

# Package ‘RSurveillance’

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**Type** Package

**Title** Design and Analysis of Disease Surveillance Activities

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**Description** A range of functions for the design and analysis of disease surveillance activities. These functions were originally developed for animal health surveillance activities but can be equally applied to aquatic animal, wildlife, plant and human health surveillance activities. Utilities are included for sample size calculation and analysis of representative surveys for disease freedom, risk-based studies for disease freedom and for prevalence estimation.

This package is based on Cameron A., Conraths F., Frohlich A., Schauer B., Schulz K., Sergeant E., Sonnenburg J., Staubach C. (2015). R package of functions for risk-based surveillance. Deliverable 6.24, WP 6 - Decision making tools for implementing risk-based surveillance, Grant Number no. 310806, RISKSUR (<[https://www.fp7-risksur.eu/sites/default/files/documents/Deliverables/RISKSUR\\_%28310806%29\\_D6.24.pdf](https://www.fp7-risksur.eu/sites/default/files/documents/Deliverables/RISKSUR_%28310806%29_D6.24.pdf)>).

Many of the 'RSurveillance' functions are incorporated into the 'epitools' website: Sergeant, ESG, 2019. Epitools epidemiological calculators.

Ausvet Pty Ltd. Available at: <<http://epitools.ausvet.com.au>>.

**License** GPL-2 | GPL-3

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**URL** <https://github.com/roStats/RSurveillance>

**Imports** epitools, epiR, stats, mc2d

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'freedom\_functions\_2.R' 'n\_rb2stage.R' 'prevalence\_functions.R'  
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adj.risk	<i>Adjusted risk</i>
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## Description

Calculates adjusted risk for given relative risk and population proportions. This is an intermediate calculation in the calculation of effective probability of infection for risk-based surveillance activities

## Usage

```
adj.risk(rr, ppr)
```

## Arguments

- |     |  |
|-----|--|
| rr  | relative risk values (vector of values corresponding to the number of risk strata) |
| ppr | population proportions corresponding to rr values (vector of equal length to rr)   |

## Value

vector of adjusted risk values (in order corresponding to rr)

## Examples

```
# examples for adj.risk
adj.risk(c(5, 1), c(0.1, 0.9))
adj.risk(c(5, 3, 1), c(0.1, 0.1, 0.8))
```

**adj.risk.sim***Adjusted risk for simulation models***Description**

Calculates adjusted risk for given relative risk and population proportions. This is an intermediate calculation in the calculation of effective probability of infection for risk-based surveillance activities. This function is similar to Adj.risk, except is adapted for use with multiple simulations instead of single RR values.

**Usage**

```
adj.risk.sim(rr, ppr)
```

**Arguments**

- |            |   |
|------------|---|
| <b>rr</b>  | relative risk values (matrix of values, columns corresponding to the number of risk strata, rows corresponding to number of iterations for simulation ) |
| <b>ppr</b> | population proportions corresponding to rr values (vector of equal length to columns in rr)   |

**Value**

matrix of adjusted risk values (in order corresponding to rr)

**Examples**

```
# examples for adj.risk.sim
its<- 10
risk.cat<- 3
rr<- matrix(0, nrow=its, ncol=risk.cat)
rr[,1]<- mc2d::rpert(its, 5,10,20)
rr[,2]<- mc2d::rpert(its, 2,3,5)
rr[,3]<- 1
ppr<- c(0.05, 0.2, 0.75)
adj.risk.sim(rr, ppr)
adj.risk.sim(matrix(c(5, 3, 1), nrow=1), matrix(c(0.1, 0.1, 0.8), nrow=1))
```

ap

*Apparent prevalence***Description**

Estimates apparent prevalence and confidence limits for given sample size and result, assuming representative sampling

**Usage**

```
ap(x, n, type = "wilson", conf = 0.95)
```

**Arguments**

x	number of positives in sample
n	sample size, note: either x or n can be a vector, but at least one must be scalar
type	method for estimating CI, one of c("normal", "exact", "wilson", "jeffreys", "agresti-coull", "all"), default = "wilson"
conf	level of confidence required, default = 0.95 (scalar)

**Value**

either 1) if type = "all", a list with 5 elements, each element a matrix with 6 columns, x, n, proportion, lower confidence limit, upper confidence limit, confidence level and CI method; or 2) a matrix of results for the chosen method

**Examples**

```
# examples for ap function
n<- 200
x<- 25
conf<- 0.95
ap(x, n)
ap(seq(10, 100, 10), 200, type = "agresti")
ap(seq(10, 100, 10), 200, type = "all")
```

binom.agresti

*Agresti-Coull confidence limits***Description**

Calculates Agresti-Coull confidence limits for a simple proportion (apparent prevalence)

**Usage**

```
binom.agresti(x, n, conf = 0.95)
```

**Arguments**

x	number of positives in sample
n	sample size, note: either x or n can be a vector, but at least one must be scalar
conf	level of confidence required, default 0.95 (scalar)

**Value**

a data frame with 6 columns, x, n, proportion, lower confidence limit, upper confidence limit, confidence level and CI method

## Examples

```
# test binom.agresti
binom.agresti(25, 200)
binom.agresti(seq(10, 100, 10), 200)
binom.agresti(50, seq(100, 1000, 100))
```

**binom.cp**

*Clopper-Pearson exact confidence limits*

## Description

Calculates Clopper-Pearson exact binomial confidence limits for a simple proportion (apparent prevalence)

## Usage

```
binom.cp(x, n, conf = 0.95)
```

## Arguments

- |      |   |
|------|---|
| x    | number of positives in sample   |
| n    | sample size, note: either x or n can be a vector, but at least one must be scalar |
| conf | level of confidence required, default = 0.95 (scalar)                             |

## Value

a dataframe with 6 columns, x, n, proportion, lower confidence limit, upper confidence limit, confidence level and CI method

## Examples

```
# test binom.cp
binom.cp(25, 200)
binom.cp(seq(10, 100, 10), 200)
binom.cp(50, seq(100, 1000, 100))
```

<code>binom.jeffreys</code>	<i>Jeffreys confidence limits</i>
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### Description

Calculates Jeffreys confidence limits for a simple proportion (apparent prevalence)

### Usage

```
binom.jeffreys(x, n, conf = 0.95)
```

### Arguments

<code>x</code>	number of positives in sample
<code>n</code>	sample size, note: either <code>x</code> or <code>n</code> can be a vector, but at least one must be scalar
<code>conf</code>	level of confidence required, default = 0.95 (scalar)

### Value

a dataframe with 6 columns, `x`, `n`, proportion, lower confidence limit, upper confidence limit, confidence level and CI method

### Examples

```
# test binom.jeffreys
binom.jeffreys(25, 200)
binom.jeffreys(seq(10, 100, 10), 200)
binom.jeffreys(50, seq(100, 1000, 100))
```

<code>disc.prior</code>	<i>Discounted prior probability of freedom</i>
-------------------------	--

### Description

Calculates the discounted prior probability of disease freedom, after adjusting for the probability of disease exceeding the design prevalence during the time period of the surveillance data being analysed

### Usage

```
disc.prior(prior, p.intro)
```

### Arguments

<code>prior</code>	prior probability of freedom before surveillance
<code>p.intro</code>	probability of introduction (or of prevalence exceeding the design prevalence) during the time period (scalar or vector equal length to <code>prior</code> )

**Value**

vector of discounted prior probabilities of freedom

**Examples**

```
# examples for disc.prior
disc.prior(0.5, 0.01)
disc.prior(0.95, c(0.001, 0.005, 0.01, 0.02, 0.05))
disc.prior(c(0.5, 0.6, 0.7, 0.8, 0.9, 0.95), 0.01)
```

---

**epi.calc**

*Effective probability of infection (EPI)*

---

**Description**

Calculates effective probability of infection (adjusted design prevalence) for each risk group for risk-based surveillance activities

**Usage**

```
epi.calc(pstar, rr, ppr)
```

**Arguments**

<b>pstar</b>	design prevalence (scalar)
<b>rr</b>	relative risk values (vector of values corresponding to the number of risk strata)
<b>ppr</b>	population proportions corresponding to rr values (vector of equal length to rr)

**Value**

list of 2 elements, a vector of EPI values and a vector of corresponding adjusted risks (in corresponding order to rr)

**Examples**

```
# examples for epi.calc
epi.calc(0.1, c(5, 1), c(0.1, 0.9))
epi.calc(0.02, c(5, 3, 1), c(0.1, 0.1, 0.8))
```

