# Package 'DiagTestKit'

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<b>Description</b> A package written by CVB Statistics to estimate the sensitivity and specificity of an experimental diagnostic test kit in accordance with CVB STATWI0002 supporting the 2018 revision to VSM 800.73.			
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DiagTestKit-package DiagTestKit Package.

## **Description**

A package written by CVB Statistics to estimate the sensitivity and specificity of an experimental diagnostic test kit in accordance with CVB STATWI0002 supporting the 2018 revision to VSM 800.73.

## **Details**

Functions used in evaluating sensitivity and specificity at CVB Statistics

Package: DiagTestKit-package

Type: Package Version: 0.5.3 Date: 26 July 2018

License: MIT LazyLoad: yes LazyData: yes

## Resources

- GUIDANCE: https://www.aphis.usda.gov/aphis/ourfocus/animalhealth/veterinary-biologics/biologics-regulations-and-guidance/ct\_vb\_statwi
- $\bullet \ QUICK \ START: \ https://github.com/ABS-dev/DiagTestKit/blob/master/README.md$
- BUG REPORTS: https://github.com/ABS-dev/DiagTestKit/issues

## Author(s)

CVB Statistics <CVB.Data.Help@aphis.usda.gov>

betaParm Convert Beta Parameterizations

#### **Description**

Convert between the paramaterizations of a beta distribution.

#### Usage

```
betaParm(B, to = "alpha.beta")
```

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#### **Arguments**

B vector A named vector specifying non-NULL values for 2 parameters. e.g.

c(alpha=NA, beta=NA, mu=.6, theta=NA, phi=1.6, sigma2=NA) or just c(mu=.6,

phi=1.6)

to Specification of desired parameters, options are one of 'alpha.beta', 'mu.phi',

'mu.theta' or 'mu.sigma2'.

#### Value

vector A named vector with values for the parameters specified in the 'to' argument of the input.

#### Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

cellS Obtain cell counts

#### **Description**

This function creates expected cell counts (and probabilities) for a specific test pattern based on the diagnostic characteristics of the reference test(s) and experimental test.

## Usage

```
cellS(SnR, SpR, Prev, SnE, SpE, sus.perc, N, nstates)
```

#### **Arguments**

SnR data.frame Each column corresponds to one reference test. Row 1 contains the

sensitivity for the reference test(s). Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely,  $P(T? \mid D+) = \psi = \delta * (1 - \pi)$  where  $\delta$  is the second row for a given column (reference test).  $\delta = \frac{\psi}{(1-\pi)}$ . Use a zero for a 2-state test (i.e. no

suspect region).

SpR data.frame Each column corresponds to one reference test. Row 1 contains the

specificity for each reference test. Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, P(T? | D-) =  $\phi = \gamma * (1 - \theta)$  where  $\gamma$  is the second row for a given column (reference test).  $\gamma = \frac{\phi}{(1-\theta)}$ . Use a zero for a 2-state test (i.e. no

suspect region).

Prev vector A named vector containing the prevalence for each population sampled.

SnE Sensitivity of the experimental test kit.

SpE Specificity of the experimental test kit.

sus.perc vector A vector containing 2 elements,  $c(\delta, \gamma)$  for the experimental test kit. A

vector of zeros for a 2-state experimental kit.  $\delta$  and  $\gamma$  are values between 0 and 1 (inclusive) corresponding to the proportion of the remaining probability (i.e.

1 -  $\pi$  or 1 -  $\theta$ ) that is suspect ( $\psi$  or  $\phi$ ).  $\delta = \frac{\psi}{(1-\pi)}$  and  $\gamma = \frac{\phi}{(1-\theta)}$ .

N vector A named vector containing the sample size for each population sampled.

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nstates

vector A vector with length one greater than the number of reference tests. The first element is the number of states of the experimental test and the remaining entries are the number of states of each reference test (using the same ordering as SnR and SpR).

## Value

vector A vector of expected counts corresponding to the properties of the reference and experimental tests. The expected counts are obtained based on a conditional independence assumption of all test methods.

#### Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

cloppearSnSp Binomial confidence interval, Clopper-Pearson method.	loppearSnSp	Binomial confidence interval, Clopper-Pearson method.
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## **Description**

Evaluate binomial confidence interval using Clopper-Pearson method. A function written by CVB Statistics to estimate the sensitivity and specificity of an experimental diagnostic test kit in accordance with CVB STATWI0002.

#### Usage

```
cloppearSnSp(dat, alpha = 0.05, est.Sn = TRUE)
```

## **Arguments**

dat	data.frame A data frame with a column for the experimental test results, a col-
	umn for the infallible reference test results, and a column for the corresponding
	count. The column name for the experimental test results must contatin 'exp'
	and the column name for the infallible reference test results must include 'ref'.
	The counts should be the last column.

alpha Complement of confidence level.

 $\verb| est.Sn| \qquad \qquad | logical (TRUE/FALSE) \ Indicating \ if \ the \ sensitivity \ and \ its \ confidence \ interval$ 

should be supplied (TRUE) or if the specificity and its confidence interval should

be supplied (FALSE).

#### Value

An object of type cp that extends list.

calcVal Named vector of point estimates and estimated simulated intervals. See below.

data Test and Total values of the data. See below.

**alpha** Complement of the confidence interval as provided above.

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If est.Sn == TRUE

calcVal is a list with the following elements

- Sn Sensitivity estimate.
- Sn.LL Lower confidence limit for sensitivity.
- Sn.UL Upper confidence limit for sensitivity.

data is a list with the following elements

- Test.Positive Number of experimental test positives.
- Total.Positive Total number of positive samples.

```
If est.Sn == FALSE
```

calcVal is a list with the following elements

- Sp Specificity estimate.
- Sp.LL Lower confidence limit for specificity.
- Sp.UL Upper confidence limit for specificity.

data is a list with the following elements

- Test.Negative Number of experimental test negatives.
- Total.Negative Total number of negative samples.

## Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

#### References

Clopper CJ, Pearson ES, 1934. The use of confidence or fiducial limits illustrated in the case of the binomial. *Biometrika* 26:404-413.

## **Examples**

```
CP.Sn <- cloppearSnSp(dat = dat_infal, est.Sn = TRUE)
CP.Sn
# Sn = P(T+|D+): 0.987013 (95% CI: 0.953876, 0.998423)
CP.Sp <- cloppearSnSp(dat = dat_infal, est.Sn = FALSE)
CP.Sp
# Sp = P(T-|D-): 0.970297 (95% CI: 0.915643, 0.915643)</pre>
```

emp.hpd

Calculate the empirical hpd.

#### **Description**

Empirical highest posterior density by shortest length interval.

#### Usage

```
emp.hpd(X, alpha)
```

#### **Arguments**

X vector of values alpha 1 - confidence

#### Value

highest posterior density (1-alpha) interval

#### Note

Uses type 7 quantile. Also used in package MF

#### Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

estimateSnSp

Estimate Sensitivity and Specificity

# Description

A function written by CVB Statistics to estimate the sensitivity and specificity of an experimental diagnostic test kit in accordance with CVB STATWI0002.

## Usage

```
estimateSnSp(dat, Sn.ref, Sp.ref, prev.pop, nsim = 1000, control = NULL)
```

## **Arguments**

dat

data.frame This is a data frame where the first column includes information for the population sampled (if more than one population is sampled). The next column is the possible outcomes of the experimental test followed by one column for the possible outcomes for each reference test (one column per test). The last column of the data frame provides the number of samples with each pattern of test outcomes. The columns must be included in the order described. If more than one population is sampled, the column name for the column containing the population information must be 'population'. The column containing the test

results for the experimental test must have 'exp' in the name, such as experimental, experiment, exp, Exp, etc. The column names containing the reference test results much contain 'ref' in the name, such as Ref1, Ref2, ref1\_results, Reference2, etc.

