

converge Convergence criterion

rnd Number of digits for rounding. Affects display only, not estimates.

#### **Details**

Estimates are returned for three estimators based on the score statistic. The score method was introduced by Koopman (1984). Gart and Nam's modification (1988) includes a skewness correction. The method of Miettinen and Nurminen (1985) is a version made slightly more conservative than Koopman's by including a factor of (N-1)/N. The starting estimate for the DUD algorithm is obtained by the modified Katz method (log method with 0.5 added to each cell). Both forms of the Katz estimate may be retrieved from the returned object using RRsc()\$estimate.

The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1-y1, y2, n2-y2), 2, 2, byrow = TRUE).

#### Value

A rrsc object with the following fields.

estimate matrix of point and interval estimates - see details

estimator either "PF" or "RR"

y data.frame with "y1", "n1", "y2", "n2" values.

rnd how many digits to round the display alpha complement of confidence level

#### Author(s)

PF-package

### References

Gart JJ, Nam J, 1988. Approximate interval estimation of the ratio of binomial parameters: a review and corrections for skewness. *Biometrics* 44:323-338.

Koopman PAR, 1984. Confidence intervals for the ratio of two binomial proportions. *Biometrics* 40:513-517.

Miettinen O, Nurminen M, 1985. Comparative analysis of two rates. *Statistics in Medicine* 4:213-226.

Ralston ML, Jennrich RI, 1978. DUD, A Derivative-Free Algorithm for Nonlinear Least Squares. *Technometrics* 20:7-14.

#### See Also

rrsc

rrsc-class 27

```
# All examples represent the same observation, with data entry by using
# multiple notation options.
y_{\text{vector}} < -c(4, 24, 12, 28)
RRsc(y_vector)
# PF
# 95% interval estimates
# PF
         LL
               UL
# MN method
              0.611 0.0251 0.857
# score method 0.611 0.0328 0.855
# skew corr 0.611 0.0380 0.876
y_{matrix} \leftarrow matrix(c(4, 20, 12, 16), 2, 2, byrow = TRUE)
# [, 1] [, 2]
# [1, ] 4 20
# [2, ] 12 16
RRsc(y_matrix)
# PF
# 95% interval estimates
# PF
         LL
               UL
# MN method 0.611 0.0251 0.857
# score method 0.611 0.0328 0.855
# skew corr 0.611 0.0380 0.876
require(dplyr)
data1 <- data.frame(group = rep(c("treated", "control"), each = 2),</pre>
 y = c(1, 3, 7, 5),
 n = c(12, 12, 14, 14),
 cage = rep(paste('cage', 1:2), 2))
data2 <- data1 |>
  group_by(group) |>
  summarize(sum_y = sum(y),
   sum_n = sum(n)
RRsc(data = data2, formula = cbind(sum_y, sum_n) ~ group,
 compare = c("treated", "control"))
# PF
# 95% interval estimates
# PF
         LL
               UL
              0.611 0.0251 0.857
# MN method
# score method 0.611 0.0328 0.855
# skew corr 0.611 0.0380 0.876
```

28 rrsi-class

### Description

data class rrsc

### **Fields**

- [estimate] vector with point and interval estimate
- [rnd] how many digits to round display
- [alpha] complement of c.i.
- [estimator] either "PF" or "RR"
- [y] data.frame with restructured input

### Author(s)

PF-package

### See Also

RRsc

rrsi-class

Data class rrsi

### Description

data class rrsi

#### **Fields**

- [y] data.frame with restructured input
- [k] likelihood ratio criterion
- [rnd] digits to round display
- [alpha] complement of c.i.
- [estimate] vector with point and interval estimate
- [estimator] either "PF" or "IDR"

### Author(s)

PF-package

### See Also

IDRlsi, RRlsi

RRstr 29

RRstr

Gart-Nam method, CI for common RR over strata or clusters.

### Description

Estimates confidence intervals for the risk ratio or prevented fraction from clustered or stratified data.