---

n.2stage	<i>2-stage freedom sample size</i>
----------	------------------------------------

---

## Description

Calculates sample sizes for a 2-stage representative survey (sampling of clusters and units within clusters) for disease freedom or detection, assuming imperfect test sensitivity, perfect test specificity and representative sampling

## Usage

```
n.2stage(H = NA, N = NA, sep.sys = 0.95, sep.c, pstar.c, pstar.u,
se = 1)
```

## Arguments

H	population size = number of clusters or NA if not known, default = NA
N	population sizes for clusters, default = NA, scalar or vector of population sizes for clusters
sep.sys	desired population sensitivity (scalar)
sep.c	desired cluster-level sensitivity (scalar)
pstar.c	specified cluster-level design prevalence as proportion or integer (scalar)
pstar.u	specified population-level design prevalence as proportion or integer (scalar)
se	unit sensitivity (scalar)

## Value

a list of 2 elements, the number of clusters to sample and a vector of sample sizes per cluster

## Examples

```
# examples of n.2stage - checked
n.2stage(NA, NA, 0.95, 0.5, 0.01, 0.1, 0.95)
n.2stage(500, NA, 0.95, 0.5, 10, 0.1, 0.95)
n.2stage(1000, c(50, 100, 200, 500, 1000, 5000, NA), 0.95, 0.5, 0.01, 0.05, 0.8)
n.2stage(1000, c(50, 100, 200, 500, 1000, 5000, NA), 0.95, 0.5, 0.01, 1, 0.8)
n.2stage(1000, c(50, 100, 200, 500, 1000, 5000, NA), 0.9, 0.95, 1, 0.1, 0.8)
```

**n.ap***Sample size for apparent prevalence***Description**

Calculates sample size for estimating apparent prevalence (simple proportion)

**Usage**

```
n.ap(p, precision, conf = 0.95)
```

**Arguments**

<b>p</b>	expected proportion, scalar or vector of values
<b>precision</b>	absolute precision, +/- proportion equivalent to half the width of the desired confidence interval, scalar or vector of values, note: at least one of p and precision must be a scalar
<b>conf</b>	level of confidence required, default = 0.95 (scalar)

**Value**

a vector of sample sizes

**Examples**

```
# examples of n.ap
n.ap(0.5, 0.1)
n.ap(0.5, 0.1, conf=0.99)
n.ap(seq(0.1, 0.5, by = 0.1), 0.05)
n.ap(0.2, c(0.01, 0.02, 0.05, 0.1))
```

**n.binom***Binomial sample size***Description**

Calculates sample size for demonstrating freedom or detecting disease using binomial approach and assuming imperfect test sensitivity, perfect test specificity and representative sampling

**Usage**

```
n.binom(sep, pstar, se = 1)
```

**Arguments**

sep	desired population sensitivity (scalar or vector)
pstar	specified design prevalence (scalar or vector of same length as sep)
se	unit sensitivity, default = 1 (scalar or vector of same length as sep)

**Value**

vector of sample sizes

**Examples**

```
# examples for n.binom - checked
n.binom(sep=0.95, pstar=c(0.01, 0.02, 0.05, 0.1, 0.2))
n.binom(c(0.5, 0.8, 0.9, 0.95), 0.01)
```

n.c.freecalc

*Freecalc optimum sample size and cut-point number of positives*

**Description**

Calculates optimum sample size and cut-point number of positives to achieve specified population sensitivity, for given population size and other parameters, using freecalc algorithm, all paramaters must be scalars

**Usage**

```
n.c.freecalc(N, sep = 0.95, c = 1, se, sp = 1, pstar,
minSpH = 0.95)
```

**Arguments**

N	population size
sep	target population sensitivity
c	The maximum allowed cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive
se	test unit sensitivity
sp	test unit specificity, default=1
pstar	design prevalence as a proportion or integer (number of infected units)
minSpH	minimum desired population specificity

**Value**

a list of 3 elements, a dataframe with 1 row and six columns for the recommended sample size and corresponding values for population sensitivity (SeP), population specificity (SpP), N, c and pstar, a vector of SeP values and a vector of SpP values, for n = 1:N

## Examples

```
# examples for n.c.hp
n.c.freecalc(120,0.95,c=5,se=0.9,sp=0.99,pstar=0.1, minSpH=0.9)[[1]]
n.c.freecalc(65,0.95,c=5,se=0.95,sp=0.99,pstar=0.05, minSpH=0.9)
```

n.c.hp

*Hypergeometric (HerdPlus) optimum sample size and cut-point number of positives*

## Description

Calculates optimum sample size and cut-point positives to achieve specified population sensitivity, for given population size and other parameters, all paramaters must be scalars

## Usage

```
n.c.hp(N, sep = 0.95, c = 1, se, sp = 1, pstar, minSpH = 0.95)
```

## Arguments

N	population size
sep	target population sensitivity
c	The maximum allowed cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive
se	test unit sensitivity
sp	test unit specificity, default=1
pstar	design prevalence as a proportion or integer (number of infected units)
minSpH	minimum desired population specificity

## Value

a list of 3 elements, a datafame with 1 row and six columns for the recommended sample size and corresponding values for population sensitivity (SeP), population specificity (SpP), N, c and pstar, a vector of SeP values and a vector of SpP values, for n = 1:N

## Examples

```
# examples for n.c.hp
n.c.hp(65,0.95,c=5,se=0.95,sp=0.99,pstar=0.05, minSpH=0.9)[[1]]
tmp<- n.c.hp(120,0.95,c=5,se=0.9,sp=0.99,pstar=0.1, minSpH=0.9)
```

---

n.freecalc	<i>Freecalc sample size for a finite population and specified cut-point number of positives</i>
------------	---

---

## Description

Calculates sample size required for a specified population sensitivity, for a given population size, cut-point number of positives and other parameters, using Freecalc algorithm. All paramaters must be scalars

## Usage

```
n.freecalc(N, sep = 0.95, c = 1, se, sp = 1, pstar, minSpH = 0.95)
```

## Arguments

N	population size
sep	target population sensitivity
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive
se	test unit sensitivity
sp	test unit specificity, default=1
pstar	design prevalence as a proportion or integer (number of infected units)
minSpH	minimum desired population specificity

## Value

a list of 2 elements, a dataframe with 1 row and six columns for the recommended sample size and corresponding values for population sensitivity (SeP), population specificity (SpP), N, c and pstar and a dataframe of n rows with SeP and SpP values for each value of n up to the recommended value

## Examples

```
# examples for n.freecalc
n.freecalc(65,0.95,c=1,se=0.95,sp=0.99,pstar=0.05, minSpH=0.9)[[1]]
n.freecalc(65,0.95,c=2,se=0.95,sp=0.99,pstar=0.05, minSpH=0.9)[[1]]
n.freecalc(65,0.95,c=3,se=0.95,sp=0.99,pstar=0.05, minSpH=0.9)
```

---

n.freedom	<i>Freedom sample size</i>
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---

### Description

Calculates sample size for demonstrating freedom or detecting disease using the appropriate method, depending on whether or not N provided (hypergeometric if N provided, binomial otherwise), assuming imperfect test sensitivity, perfect test specificity and representative sampling

### Usage

```
n.freedom(N = NA, sep = 0.95, pstar, se = 1)
```

### Arguments

N	population size, default = NA (unknown) (scalar or vector of same length as sep)
sep	desired population sensitivity (scalar or vector)
pstar	specified design prevalence as proportion or integer (scalar or vector of same length as sep)
se	unit sensitivity (scalar or vector of same length as sep)

### Value

vector of sample sizes, NA if N is specified and n>N

### Examples

```
# examples for n.freedom - checked
n.freedom(NA, sep=0.95, pstar=0.01, se=1)
n.freedom(500, sep=0.95, pstar=0.01, se=1)
n.freedom(N=c(100, 500, 1000, 5000, 10000, 100000, NA), sep=0.95, pstar=0.01, se=1)
n.freedom(500, sep=0.95, pstar=0.01, se=c(0.5, 0.6, 0.7, 0.8, 0.9, 0.99, 1))
```

---

n.hp	<i>Hypergeometric (HerdPlus) sample size for finite population and specified cut-point number of positives</i>
------	--

---

### Description

Calculates sample size to achieve specified population sensitivity with population specificity >= specified minimum value, for given population size, cut-point number of positives and other parameters, all paramaters must be scalars

### Usage

```
n.hp(N, sep = 0.95, c = 1, se, sp = 1, pstar, minSpH = 0.95)
```

**Arguments**

N	population size
sep	target population sensitivity
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, $\geq c$ is positive
se	test unit sensitivity
sp	test unit specificity, default=1
pstar	design prevalence as a proportion or integer (number of infected units)
minSpH	minimum desired population specificity

**Value**

A list of 2 elements, a data frame with 1 row and six columns for the recommended sample size and corresponding values for population sensitivity (SeP), population specificity (SpP), N, c and pstar and a data frame of n rows with SeP and SpP values for each value of n up to the recommended value. Returns sample size for maximum achievable sep if it is not possible to achieve target sep AND SpP  $\geq$  minSpH.