Sn.ref

data.frame Each column corresponds to one reference test. Row 1 contains the sensitivity for the reference test(s). Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely,  $P(T? \mid D+) = \psi = \delta * (1-\pi)$  where  $\delta$  is the second row for a given column (reference test).  $\delta = \frac{\psi}{(1-\pi)}$ . Use a zero for a 2-state test (i.e. no suspect region). Alternatively, if all reference tests are 2-state tests, the sensitivities can be input as a named vector. Specifically, each element in the vector must be given a name which includes 'ref' (see above) and the column names (or names of the elements within the vector) must match those for Sp.ref.

Sp.ref

data. frame Each column corresponds to one reference test. Row 1 contains the specificity for each reference test. Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, P(T? | D-) =  $\phi = \gamma * (1 - \theta)$  where  $\gamma$  is the second row for a given column (reference test).  $\gamma = \frac{\phi}{(1-\theta)}$ . Use a zero for a 2-state test (i.e. no suspect region). Alternatively, if all reference tests are 2-state tests, the specificities can can be input as a named vector. Specifically, each element in the vector must be given a name which includes 'ref' (see above) and the column names (or names of the elements within the vector) must match those for Sn.ref.

prev.pop

vector A named vector containing the prevalence for each population sampled. The names in the vector must match the population labels used in 'dat'.

nsim

The number of simulations to draw from the sensitivity and specificity distribution(s) for each reference test and the prevalence distribution from each population.

control

list of control values to replace defaults. See estimateSnSpControl for details.

#### Value

An object of type snsp that extends list.

calcVal Point estimates and estimated simulated intervals for properties of the experimental kit. See below.

detailOut Detailed output values. See below.

input Simulated values. See below.

## calcVal

A list with the following values which will include the following for both 2- and 3-state experimental tests –

- Nsim Number of simulations performed.
- Confidence 1  $\alpha$ .
- SnPE Sensitivity point estimate obtained as the median of the estimated values.
- SnInterval Estimated simulated interval for sensitivity.

- SpPE Specificity point estimate obtained as the median of the estimated values.
- **SpInterval** Estimated simulated interval for specificity.

If three states, the list will also include –

- **SusDisPosPE** Point estimate for the probability of test suspect given disease positive  $(\psi)$  which is the median of the calculated values  $(\psi = \delta^*(1-\pi))$ .
- **SusDisPosInterval** Estimated simulated interval for the probability of test suspect given disease positive  $(\psi)$ .
- **SusDisNegPE** Point estimate for the probability of test suspect given disease negative  $(\phi)$  which is the median of the calculated values  $(\phi = \gamma^*(1-\theta))$ .
- **SusDisNegInterval** Estimated simulated interval for the probability of test suspect given disease negative  $(\phi)$ .

#### detailOut

A list with the following detailed output values which will include the following for both 2- and 3-state experimental tests –

- Exp.Sn vector The optimized values for the sensitivity of the experimental test kit.
- Exp.Sp vector The optimized values for the specificity of the experimental test kit.
- **Converge** vector Each entry is an integer code detailing the convergence of the optimization for each iteration. 0 indicates successful completion. See also optim.
- **Message** vector Each entry includes a character string providing any additional information returned by the optimizer or NULL. See also optim.

If three states, the list will also inlcude –

- **Exp.pos.p** vector The optimized values for the proportion of the remaining probability (1-Sn) that corresponds to a suspect region for diseased samples, namely  $\delta$ .
- **Exp.sus.pos** vector The values for P(T? | D+) ( $\psi$ ) calculated from Exp.sn and Exp.pos.p. P(T?|D+) =  $\delta * (1 \pi)$ .
- **Exp.neg.p** vector The optimized value for the proportion of the remaining probability (1-Sp) that corresponds to a suspect region for non-diseased samples, namely  $\gamma$ .
- **Exp.sus.neg** vector The values for P(T? | D-) ( $\phi$ ) calculated from Exp.sp and Exp.neg.p. P(T?|D-) =  $\gamma$  \* (1  $\theta$ ).

#### input

A list containing the seed used and the simulated values.