### Usage

```
RRstr(
  formula = NULL,
  data = NULL,
  compare = c("vac", "con"),
  Y,
  alpha = 0.05,
  pf = TRUE,
  trace.it = FALSE,
  iter.max = 24,
  converge = 1e-06,
  rnd = 3,
  multiplier = 0.7,
  divider = 1.1
)
```

### Arguments

formula	Formula of the form $cbind(y, n) \sim x + cluster(w)$ , where y is the number positive, n is the group size, x is a factor with two levels of treatment, and w is a factor indicating the clusters.
data	data.frame containing variables of formula
compare	Text vector stating the factor levels: compare[1] is the control or reference group to which compare[2] is compared
Υ	Matrix of data. Each row is a stratum or cluster. The columns are y2, n2, y1, n1. If data entered by formula and dataframe, Y is generated automatically.
alpha	Size of the homogeneity test and complement of the confidence level.
pf	Estimate RR or its complement PF?
trace.it	verbose tracking of the iterations?
iter.max	Maximum number of iterations
converge	Convergence criterion
rnd	Number of digits for rounding. Affects display only, not estimates.
multiplier	internal control parameter for algorithm
divider	internal control parameter for algorithm

### **Details**

Uses the DUD algorithm to estimate confidence intervals by the method of Gart.

30 RRstr

#### Value

A rrstr object with the following fields:

estimate matrix of point and interval estimates - starting value, MLE, and skewness cor-

rected

hom list of homogeneity statistic, p-value, and degrees of freedom, or error message

if appropriate.

estimator either "PF" or "RR"

y data.frame of restructured input

compare groups compared

rnd how many digits to round the display

alpha size of test; complement of confidence level

#### Note

Vignette Examples for Stratified Designs forthcoming with more examples.

Call to this function may be one of two formats: (1) specify data and formula or (2) as a matrix Y

```
RRstr(formula, data, compare = c('b', 'a'), pf = TRUE, alpha = 0.05, trace.it = FALSE, iter.max = 24, converge = 1e-6, rnd = 3, multiplier = 0.7, divider = 1.1)
```

```
RRstr(Y, compare = c('b', 'a'), pf = TRUE, alpha = 0.05, trace.it = FALSE, iter.max = 24, converge = 1e-6, rnd = 3, multiplier = 0.7, divider = 1.1)
```

#### Author(s)

PF-package

### References

Gart JJ, 1985. Approximate tests and interval estimation of the common relative risk in the combination of  $2x^2$  tables. *Biometrika* 72:673-677.

Gart JJ, Nam J, 1988. Approximate interval estimation of the ratio of binomial parameters: a review and corrections for skewness. *Biometrics* 44:323-338.

Ralston ML, Jennrich RI, 1978. DUD, A Derivative-Free Algorithm for Nonlinear Least Squares. *Technometrics* 20:7-14.

### See Also

rrstr

rrstr-class 31

```
# stat
          0.954
# df
          3
# p
          0.812
# RR estimates
        RR LL UL
# starting 2.66 1.37 5.18
# mle 2.65 1.39 5.03
# skew corr 2.65 1.31 5.08
## or as matrix
RRstr(Y = table6, pf = FALSE)
tst <- data.frame(y = c(0, 2, 0, 4, 0, 3, 0, 7),
n = rep(10, 8),
tx = rep(c('a', 'b'), 4),
clus = rep(paste('Row', 1:4, sep = ''), each = 2))
```

rrstr-class

Data class rrstr

### **Description**

data class rrstr

### **Fields**

- [estimate] vector with point and interval estimate
- [rnd] how many digits to round display
- [alpha] complement of c.i.
- [estimator] either "PF" or "RR"
- [hom]list of homogeneity statistic, p-value, and degrees of freedom. If Phi == 0 | Phi == 1, homogeneity test is not possible and error message displays
- [y]data.frame of restructured input
- [compare] groups compared

### Author(s)

PF-package

#### See Also

RRstr

RRtosst

RRtosst

RR exact CI, TOSST method.

### Description

Estimates confidence interval for the risk ratio or prevented fraction; exact method based on the score statistic (inverts two one-sided tests).