**Examples**

```
# examples for n.hp
n.hp(65, 0.95, c=1, se=0.95, sp=0.99, pstar=0.05, minSpH=0.9)[[1]]
n.hp(65, 0.95, c=2, se=0.95, sp=0.99, pstar=0.05, minSpH=0.9)
```

n.hypergeo

*Hypergeometric sample size***Description**

Calculates sample size for demonstrating freedom or detecting disease using hypergeometric approximation and assuming imperfect test sensitivity, perfect test specificity and representative sampling

**Usage**

```
n.hypergeo(sep, N, d, se = 1)
```

**Arguments**

sep	desired population sensitivity (scalar or vector)
N	population size (scalar or vector of same length as sep)
d	expected number of infected units in population, = design prevalence*N rounded to next integer (scalar or vector of same length as sep)
se	unit sensitivity, default = 1 (scalar or vector of same length as sep)

**Value**

vector of sample sizes, NA if n>N

**Examples**

```
# examples for n.hypergeo - checked
n.hypergeo(0.95, N=100, d=1, se = 0.95)
n.hypergeo(sep=0.95, N=c(100, 200, 500, 1000, 10000), d=ceiling(0.01*c(100, 200, 500, 1000, 10000)))
n.hypergeo(c(0.5, 0.8, 0.9, 0.95), N=100, d=5)
n.hypergeo(0.95, N=80, d=c(1, 2, 5, 10))
n.hypergeo(0.95, N=80, d=c(1, 2, 5, 10), se = 0.8)
```

n.pfree

*Sample size to achieve desired (posterior) probability of freedom*

**Description**

Calculates the sample size required to achieve a given value for probability of disease freedom

**Usage**

```
n.pfree(pfree, prior, p.intro, pstar, se, N = NA)
```

**Arguments**

pfree	desired probability of freedom (scalar or vector)
prior	prior probability of freedom before surveillance (scalar or vector of same length as pfree)
p.intro	probability of introduction for time period (scalar or vector of same length as pfree)
pstar	design prevalence (scalar or vector of same length as pfree)
se	unit sensitivity (scalar or vector of same length as pfree)
N	population size (scalar or vector of same length as pfree)

**Value**

a list of 3 elements, the first a vector of sample sizes and the second a corresponding vector of population sensitivity values and the third a vector of adjusted priors

**Examples**

```
# examples for n.pfree
n.pfree(0.95, 0.5, 0.01, 0.05, 0.9)
n.pfree(0.95, 0.5, 0.01, 0.05, 0.9, N=300)
n.pfree(pfree = c(0.9, 0.95, 0.98, 0.99), prior = 0.7, 0.01, 0.01, 0.8, 1000)
n.pfree(0.95, 0.7, 0.01, 0.1, 0.96)
```

---

n.pooled	<i>Sample size for pooled testing for freedom</i>
----------	---

---

**Description**

Calculates sample size to achieve desired population-level sensitivity, assuming pooled sampling and allowing for imperfect sensitivity and specificity of the pooled test

**Usage**

```
n.pooled(sep, k, pstar, pse, psp = 1)
```

**Arguments**

sep	desired population sensitivity (scalar or vector)
k	pool size (constant across pools) (scalar or vector of same length as sep)
pstar	design prevalence (scalar or vector of same length as sep)
pse	pool-level sensitivity (scalar or vector of same length as sep)
psp	pool-level specificity (scalar or vector of same length as sep)

**Value**

vector of sample sizes

**Examples**

```
# examples for n.pooled
n.pooled(0.95, 5, 0.01, 1, 1)
n.pooled(0.95, 10, 0.1, 0.9, 1)
n.pooled(0.95, c(2, 5, 10, 20), 0.1, c(0.99, 0.98, 0.97, 0.95), 1)
```

---

n.rb	<i>Risk-based sample size</i>
------	-------------------------------

---

**Description**

Calculates sample size for risk-based sampling for a single risk factor and using binomial method

**Usage**

```
n.rb(pstar, rr, ppr, spr, se, sep)
```

**Arguments**

pstar	design prevalence (scalar)
rr	relative risk values (vector, length equal to the number of risk strata)
ppr	population proportions corresponding to rr values (vector of equal length to rr)
spr	planned surveillance proportion for each risk group (vector equal length to rr, ppr)
se	unit sensitivity (fixed or vector same length as rr, ppr, n)
sep	required population sensitivity (scalar)

**Value**

list of 2 elements, a vector of sample sizes for each risk group a scalar of total sample size, a vector of EPI values and a vector of adjusted risks

**Examples**

```
# examples for n.rb
n.rb(0.1, c(5, 3, 1), c(0.1, 0.10, 0.80), c(0.5, 0.3, 0.2), 0.9, 0.95)
n.rb(0.01, c(5, 1), c(0.1, 0.9), c(0.8, 0.2), c(0.9, 0.95), 0.95)
```

n.rb.2stage.1	<i>sample size for 2-stage risk-based surveillance, risk factor at cluster level only</i>
---------------	---

**Description**

Calculates sample size required (clusters and units) for a 2-stage risk-based survey with a single risk factor at the cluster level only.

**Usage**

```
n.rb.2stage.1(rr, ppr, spr, pstar.c, pstar.u, se = 1, sep.c = 0.95,
sep.sys = 0.95)
```

**Arguments**

rr	relative risk values (vector of values, corresponding to the number of risk strata)
ppr	population proportions corresponding to rr values (vector of equal length to rr)
spr	planned surveillance proportions corresponding to rr values - the proportions of the total sample to be collected from each risk stratum (vector of equal length to rr).
pstar.c	cluster (herd) level design prevalence, scalar, either proportion or integer
pstar.u	unit (animal) level design prevalence, scalar, either proportion or integer
se	unit sensitivity of test (proportion), scalar, default = 1
sep.c	desired cluster-level sensitivity (proportion), scalar, default = 0.95
sep.sys	desired population-level sensitivity (proportion), scalar, default = 0.95

### Value

A list of seven elements: 1) a vector (of the same length as rr) of the numbers of clusters to sample from each risk stratum, 2) the total number of clusters to be sampled, 3) a vector of EPI values for each risk stratum, 4) a vector of adjusted risk values for each risk stratum, 5) the number of units to be sampled per cluster 6) a vector of the total numbers of units to be sampled for each risk stratum 7) the overall total number of units to be sampled

### Examples

```
rr<- c(5,3,1)
ppr<- c(0.1, 0.2, 0.7)
spr<- c(0.4, 0.4, 0.2)
n.rb.2stage.1(rr, ppr, spr, pstar.c=0.01, pstar.u=0.1, se =0.9, sep.c=0.8, sep.sys=0.95)
n.rb.2stage.1(c(3,1), c(0.2,0.8), c(0.7,0.3),0.05, 0.1, 0.9, 0.95, 0.99)
```

n.rb.2stage.2

*Sample size for 2-stage risk-based surveillance, allowing for risk factors at either or both cluster and unit level*

### Description

Calculates sample size required (clusters and units) for a 2-stage risk-based survey with risk factors at either cluster level or unit level, or both.

### Usage

```
n.rb.2stage.2(rr.c, ppr.c, spr.c, pstar.c, rr.u = 1, ppr.u = 1,
spr.u = 1, pstar.u, se = 1, sep.c = 0.95, sep.sys = 0.95)
```

### Arguments

rr.c	relative risk values at the cluster level (vector of values, corresponding to the number of risk strata)
ppr.c	population proportions at the cluster level, corresponding to rr.c values (vector of equal length to rr.c)
spr.c	planned surveillance proportions at the cluster level, corresponding to rr.c values - the proportions of the total sample to be collected from each risk stratum (vector of equal length to rr.c).
pstar.c	cluster (herd) level design prevalence, scalar, either proportion or integer
rr.u	relative risk values at the unit level (vector of values, corresponding to the number of risk strata)
ppr.u	population proportions at the unit level, corresponding to rr.u values (vector of equal length to rr.u)
spr.u	planned surveillance proportions at the unit level, corresponding to rr.u values - the proportions of the total sample to be collected from each risk stratum (vector of equal length to rr.u).