- **seed** The seed used in the random generation of the distributions of sensitivity and specificity for all reference tests and prevalence of each population. See also **set.seed**
- Sn.sims matrix The simulated values for the sensitivity of each reference test and  $\psi$  where  $\psi$  was specified in the second row of Sn.ref (or zero if Sn.ref was a vector). The first two columns correspond to the first reference test, columns 3 and 4 to the second reference test if it exists, etc.

• **Sp.sims** matrix The simulated values for the specificity of each reference test and  $\phi$  where  $\phi$  was specified in the second row of Sp.ref (or zero is Sp.ref was a vector). The first two columns correspond to the first reference test, columns 3 and 4 to the second reference test if it exists, etc.

• **prev.sims** matrix The simulated values of prevalence for each population. Each column correspond to one population.

#### Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

#### See Also

estimateSnSpControl

## **Examples**

```
data.1 <- data.frame(exp_result = rep(c('positive', 'negative'), each = 2),</pre>
                      ref1_result = rep(c('positive', 'negative'), 2),
                      count = c(82, 11, 5, 22))
example.1 <- estimateSnSp(dat = data.1, Sn.ref = data.frame(ref = c(0.90, 0)),
                      Sp.ref = data.frame(ref=c(0.99, 0)), prev.pop=c(A=0.80),
                      control = estimateSnSpControl(seed = 64725))
example.1
# 1000 simulations
# 95 % Interval Estimates
#
                Point.Estimate
                                    Lower Upper
\# Sn = P(T+|D+)
                     0.9449821 0.9019639
\# Sp = P(T-|D-)
                      0.9062769 0.7523346
## Not run:
data.2 <- data.frame(Population = rep(LETTERS[1:3], each = 24),</pre>
                 exp_result = rep(rep(c('negative', 'positive', 'suspect'), each = 8), 3),
                      ref1_result = rep(rep(c('negative', 'positive'), each = 4), 9),
                      ref2_result = rep(rep(c('negative', 'positive'), each = 2), 18),
                      ref3_result = rep(c('negative', 'positive'), 36),
                      count = c(3, 0, 0, 0, 1, 0, 0, 1, 0, 1, 1, 5, 1, 8, 11, 62,
                                0, 0, 0, 0, 0, 0, 0, 2, 27, 2, 3, 0, 4, 0, 1, 1, 0,
                                0,\ 1,\ 4,\ 1,\ 6,\ 7,\ 41,\ 0,\ 0,\ 0,\ 0,\ 0,\ 0,\ 0,\ 2,\ 57,\ 5,
                                6,\ 1,\ 9,\ 1,\ 1,\ 0,\ 0,\ 0,\ 1,\ 0,\ 2,\ 2,\ 12,\ 1,\ 0,\ 0,
                                0, 0, 0, 0, 0))
example.2 <- estimateSnSp(dat = data.2,</pre>
                  Sn.ref = data.frame(ref1 = c(0.92, 0), ref2 = c(0.88, 0), ref3 = c(0.85, 0)),
                  Sp.ref = data.frame(ref1 = c(0.86, 0), ref2 = c(0.90, 0), ref3 = c(0.92, 0)),
                           prev.pop = c(A = 0.95, B = 0.62, C = 0.18),
                           control = estimateSnSpControl(seed = 865213))
# 1000 simulations
# 95 % Interval Estimates
                  Point.Estimate
                                     Lower
                                                 Upper
\# Sn = P(T+|D+)
                      0.96541704 0.8879949 1.00000000
\# Sp = P(T-|D-)
                      0.98351924 0.9016964 1.00000000
\# SsP = P(T?|D+)
                      0.02568427 0.0000000 0.06339616
```

```
# SsN = P(T?|D-) 0.01534125 0.0000000 0.05604950 ## End(Not run)
```

 ${\tt estimateSnSpControl}$ 

control values for estimateSnSp

## **Description**

The values supplied in the function—call replace the defaults and a list with all possible arguments is returned. The returned list is used as the control argument to the function estimateSnSp.