### Usage

```
RRtosst(
  y = NULL,
  formula = NULL,
  data = NULL,
  compare = c("vac", "con"),
  alpha = 0.05,
  pf = TRUE,
  stepstart = 0.1,
  iter.max = 36,
  converge = 1e-06,
  rnd = 3,
  trace.it = FALSE,
  nuisance.points = 120,
  gamma = 1e-06
)
```

### Arguments

У	Data vector $c(y1, n1, y2, n2)$ where y are the positives, n are the total, and group 1 is compared to group 2 (control or reference group).				
formula	Formula of the form $cbind(y, n) \sim x$ , where y is the number positive, n is the group size, x is a factor with two levels of treatment.				
data	data.frame containing variables of formula.				
compare	Text vector stating the factor levels: compare[1] is the vaccinate group to which compare[2] (control or reference) is compared.				
alpha	Complement of the confidence level.				
pf	Estimate <i>RR</i> or its complement <i>PF</i> ?				
stepstart	starting interval for step search				
iter.max	Maximum number of iterations				
converge	Convergence criterion				
rnd	Number of digits for rounding. Affects display only, not estimates.				
trace.it	Verbose tracking of the iterations?				
nuisance.points					
	number of points over which to evaluate nuisance parameter				
gamma	parameter for Berger-Boos correction (restricts range of nuisance parameter evaluation)				

RRtosst 33

#### **Details**

Estimates confidence intervals based on the score statistic that are 'exact' in the sense of accounting for discreteness. Inverts two one-sided score tests. The score statistic is used to select tail area tables, and the binomial probability is estimated over the tail area by taking the maximum over the nuisance parameter. Algorithm is a simple step search.

The data may also be a matrix. In that case y would be entered as matrix(c(y1, n1-y1, y2, n2-y2), 2, 2, byrow = TRUE).

### Value

A rr1 object with the following fields.

```
estimate vector with point and interval estimate
estimator either "PF" or "RR"

y data.frame with "y1", "n1", "y2", "n2" values.
rnd how many digits to round the display
alpha complement of confidence level
```

### Author(s)

PF-package

#### References

Koopman PAR, 1984. Confidence intervals for the ratio of two binomial proportions. *Biometrics* 40:513-517.

Agresti A, Min Y, 2001. On small-sample confidence intervals for parameters in discrete distribution. *Biometrics* 57: 963-971.

Berger RL, Boos DD, 1994. P values maximized over a confidence set for the nuisance parameter. *Journal of the American Statistical Association* 89:214-220.

#### See Also

```
RRotsst, rr1
```

```
# Both examples represent the same observation, with data entry by vector
# and matrix notation.

y_vector <- c(4, 24, 12, 28)
RRtosst(y_vector)

# PF
# 95% interval estimates

# PF LL UL
# 0.611 0.012 0.902

y_matrix <- matrix(c(4, 20, 12, 16), 2, 2, byrow = TRUE)
# [, 1] [, 2]
# [1, ] 4 20</pre>
```

34 rsb

```
# [2, ] 12 16
RRtosst(y_matrix)
# PF
# 95% interval estimates
          LL
                 UL
# 0.611 0.012 0.902
require(dplyr)
data1 <- data.frame(group = rep(c("treated", "control"), each = 2),</pre>
 y = c(1, 3, 7, 5),
 n = c(12, 12, 14, 14),
  cage = rep(paste('cage', 1:2), 2))
data2 <- data1 |>
  group_by(group) |>
  summarize(sum_y = sum(y),
   sum_n = sum(n)
RRtosst(data = data2, formula = cbind(sum_y, sum_n) ~ group,
 compare = c("treated", "control"))
# PF
# 95% interval estimates
    PF
          LL
# 0.611 0.012 0.902
```

rsb

Rao-Scott weights.

### **Description**

Rao-Scott weights.

### Usage

```
rsb(y = NULL, n = NULL, formula = NULL, data = NULL, id = NULL)
```

### **Arguments**

y vector of number positive.

n vector of total number.

formula Formula of the form cbind(y, n) ~ id, where y is the number positive, n is the total number, id is a factor for estimating the weights by subset.

data data.frame containing variables of formula.

id vector of factor for estimating the weights by subset.

### **Details**

Estimates the cluster design effect  $d_i$  as the variance inflation due to clustering by the method of Rao and Scott. rsb estimates the  $d_i$  for use by rsbWt or other functions.

rsbWt 35

#### Value

A list with the following elements.

```
w vector of weights  {\sf d} \qquad \qquad {\sf vector} \ {\sf of} \ d_i \ {\sf estimates}
```

### Author(s)

```
PF-package
```

#### References

Rao JNK, Scott AJ, 1992. A simple method for the analysis of clustered binary data. *Biometrics* 48:577-585.

#### See Also

rsbWt.

### **Examples**

```
# Weil's rat data (Table 1 of Rao and Scott)
rsb(rat$y, rat$n, id = rat$group)$d
# control treated
# 1.232495 3.952861
rsb(data = rat, formula = cbind(y, n) ~ group)$d
# control treated
# 1.232495 3.952861
```

rsbWt

Rao-Scott weighting.