<code>pstar.u</code>	unit (animal) level design prevalence, scalar, either proportion or integer
<code>se</code>	unit sensitivity of test (proportion), scalar, default = 1
<code>sep.c</code>	desired cluster-level sensitivity (proportion), scalar, default = 0.95
<code>sep.sys</code>	desired population-level sensitivity (proportion), scalar, default = 0.95

**Value**

A list of cluster and unit level results number of clusters/units to sample per risk stratum, the total number of clusters or units per cluster to be sampled and vectors of EPI and adjusted risk values for each risk stratum.

**Examples**

```
rr.c<- c(5,3,1)
ppr.c<- c(0.1, 0.2, 0.7)
spr.c<- c(0.4, 0.4, 0.2)
rr.u<- c(4,1)
ppr.u<- c(0.1, 0.9)
spr.u<- c(1, 0)
n.rb.2stage.2(rr.c, ppr.c, spr.c, pstar.c=0.02, rr.u, ppr.u,
               spr.u, 0.1, se=0.9, sep.c=0.5, sep.sys=0.95)
n.rb.2stage.2(c(3,1), c(0.2,0.8), c(0.7,0.3), pstar.c=0.05,
               pstar.u=0.1, se=0.9, sep.c=0.95, sep.sys=0.99)
```

**n.rb.varse***Risk-based sample size for varying unit sensitivity***Description**

Calculates sample size for risk-based sampling for a single risk factor and varying unit sensitivity, using binomial method

**Usage**

```
n.rb.varse(pstar, rr, ppr, spr, se, spr.rg, sep)
```

**Arguments**

<code>pstar</code>	design prevalence (scalar)
<code>rr</code>	relative risk values (vector, length equal to the number of risk strata)
<code>ppr</code>	population proportions for each risk group, vector of same length as rr
<code>spr</code>	planned surveillance proportions for each risk group, vector of same length as rr
<code>se</code>	unit sensitivities (vector of group values)
<code>spr.rg</code>	proportions of samples for each sensitivity value in each risk group (matrix with rows = risk groups, columns = sensitivity values), row sums must equal 1
<code>sep</code>	required population sensitivity (scalar)

**Value**

list of 3 elements, a matrix of sample sizes for each risk and sensitivity group, a vector of EPI values and a vector of mean sensitivity for each risk group

**Examples**

```
# examples for n.rb.varse
m<- rbind(c(0.8, 0.2), c(0.5, 0.5), c(0.7, 0.3))
n.rb.varse(0.01, c(5, 3, 1), c(0.1, 0.1, 0.8), c(0.4, 0.4, 0.2), c(0.92, 0.8), m, 0.95)

m<- rbind(c(0.8, 0.2), c(0.6, 0.4))
n.rb.varse(0.05, c(3, 1), c(0.2, 0.8), c(0.7, 0.3), c(0.95, 0.8), m, 0.95)

m<- rbind(c(1), c(1))
n.rb.varse(0.05, c(3, 1), c(0.2, 0.8), c(0.7, 0.3), c(0.95), m, 0.99)
```

n.tp

*Sample size for true prevalence***Description**

Calculates sample size for estimating true prevalence using normal approximation

**Usage**

```
n.tp(p, se, sp, precision, conf = 0.95)
```

**Arguments**

p	estimated true prevalence (scalar or vector)
se	test sensitivity (scalar or vector)
sp	test specificity (scalar or vector)
precision	absolute precision, +/- proportion equal to half the width of the desired confidence interval (scalar or vector)
conf	desired level of confidence for CI, default = 0.95 (scalar or vector)

**Value**

a vector of sample sizes

**Examples**

```
# examples for n.tp
n.tp(0.1, 0.9, 0.99, 0.05)
n.tp(0.1, 0.9, 0.99, 0.05, conf = 0.99)
n.tp(c(0.05, 0.1, 0.2, 0.3, 0.4, 0.5), 0.9, 0.99, 0.05)
n.tp(0.5, 0.9, 0.99, c(0.01, 0.02, 0.05, 0.1, 0.2))
```

**pfree.1***Probability of freedom for single time period***Description**

Calculates the posterior probability (confidence) of disease freedom (negative predictive value) for a single time period

**Usage**

```
pfree.1(sep, p.intro, prior = 0.5)
```

**Arguments**

<code>sep</code>	population sensitivity for time period (scalar or vector)
<code>p.intro</code>	probability of introduction for time period (scalar or vector of same length as <code>sep</code> )
<code>prior</code>	prior probability of freedom before surveillance (scalar or vector of same length as <code>sep</code> )

**Value**

`data.frame` with columns for `sep`, `p.intro`, discounted prior, `pfree`, `pfree.equ` and `prior.equ`

**Examples**

```
# examples for pfree.1
pfree.1(0.8, 0.01, 0.5)
pfree.1(0.6, c(0.001, 0.005, 0.01, 0.02, 0.05), 0.5)
pfree.1(runif(10, 0.4, 0.6), 0.01, 0.5)
pfree.1(runif(10, 0.4, 0.6), runif(10, 0.005, 0.015), 0.5)
```

**pfree.calc***Probability of freedom over time***Description**

Calculates the probability (confidence) of disease freedom for given prior, `sep` and `p.intro` over 1 or more time periods

**Usage**

```
pfree.calc(sep, p.intro, prior = 0.5, discount.1 = TRUE)
```

**Arguments**

sep	population sensitivity for each time period (vector)
p.intro	probability of introduction for each time period (scalar or vector of same length as sep)
prior	prior probability of freedom before surveillance (scalar)
discount.1	logical variable whether or not to discount first time period.

**Value**

data.frame with columns for sep, p.intro, discounted prior, probability of freedom, equilibrium probability of freedom and equilibrium prior

**Examples**

```
# examples for pfree.calc
pfree.calc(0.8, 0.01, 0.5)
pfree.calc(0.8, 0.01, 0.5, FALSE)
pfree.calc(rep(0.6,24), 0.01, 0.5)
pfree.calc(rep(0.6,24), 0.01, 0.5, FALSE)
pfree.calc(runif(10, 0.4, 0.6), 0.01, 0.5)
pfree.calc(runif(10, 0.4, 0.6), runif(10, 0.005, 0.015), 0.5)
```

**pfree.equ***Equilibrium probability of freedom***Description**

Calculates equilibrium probability of disease freedom and equilibrium prior probability of freedom, after discounting for probability of introduction

**Usage**

```
pfree.equ(sep, p.intro)
```

**Arguments**

sep	population sensitivity for time period (scalar or vector)
p.intro	probability of introduction for time period (scalar or vector of same length as sep)

**Value**

a list of 2 vectors, equilibrium posterior probability of freedom and equilibrium prior (discounted) probability of freedom

## Examples

```
# examples of pfree.equ
pfree.equ(runif(10, 0.4, 0.6), 0.01)
pfree.equ(0.8, 0.05)
pfree.equ(rep(0.9, 6), c(0.0001, 0.0005, 0.001, 0.005, 0.01, 0.05))
```

**pstar.calc**

*Design prevalence back-calculation*

## Description

Calculates design prevalence required for given sample size and desired population-level sensitivity, assuming imperfect test sensitivity, perfect test specificity and representative sampling

## Usage

```
pstar.calc(N = NA, n, sep, se)
```

## Arguments

N	populaton size if known (scalar or vector of same length as n)
n	sample size (scalar or vector)
sep	desired population sensitivity (scalar or vector of same length as n)
se	unit sensitivity (scalar or vector of same length as n)

## Value

vector of design prevalence values

## Examples

```
# examples of pstar.calc- checked
pstar.calc(NA, 280, 0.95, 0.98)
pstar.calc(500, 250, sep=0.95, se=1)
pstar.calc(N=c(100, 500, 1000, 5000, 10000, 100000, NA), n=30, sep=0.95, se=1)
pstar.calc(500, n=30, sep=0.95, se=c(0.5, 0.6, 0.7, 0.8, 0.9, 0.99, 1))
```

---

se.parallel                  *Sensitivity of tests in parallel*

---

**Description**

Calculates the combined sensitivity for multiple tests interpreted in parallel (assuming independence)

**Usage**

```
se.parallel(se)
```

**Arguments**

se                  vector of unit sensitivity values

**Value**

scalar of combined sensitivity, assuming independence

**Examples**

```
# examples for se.parallel  
se.parallel(c(0.99, 0.95, 0.8))
```

---

se.series                  *Sensitivity of tests in series*

---

**Description**

Calculates the combined sensitivity for multiple tests interpreted in series (assuming independence)

**Usage**

```
se.series(se)
```

**Arguments**

se                  vector of unit sensitivity values

**Value**

scalar of combined sensitivity, assuming independence

**Examples**

```
# examples for se.series  
se.series(c(0.99, 0.95, 0.8))
```

---

sep	<i>Population sensitivity</i>
-----	-------------------------------

---

## Description

Calculates population sensitivity using appropriate method, depending on whether or not N provided (hypergeometric if N provided, binomial otherwise), assuming perfect test specificity and representative sampling

## Usage

```
sep(N = NA, n, pstar, se = 1, dig = 5)
```

## Arguments

N	population size, NA or vector of same length as n
n	sample size (number tested), scalar or vector
pstar	design prevalence as a proportion or integer, scalar or vector of same length as n
se	unit sensitivity, scalar or vector of same length as n
dig	number of digits for rounding of results

## Value

a vector of population-level sensitivities

## Examples

```
# examples for sep - checked
sep(n=300, pstar=0.01, se=1)
sep(NA, 300, 0.01, 1)
sep(10000, 150, 0.02, 1)
sep(n=1:100, pstar = 0.05, se=0.95)
N<- seq(30, 100, by = 5)
se<- 0.95
pstar<- 0.1
n<- rep(30, length(N))
sep(N, n, pstar, se = se)
sep(rep(100, 10), seq(10, 100, by = 10), pstar = 1, se=0.99)
N<- c(55, 134, NA, 44, 256)
n<- c(15, 30, 28, 15, 33)
sep(N, n, 0.1, 0.95)
```

sep.binom

*Binomial Population sensitivity***Description**

Calculates population sensitivity for detecting disease, assuming imperfect test sensitivity and specificity and representative sampling, using binomial distribution (assumes large or unknown population size and that cut-point number of reactors for a positive result = 1)

**Usage**

```
sep.binom(n, pstar, se = 1, sp = 1, dig = 5)
```

**Arguments**

n	sample size = number of units tested (integer), scalar or vector
pstar	design prevalence as a proportion (scalar or vector of same length as n)
se	unit sensitivity of test (proportion), default = 1 (scalar or vector of same length as n)
sp	unit specificity of test (proportion), default = 1 (scalar or vector of same length as n)
dig	number of digits for rounding of results

**Value**

vector of population-level sensitivities

**Examples**

```
# examples for sep.binom - checked
sep.binom(n=300, pstar = 0.02, se = 0.92)
tested<- seq(10,100, by=10)
prev<- 0.05
sens<- 0.9
sep.binom(tested, prev, sens)
```

sep.binom.imperfect

*Binomial population sensitivity for imperfect test***Description**

Calculates population sensitivity for a large or unknown population and allowing for imperfect test sensitivity and specificity, using Binomial distribution an allowing for a variable cut-point number of positives to classify as positive

**Usage**

```
sep.binom.imperfect(n, c = 1, se, sp = 1, pstar)
```

**Arguments**

n	sample size (scalar or vector)
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive (scalar or vector of same length as n)
se	test unit sensitivity (scalar or vector of same length as n)
sp	test unit specificity, default=1 (scalar or vector of same length as n)
pstar	design prevalence as a proportion (scalar or vector of same length as n)

**Value**

a vector of population-level sensitivities

**Examples**

```
# examples for sep.imperfect.binom
sep.binom.imperfect(1:10*5, 2, 0.95, 0.98, 0.1)
sep.binom.imperfect(50, 1:5, 0.95, 0.98, 0.1)
sep.binom.imperfect(30, 2, 0.9, 0.98, 0.1)
sep.binom.imperfect(30, 1, 0.9, 0.98, 0.1)
```

**sep.exact**

*Population sensitivity for census (all units tested)*

**Description**

Calculates population sensitivity for detecting disease assuming imperfect test sensitivity, perfect test specificity and a census of all units in the population

**Usage**

```
sep.exact(d = 1, se = 1, dig = 5)
```

**Arguments**

d	expected number of infected units in population (=design prevalence*N rounded to next integer), scalar or vector of same length as se
se	unit sensitivity of test (proportion), scalar or vector
dig	number of digits for rounding of results

**Value**

vector of population-level sensitivities

## Examples

```
# examples for sep.exact - checked
sep.exact(d=1, se = 0.92)
inf<- 1:5
sens<- 0.8
sep.exact(d=inf, se=sens)
sep.exact(se=0.8, d = ceiling(0.01*c(10, 50, 100, 250, 500)))
```

**sep.freecalc**

*FreeCalc population sensitivity for imperfect test*

## Description

Calculates population sensitivity for a finite population and allowing for imperfect test sensitivity and specificity, using Freecalc method

## Usage

```
sep.freecalc(N, n, c = 1, se, sp = 1, pstar)
```

## Arguments

N	population size (scalar)
n	sample size (scalar)
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive (scalar)
se	test unit sensitivity (scalar)
sp	test unit specificity, default=1 (scalar)
pstar	design prevalence as a proportion - assumed or target prevalence for detection of disease in the population (scalar)

## Value

population-level sensitivity

## Examples

```
# examples of sep.freecalc
sep.freecalc(150, 30, 2, 0.9, 0.98, 0.1)
sep.freecalc(150, 30, 1, 0.9, 0.98, 0.1)
```

---

sep.hp*Hypergeometric (HerdPlus) population sensitivity for imperfect test*

---

**Description**

Calculates population sensitivity for a finite population and allowing for imperfect test sensitivity and specificity, using Hypergeometric distribution

**Usage**

```
sep.hp(N, n, c = 1, se, sp = 1, pstar)
```

**Arguments**

N	population size (scalar or vector of same length as n)
n	sample size (scalar or vector)
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive (scalar)
se	test unit sensitivity (scalar)
sp	test unit specificity, default=1 (scalar)
pstar	design prevalence as a proportion (scalar)

**Value**

a vector of population-level sensitivities

**Examples**

```
# examples of sep.hp
sep.hp(150, 1:5*10, 2, 0.9, 0.98, 0.1)
sep.hp(150, 30, 2, 0.9, 0.98, 15)
sep.hp(150, 30, 1, 0.9, 0.98, 15)
sep.hp(150, 30, 1, 0.9, 0.98, 0.1)
```

---

sep.hypergeo*Hypergeometric Population sensitivity*

---

**Description**

Calculates population sensitivity for detecting disease, assuming imperfect test sensitivity, perfect test specificity and representative sampling, using hypergeometric approximation (assumes known population size)

**Usage**

```
sep.hypergeo(N, n, d, se = 1, dig = 5)
```

**Arguments**

N	population size, scalar or vector of same length as n
n	sample size (number tested), scalar or vector
d	expected number of infected units in population (=design prevalence*N rounded to next integer)
se	unit sensitivity of test (proportion), scalar or vector of same length as n
dig	number of digits for rounding of results

**Value**

a vector of population-level sensitivities. if all n <= corresponding N then vector is numeric, otherwise vector is character and elements where n>N are recorded as such

**Examples**

```
# examples for sep.hypergeo - checked
sep.hypergeo(N=100, n=50, d=1, se = 0.92)
inf<- 1:5
sens<- 0.8
sep.hypergeo(N=100, n=50, d=inf, se=sens)
N<- c(10, 50, 100, 250, 500)
sep.hypergeo(se=0.8, N=N, n=c(5, 25, 50, 125, 250), d = ceiling(0.01*N))
```

sep.passive

*Passive surveillance sensitivity***Description**

Estimates the population sensitivity of a passive surveillance system. Assumes comprehensive population coverage and samling of representative affected units from infected clusters

**Usage**

```
sep.passive(step.p, p.inf.u, se, N, n, pstar.c)
```

**Arguments**

step.p	vector or matrix of detection probabilities for each step in the detection process. If a vector each value represents a step probability for a single calculation. If a matrix, columns are step probabilities and rows are simulation iterations.
p.inf.u	the probability of infection in units sampled, equivalent to the positive predictive value of clinical signs of disease (for a given prior probability of infection). Either a scalar or vector with length equal to number of rows in step.p.