### **Description**

Rao-Scott weighting of clustered binomial observations.

### Usage

```
rsbWt(fit = NULL, subset.factor = NULL, fit.only = TRUE)
```

### **Arguments**

fit A glm object.

subset.factor Factor for estimating the weights by subset.

fit.only Return only the new fit? If FALSE, also returns the weights and phi estimates.

### **Details**

Estimates the cluster design effect  $d_i$  as the variance inflation due to clustering by the method of Rao and Scott. Observations are then weighted by the inverse of the  $d_i$ .

36 Set1

#### Value

A list with the following elements.

fit the new model fit, updated by the estimated weights weights vector of weights  ${\rm d} \qquad {\rm vector\ of}\ d_i \ {\rm estimates}$ 

#### Author(s)

PF-package

#### References

Rao JNK, Scott AJ, 1992. A simple method for the analysis of clustered binary data. *Biometrics* 48:577-585.

#### See Also

```
RRor, rsb.
```

### **Examples**

```
birdm.fit <- glm(cbind(y,n-y)~tx-1,binomial,birdm)
RRor(rsbWt(birdm.fit))
#
# 95% t intervals on 4 df
#
# PF
# PF LL UL
# 0.479 -1.061 0.868
#
# mu.hat LL UL
# txcon 0.768 0.968 0.2659
# txvac 0.400 0.848 0.0737
#</pre>
```

Set1

Set1 dataset

### Description

Set1 dataset

#### **Format**

```
a data.frame with 6 observation of the following 4 variables, no NAs
```

```
\mathbf{y} number positive
```

 ${\bf n}$  total number in group tx x clus

```
tx treatment 'vac' or 'con'
```

clus cluster ID

set1 37

#### References

We need some references

set1

set1 dataset

### Description

set1 dataset

### **Format**

a 3 x 4 matrix of data in Set1

### References

we need some references!

Table6

Table6 dataset

### Description

Table6 dataset

### **Format**

a data.frame with 8 observations of the following 4 variables, no NAs

y number positive

 ${\bf n}$  total number in group tx x clus

tx treatment 'a' or 'b'

clus cluster ID

#### References

Table 1 from Gart (1985)

table6

table6 dataset

### Description

table6 dataset

#### **Format**

matrix for of data in Table6

38 tauWt

tauWt

Binomial dispersion: intra-cluster correlation parameter.

#### **Description**

MME estimates of binomial dispersion parameter tau (intra-cluster correlation).

#### Usage

```
tauWt(
  fit,
  subset.factor = NULL,
  fit.only = TRUE,
  iter.max = 12,
  converge = 1e-06,
  trace.it = FALSE
)
```

#### **Arguments**

fit A glm object.

subset.factor Factor for estimating tau by subset.

fit.only Return only the final fit? If FALSE, also returns the weights and tau estimates.

iter.max Maximum number of iterations.

converge Convergence criterion: difference between model degrees of freedom and Pear-

son's chi-square. Default 1e-6.

trace.it Display print statments indicating progress

### **Details**

Estimates binomial dispersion parameter  $\tau$  by the method of moments. Iteratively refits the model by the Williams procedure, weighting the observations by  $1/\phi_{ij}$ , where  $\phi_{ij} = 1 + \tau_j(n_{ij} - 1)$ , j indexes the subsets, and i indexes the observations.

### Value

A list with the following elements.

fit the new model fit, updated by the estimated weights

weights vector of weights
phi vector of phi estimates

### Author(s)

PF-package

#### References

Williams DA, 1982. Extra-binomial variation in logistic linear models. *Applied Statistics* 31:144-148

Wedderburn RWM, 1974. Quasi-likelihood functions, generalized linear models, and the Gauss-Newton method. *Biometrika* 61:439-447.

tauWt 39

### See Also

```
phiWt, RRor.
```

```
birdm.fit <- glm(cbind(y,n-y)~tx-1, binomial, birdm)
RRor(tauWt(birdm.fit))
# 95% t intervals on 4 df
#
# PF
# PF LL UL
# 0.489 -0.578 0.835
#
# mu.hat LL UL
# txcon 0.737 0.944 0.320
# txvac 0.376 0.758 0.104
#</pre>
```

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