<b>se</b>	unit sensitivity of test (proportion). Either a scalar or vector with length equal to number of rows in step.p.
<b>N</b>	population size. Either a scalar or vector with length equal to number of rows in step.p
<b>n</b>	number of units tested per cluster reporting suspected disease. Either a scalar or vector with length equal to number of rows in step.p
<b>pstar.c</b>	cluster-level design prevalence (proportion). Either a scalar or vector with length equal to number of rows in step.p

**Value**

a list of 2 elements, the estimated cluster-level and population-level sensitivities. If step.p is a vector, values are scalars, if step.p is a matrix, values are vectors with length equal to the number of rows in step.p

**Examples**

```
# examples for sep.passive
sep.passive(c(0.1, 0.2, 0.9, 0.99), 0.98, 0.9, 1000, 5, 0.01)
sep.passive(c(0.1, 0.5, 0.95, 0.99), 0.98, 0.9, 1000, 5, 0.01)
step.p<- matrix(runif(30), nrow=10)
p.inf.u<- runif(10, 0.98, 0.999)
se<- mc2d::rpert(10, 0.9, 0.95, 0.98)
sep.passive(step.p, p.inf.u, se, 10000, 10, 0.02)
```

**sep.pfree**

*Population sensitivity to achieve desired (posterior) probability of freedom*

**Description**

Calculates the population sensitivity required to achieve a given value for probability of disease freedom

**Usage**

```
sep.pfree(prior, pfree)
```

**Arguments**

<b>prior</b>	prior probability of freedom before surveillance (scalar or vector)
<b>pfree</b>	desired probability of freedom (scalar or vector)

**Value**

a vector of population-level sensitivities

**Examples**

```
# examples of sep.pfree
sep.pfree(0.5, 0.95)
sep.pfree(c(0.5, 0.6, 0.7, 0.8, 0.9, 0.95), 0.99)
sep.pfree(0.5, c(0.8, 0.9, 0.95, 0.99))
```

sep.pooled

*Pooled population sensitivity***Description**

Calculates population sensitivity (sep) and population specificity (spp) assuming pooled sampling and allowing for imperfect sensitivity and specificity of the pooled test

**Usage**

```
sep.pooled(r, k, pstar, pse, psp = 1)
```

**Arguments**

- |       |   |
|-------|---|
| r     | number of pools sampled (scalar or vector)                    |
| k     | pool size (scalar or vector of same length as r)              |
| pstar | design prevalence (scalar or vector of same length as r)      |
| pse   | pool-level sensitivity (scalar or vector of same length as r) |
| psp   | pool-level specificity (scalar or vector of same length as r) |

**Value**

list of 2 elements, vector of sep values and vector of spp values

**Examples**

```
# examples for sep.pooled
sep.pooled(60, 5, 0.01, 1, 1)
sep.pooled(4, 10, 0.1, 0.9, 1)
sep.pooled(1:10*5, 5, 0.02, 0.9, 0.99)
sep.pooled(10, 5, 0.05, c(0.8, 0.9, 0.95, 0.99), 1)
```

---

<code>sep.prior</code>	<i>Population sensitivity to achieve desired prior probability of freedom</i>
------------------------	---

---

**Description**

Calculates the population sensitivity required to achieve a given value for the prior (discounted) probability of disease freedom

**Usage**

```
sep.prior(prior, p.intro)
```

**Arguments**

- |                      |  |
|----------------------|--|
| <code>prior</code>   | prior probability of freedom before surveillance (scalar or vector)                |
| <code>p.intro</code> | probability of introduction for time period (scalar or vector equal length to sep) |

**Value**

a vector of population-level sensitivities

**Examples**

```
# examples of sep.prior
sep.prior(0.95, 0.01)
sep.prior(c(0.9, 0.95, 0.98, 0.99), 0.01)
sep.prior(0.95, c(0.001, 0.005, 0.01, 0.02, 0.05))
```

---

<code>sep.rb.bin</code>	<i>Binomial risk-based population sensitivity</i>
-------------------------	---

---

**Description**

Calculates risk-based population sensitivity with a single risk factor, using binomial method (assumes a large population), allows for unit sensitivity to vary among risk strata

**Usage**

```
sep.rb.bin(pstar, rr, ppr, n, se)
```

**Arguments**

- |                    |  |
|--------------------|--|
| <code>pstar</code> | design prevalence (scalar)   |
| <code>rr</code>    | relative risk values (vector of values corresponding to the number of risk strata)             |
| <code>ppr</code>   | population proportions corresponding to rr values (vector of equal length to rr)               |
| <code>n</code>     | sample size per risk category (vector same length as rr and ppr)                               |
| <code>se</code>    | unit sensitivity, can vary among risk strata (fixed value or vector same length as rr, ppr, n) |

**Value**

list of 3 elements, a scalar of population-level sensitivity a vector of EPI values and a vector of corresponding adjusted risks

**Examples**

```
# examples for sep.rb.bin
sep.rb.bin(0.1, c(5, 3, 1), c(0.1, 0.1, 0.8), c(5, 5, 5), 0.9)
sep.rb.bin(0.1, c(5, 1), c(0.1, 0.9), c(10, 5), c(0.95, 0.9))
sep.rb.bin(0.1, c(5, 1), c(0.1, 0.9), c(10, 5), c(0.9, 0.9))
sep.rb.bin(0.01, c(5, 1), c(0.1, 0.9), c(90, 50), c(0.9, 0.9))
```

sep.rb.bin.varse

*Binomial risk-based population sensitivity for varying unit sensitivity***Description**

Calculates population sensitivity for a single risk factor and varying unit sensitivity using binomial method (assumes large population)

**Usage**

```
sep.rb.bin.varse(pstar, rr, ppr, df)
```

**Arguments**

pstar	design prevalence (scalar)
rr	relative risk values (vector of values corresponding to the number of risk strata)
ppr	population proportions corresponding to rr values (vector of equal length to rr)
df	dataframe of values for each combination of risk stratum and sensitivity level, col 1 = risk group index, col 2 = unit Se, col 3 = n (sample size for that risk group and unit sensitivity)

**Value**

list of 3 elements, a scalar of population-level sensitivity a vector of EPI values and a vector of corresponding adjusted risks

**Examples**

```
# examples for sep.rb.bin.varse
rg<- c(1, 1, 2, 2)
se<- c(0.92, 0.85, 0.92, 0.85)
n<- c(80, 30, 20, 30)
df<- data.frame(rg, se, n)
sep.rb.bin.varse(0.01, c(5, 1), c(0.1, 0.9), df)
```

```

rg<- c(1, 1, 2, 2)
se<- c(0.95, 0.8, 0.95, 0.8)
n<- c(20, 10, 10, 5)
df<- data.frame(rg, se, n)
sep.rb.bin.varse(0.05, c(3, 1), c(0.2, 0.8), df)

rg<- c(rep(1, 30), rep(2, 15))
se<- c(rep(0.95, 20), rep(0.8, 10), rep(0.95, 10), rep(0.8, 5))
n<- rep(1, 45)
df<- data.frame(rg, se, n)
sep.rb.bin.varse(0.02, c(3, 1), c(0.2, 0.8), df)

rg<- c(1, 2, 3, 1, 2, 3)
se<- c(0.95, 0.95, 0.95, 0.8, 0.8, 0.8)
n<- c(20, 10, 10, 30, 5, 5)
df<- data.frame(rg, se, n)
sep.rb.bin.varse(0.01, c(5, 3, 1), c(0.1, 0.3, 0.6), df)

```

**sep.rb.hypergeo***Hypergeometric risk-based population sensitivity***Description**

Calculates risk-based population sensitivity with a single risk factor, using the hypergeometric method (assuming a finite and known population size), allows for unit sensitivity to vary among risk strata

**Usage**

```
sep.rb.hypergeo(pstar, rr, N, n, se)
```

**Arguments**

pstar	design prevalence (scalar)
rr	relative risk values (vector of values corresponding to the number of risk strata)
N	Population size per risk category (vector same length as rr and ppr)
n	sample size per risk category (vector same length as rr and ppr)
se	unit sensitivity, can vary among risk strata (fixed value or a vector the same length as rr, ppr, n)

**Value**

list of 3 elements, a scalar of population-level sensitivity a vector of EPI values and a vector of corresponding adjusted risks

## Examples

```
# examples for sep.rb.bin
sep.rb.hypergeo(0.1, c(5, 3, 1), c(10, 10, 80), c(5, 5, 5), 0.9)
sep.rb.hypergeo(0.1, c(5, 1), c(15, 140), c(10, 5), c(0.95, 0.9))
sep.rb.hypergeo(0.1, c(5, 1), c(23, 180), c(10, 5), c(0.9, 0.9))
sep.rb.hypergeo(0.01, c(5, 1), c(100, 900), c(90, 50), c(0.9, 0.9))
```

`sep.rb.hypergeo.varse` *Hypergeometric risk-based population sensitivity for varying unit sensitivity*

## Description

Calculates population sensitivity for a single risk factor and varying unit sensitivity using hypergeometric approximation method (assumes known population size)

## Usage

```
sep.rb.hypergeo.varse(pstar, rr, N, df)
```

## Arguments

pstar	design prevalence (scalar)
rr	relative risk values (vector of values corresponding to the number of risk strata)
N	vector of population size for each risk group, corresponding to rr values (vector of equal length to rr)
df	dataframe of values for each combination of risk stratum and sensitivity level, col 1 = risk group index, col 2 = unit Se, col 3 = n (sample size for risk group and unit sensitivity)

## Value

list of 5 elements, a scalar of population-level sensitivity a vector of EPI values, a vector of corresponding Adjusted risks a vector of sample sizes (n) per risk group and a vector of mean unit sensitivities per risk group

## Examples

```
# examples for sep.rb.hypergeo.varse
rg<- c(1, 1, 2, 2)
se<- c(0.92, 0.85, 0.92, 0.85)
n<- c(80, 30, 20, 30)
df<- data.frame(rg, se, n)
sep.rb.hypergeo.varse(0.01, c(5, 1), c(200, 1800), df)

rg<- c(1, 1, 2, 2)
se<- c(0.95, 0.8, 0.95, 0.8)
```

```

n<- c(20, 10, 10, 5)
df<- data.frame(rg, se, n)
sep.rb.hypergeo.varse(0.05, c(3, 1), c(100, 400), df)

rg<- c(rep(1, 30), rep(2, 15))
se<- c(rep(0.95, 20), rep(0.8, 10), rep(0.95, 10), rep(0.8, 5))
n<- rep(1, 45)
df<- data.frame(rg, se, n)
sep.rb.hypergeo.varse(0.02, c(3, 1), c(100, 400), df)

rg<- c(1, 2, 3, 1, 2, 3)
se<- c(0.95, 0.95, 0.95, 0.8, 0.8, 0.8)
n<- c(20, 10, 10, 30, 5, 5)
df<- data.frame(rg, se, n)
sep.rb.hypergeo.varse(0.01, c(5, 3, 1), c(100, 300, 600), df)

```

**sep.rb2.binom***Binomial risk-based population sensitivity for 2 risk factors***Description**

Calculates risk-based population sensitivity for two risk factors, using binomial method (assumes a large population)

**Usage**

```
sep.rb2.binom(pstar, rr1, ppr1, rr2, ppr2, n, se)
```

**Arguments**

pstar	design prevalence (scalar)
rr1	relative risks for first level risk factor (vector of values corresponding to the number of risk strata)
ppr1	population proportions for first level risk factor (vector of same length as rr1)
rr2	relative risks for second level risk factor, matrix, rows = levels of rr1, cols = levels of rr2
ppr2	population proportions for second level risk factor, matrix, rows = levels of rr1, cols = levels of rr2
n	matrix of number tested for each risk group (rows = levels of rr1, cols = levels of rr2)
se	test unit sensitivity (scalar)

**Value**

list of 4 elements, a scalar of population-level sensitivity a matrix of EPI values, a vector of corresponding Adjusted risks for the first risk factor and a matrix of adjusted risks for the second risk factor

## Examples

```
# examples for sep.rb2.binom
pstar<- 0.01
rr1<- c(3, 1)
ppr1<- c(0.2, 0.8)
rr2<- rbind(c(4,1), c(4,1))
ppr2<- rbind(c(0.1, 0.9), c(0.3, 0.7))
se<- 0.8
n<- rbind(c(50, 20), c(20, 10))
sep.rb2.binom(pstar, rr1, ppr1, rr2, ppr2, n, se)
```

sep.rb2.hypergeo

*Hypergeometric risk-based population sensitivity for 2 risk factors*

## Description

Calculates risk-based population sensitivity for two risk factors, using hypergeometric approximation method (assumes a known population size)

## Usage

```
sep.rb2.hypergeo(pstar, rr1, rr2, N, n, se)
```

## Arguments

pstar	design prevalence (scalar)
rr1	relative risks for first level risk factor (vector of values corresponding to the number of risk strata)
rr2	relative risks for second level risk factor, matrix, rows = levels of rr1, cols = levels of rr2
N	matrix of population size for each risk group (rows = levels of rr1, cols = levels of rr2)
n	matrix of number tested (sample size) for each risk group (rows = levels of rr1, cols = levels of rr2)
se	test unit sensitivity (scalar)

## Value

list of 6 elements, a scalar of population-level sensitivity a matrix of EPI values, a vector of corresponding Adjusted risks for the first risk factor and a matrix of adjusted risks for the second risk factor, a vector of population proportions for the first risk factor and a matrix of population proportions for the second risk factor

## Examples

```
# examples for sep.rb2.hypergeo
pstar<- 0.01
rr1<- c(3, 1)
rr2<- rbind(c(4,1), c(4,1))
N<- rbind(c(100, 500), c(300, 1000))
n<- rbind(c(50, 20), c(20, 10))
se<- 0.8
sep.rb2.hypergeo(pstar, rr1, rr2, N, n, se)
```

sep.sys

*2-stage population sensitivity*

## Description

Calculates population-level (system) sensitivity for representative 2-stage sampling (sampling of clusters and units within clusters), assuming imperfect test sensitivity and perfect test specificity

## Usage

```
sep.sys(H = NA, N = NA, n, pstar.c, pstar.u, se = 1)
```

## Arguments

H	population size = number of clusters in the population, default = NA
N	population size within clusters, scalar or a vector of same length as n, default = NA
n	sample size (vector of number tested per cluster)
pstar.c	cluster (herd) level design prevalence, scalar, either proportion or integer
pstar.u	unit (animal) level design prevalence, scalar, either proportion or integer
se	unit sensitivity of test (proportion), scalar, default = 1

## Value

list of 6 elements, 1) population level sensitivity, 2) vector of cluster-level sensitivities, 3) N, 4) n, 5) vector of design prevalences and 6) unit sensitivity

## Note

if pstar.c is not a proportion N must be provided (and N>=n)

## Examples

```
# examples for sep.sys - checked
H<- 500
N<- rep(1000, 150)
N[5]<- NA
n<- rep(30, 150)
pstar.u<- 0.1
pstar.c<- 0.01
se<- 0.98
sep.sys(H, N, n, pstar.c, pstar.u, se)
sep.sys(NA, N, n, 0.02, 0.05, 0.95)
N<- round(runif(105)*900+100)
n<- round(runif(105)*30+10)
sse<- sep.sys(1000, N, n, 0.02, 0.05, 0.9)
data.frame(N, n, sse[[2]])
```

sep.var.se

*Population sensitivity for varying unit sensitivity*

## Description

Calculates population-level sensitivity where unit sensitivity varies and using the appropriate method, depending on whether or not N provided (hypergeometric if N provided, binomial otherwise), assuming perfect test specificity and representative sampling

## Usage

```
sep.var.se(N = NA, se, pstar)
```

## Arguments

N	population size (number of units or clusters), N must be >= length(se)) or NA if unknown
se	vector of unit sensitivity values (proportion) for each unit sampled
pstar	specified design prevalence (scalar)

## Value

a scalar of population-level sensitivity

## Examples

```
# examples of sep.var.se - checked
sens<- c(rep(0.9, 50), rep(0.95, 100))
sep.var.se(NA, sens, 0.01)
sep.var.se(se=sens, pstar=0.01)
sep.var.se(N=500, sens, 0.01)
sep.var.se(NA, runif(150, 0.95, 0.99), 0.02)
sep.var.se(500, runif(150, 0.95, 0.99), 0.02)
```

**sp.parallel**                  *Specificity of tests in parallel*

### Description

Calculates the combined specificity for multiple tests interpreted in parallel (assuming independence)

### Usage

```
sp.parallel(sp)
```

### Arguments

sp                  vector of unit specificity values

### Value

scalar of combined specificity, assuming independence

### Examples

```
# examples for sp.parallel
sp.parallel(c(0.99, 0.95, 0.8))
```

**sp.series**                  *Specificity of tests in series*

### Description

Calculates the combined specificity for multiple tests interpreted in series (assuming independence)

### Usage

```
sp.series(sp)
```

### Arguments

sp                  vector of unit specificity values

### Value

scalar of combined specificity, assuming independence

### Examples

```
# examples for sp.series
sp.series(c(0.99, 0.95, 0.8))
```

sph.binom

*Binomial population specificity for imperfect test***Description**

Calculates population specificity for a large or unknown population, using the Binomial distribution and adjusting for cut-point number of positives

**Usage**

```
sph.binom(n, c = 1, sp)
```

**Arguments**

- |    |  |
|----|--|
| n  | sample size (scalar or vector)   |
| c  | The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive (scalar or vector of same length as n) |
| sp | test unit specificity (scalar or vector of same length as n)   |

**Value**

a vector of population-level specificities

**Examples**

```
# examples for sph.imperfect.sp
sph.binom(30, 2, 0.98)
sph.binom(30, 1, 0.98)
sph.binom(1:5*10, 2, 0.98)
sph.binom(100, 1:5, 0.98)
sph.binom(100, 3, 95:100/100)
sph.binom(c(5, 10, 15, 20, 30, 50, 100, 200), 2, 0.98)
```

sph.hp

*Hypergeometric population specificity calculation***Description**

Calculates population specificity for a finite population and imperfect test, using Hypergeometric distribution

**Usage**

```
sph.hp(N, n, c = 1, sp)
```

**Arguments**

N	population size (scalar or vector of same length as n)
n	sample size (scalar or vector)
c	The cut-point number of positives to classify a cluster as positive, default=1, if positives < c result is negative, >= c is positive (scalar or vector of same length as n)
sp	test unit specificity (scalar or vector of same length as n)

**Value**

a vector of population-level specificities

**Examples**

```
# examples of sph.hp
sph.hp(150, 30, 2, 0.98)
sph.hp(150, 30, 1, 0.98)
sph.hp(150, 1:5*10, 2, 0.98)
sph.hp(500, 30, 2, 95:100/100)
```

spp	<i>Population specificity</i>
-----	-------------------------------

**Description**

Calculates population specificity assuming representative sampling

**Usage**

```
spp(n, sp)
```

**Arguments**

n	sample size (number tested), integer, scalar or vector
sp	unit specificity of test (proportion), scalar or vector of same length as n

**Value**

a vector of population-level specificities

**Examples**

```
# examples for spp - checked
spp(10, 0.9)
spp(c(10, 20, 50, 100), 0.99)
spp(100, c(0.999, 0.99, 0.98, 0.95, 0.9))
```

---

sse.combined*System sensitivity by combining multiple surveillance components*

---

## Description

Calculates overall system sensitivity for multiple components, accounting for lack of independence (overlap) between components

## Usage

```
sse.combined(C = NA, pstar.c, rr, ppr, sep)
```

## Arguments

C	population sizes (number of clusters) for each risk group, NA or vector of same length as rr
pstar.c	cluster level design prevalence (scalar)
rr	cluster level relative risks (vector, length equal to the number of risk strata)
ppr	cluster level population proportions (optional), not required if C is specified (NA or vector of same length as rr)
sep	sep values for clusters in each component and corresponding risk group. A list with multiple elements, each element is a data frame of sep values from a separate component, first column= clusterid, 2nd =cluster-level risk group index, 3rd col = sep

## Value

list of 2 elements, a matrix (or vector if C not specified) of population-level (surveillance system) sensitivities (binomial and hypergeometric and adjusted vs unadjusted) and a matrix of adjusted and unadjusted component sensitivities for each component

## Examples

```
# example for sse.combined (checked in excel combined components.xlsx)
C<- c(300, 1200)
pstar<- 0.01
rr<- c(3,1)
ppr<- c(0.2, 0.8)
comp1<- data.frame(id=1:100, rg=c(rep(1,50), rep(2,50)), cse=rep(0.5,100))
comp2<- data.frame(id=seq(2, 120, by=2), rg=c(rep(1,25), rep(2,35)), cse=runif(60, 0.5, 0.8))
comp3<- data.frame(id=seq(5, 120, by=5), rg=c(rep(1,10), rep(2,14)), cse=runif(24, 0.7, 1))
sep<- list(comp1, comp2, comp3)
sse.combined(C, pstar, rr, sep = sep)
sse.combined(C=NA, pstar, rr, ppr, sep = sep)
```

sse.rb.2stage

*Two-stage risk-based system sensitivity*

## Description

Calculates system sensitivity for 2 stage risk-based sampling, allowing for a single risk factor at each stage and using either binomial or hypergeometric approximation

## Usage

```
sse.rb.2stage(C = NA, pstar.c, pstar.u, rr.c, ppr.c, rr.u, ppr.u,
N = NA, n, rg, se)
```

## Arguments

C	Population size (number of clusters), NA = unknown (default)
pstar.c	cluster level design prevalence (scalar)
pstar.u	unit level design prevalence (scalar)
rr.c	cluster level relative risks (vector with length corresponding to the number of risk strata), use rr.c = c(1,1) if risk factor does not apply
ppr.c	cluster level population proportions for risk categories (vector), NA if no cluster level risk factor
rr.u	unit level relative risks (vector with length corresponding to the number of risk strata), use rr.u = c(1,1) if risk factor does not apply
ppr.u	unit level population proportions for each risk group (optional) matrix, 1 row for each cluster, columns = unit level risk groups, not required if N is provided
N	population size per risk group for each cluster, NA or matrix of N for each risk group for each cluster, N=NA means cluster sizes not provided
n	sample size per risk group for each cluster sampled, matrix, 1 row for each cluster, columns = unit level risk groups
rg	vector of cluster level risk group (index) for each cluster
se	unit sensitivity for each cluster, scalar or vector of values for each cluster, equal in length to n

## Value

list of 2 elements, a scalar of population-level (surveillance system) sensitivity and a vector of cluster-level sensitivities

## Examples

```
# examples for sse.rb.2stage
pstar.c<- 0.02
pstar.u<- 0.1
rr.c<- c(5, 1)
ppr.c<- c(0.1, 0.9)
rr.u<- c(3, 1)
se<- 0.9
n<- cbind(rep(10, 50), rep(5, 50))
rg<- c(rep(1, 30), rep(2, 20))
ppr.u<- cbind(rep(0.2, 50), rep(0.8, 50))
N<- cbind(rep(30, 50), rep(120, 50))
C<- 500
sse.rb.2stage(C=NA, pstar.c, pstar.u, rr.c, ppr.c, rr.u, ppr.u, N=NA, n, rg, se)
sse.rb.2stage(C, pstar.c, pstar.u, rr.c, ppr.c, rr.u, ppr.u, N=NA, n, rg, se)
sse.rb.2stage(C=NA, pstar.c, pstar.u, rr.c, ppr.c, rr.u, ppr.u, N, n, rg, se)
sse.rb.2stage(C, pstar.c, pstar.u, rr.c, ppr.c, rr.u, ppr.u, N, n, rg, se)
```

tp

*True prevalence*

## Description

Estimates true prevalence and confidence limits for given sample size and result, according to specified method

## Usage

```
tp(x, n, se, sp, type = "blaker", conf = 0.95)
```

## Arguments

x	number of positive units (scalar)
n	sample size (no. units sampled) (scalar)
se	test sensitivity (scalar)
sp	test specificity (scalar)
type	method for estimating CI, one of c("normal", "c-p", "sterne", "blaker", "wilson", "all")
conf	desired level of confidence for CI, default = 0.95 (scalar)

## Value

list with 2 elements, a matrix of apparent prevalence and lower and upper confidence limits and a matrix of true prevalence and lower and upper confidence limits using the chosen method(s)

### Examples

```
# examples for tp
x<- 20
n<- 120
se<- 0.9
sp<- 0.99
conf<- 0.95
tp(x, n, se, sp, "all")
tp(x, n, se, sp, "c-p")
tp(x, n, 0.95, 0.9, "c-p")
```

**tp.normal**

*Normal approximation confidence limits for true prevalence*

### Description

Estimates true prevalence and confidence limits for estimates based on normal approximation

### Usage

```
tp.normal(x, n, se, sp, conf = 0.95)
```

### Arguments

x	number of positive results in sample (scalar or vector)
n	sample size (scalar or vector)
se	test unit sensitivity (scalar or vector)
sp	test unit specificity (scalar or vector)
conf	desired level of confidence for CI, default = 0.95 (scalar or vector)

### Value

list with 2 elements, a matrix of apparent prevalence and wilson lower and upper confidence limits and a matrix of true prevalence and normal approximation lower and upper confidence limits

### Examples

```
# examples for tp.normal
tp.normal(25, 120, 0.9, 0.99)
tp.normal(seq(5, 25, by=5), 120, 0.9, 0.99)
```

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