## Usage

```
estimateSnSpControl(seed = NULL, Sn.distn = NULL, Sn.spread = NULL,
    Sp.distn = NULL, Sp.spread = NULL, prev.distn = NULL,
    prev.spread = NULL, tolerance = 0.001, alpha = 0.05,
    step.size = 1e-06, parm = NULL, rep.iter = TRUE, iter.n = 50)
```

## **Arguments**

seed	The seed used in the random generation of the distributions of sensitivity and specificity for all reference tests and prevalence of each population. See also set.seed.
Sn.distn	vector A named vector with length equal to the number of reference tests. Determines which disibution should be used for sampling sensitivity of each reference test. Inputs are 'beta' or 'triangular'. Defaults to 'beta' for each reference test.
Sn.spread	vector A named vector with length equal to the number of reference tests. Describes the width of the distribution for the sensitivity of each reference test. Inputs are 'wide', 'medium', or 'narrow'. Defaults to 'wide' for each reference test.
Sp.distn	vector A named vector with length equal to the number of reference tests. Determines which disibution should be used for sampling specificity of each reference test. Inputs are 'beta' or 'triangular'. Defaults to 'beta' for each reference test.
Sp.spread	vector A named vector with length equal to the number of reference tests. Describes the width of the distribution for the specificity of each reference test. Inputs are 'wide', 'medium', or 'narrow'. Defaults to 'wide' for each reference test.
prev.distn	vector A named vector with length equal to the number of populations. Determines which disibution should be used for sampling the prevalence of each population. Inputs are 'beta' or 'triangular'. Defaults to 'beta'.
prev.spread	vector A named vector with length equal to the number of populations. Describes the width of the distribution for the prevalence of each population. Inputs are 'wide', 'medium', or 'narrow'. Defaults to 'wide' for each population.
tolerance	Setting a limit on the pgtol used in the optim function with the 'L-BFGS-B' method. See also optim. Defaults to 1E-03.
alpha	Significance levels. Defaults to 0.05.

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step.size Provides the level of resolution in values simulated from a triangular distribution. Defaults to 1E-06. vector Starting values for the optimization of the parameters of the experimenparm tal test. If the experimental test has 2 states, this vector is of length two with elements corresponding to sensitivity and specificity, respectively. If the experimental test has 3 states, this vector is of length 4 with elements corresponding to sensitivity  $(\pi)$ , the proportion of 1-Sn corresponding to the suspect region for disease positive samples ( $\delta$ ), specificity ( $\theta$ ), and the proportion of 1-Sp corresponding to the suspect region for disease negative samples ( $\gamma$ ). All values are between 0 and 1, inclusive. logical (TRUE/FALSE) Indicates if updates should be printed regarding the rep.iter number of iterations completed. Defaluts to TRUE. iter.n integer indicating the frequency of updates for the number of iterations com-

#### Value

A list with the following elements (as defined above): seed, Sn.disn, Sn.spread, Sp.distn, Sp.spread, prev.distn, prev.spread, tolerance, step.size, parm

## Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

pleted. Defaluts to 50.

## **Examples**

```
estimateSnSpControl()
estimateSnSpControl(seed = 64725)
```

```
get.simulated.values Get Simulated Values
```

#### **Description**

Simulate values for use in optimization. This function is used to obtain draws from a distribution for the sensitivity and specificity for each reference test and for the prevalence of each population tested.

## Usage

```
get.simulated.values(means, distn, spread, nsim, step.size, prevalence)
```

## Arguments

means	vector A named vector containing point estimates for the prevalence of each population (when prevalence = TRUE, see below) or a data frame where each
	column corresponds to a reference test and the rows are sensitivity $(\pi)$ and $\psi$ (or specificity $(\theta)$ and $\phi$ ).
distn	vector A vector of same length as means. Values may be one of NULL, 'beta', or 'triangular'. NULL will be treated as 'beta'

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vector A vector of same length as means. Values may be one of NULL, 'wide', spread 'medium', or 'narrow'. NULL will be treated as 'wide'. The number of simulations to draw from the sensitivity and specificity distribunsim tion(s) for each reference test or the prevalence distribution from each popula-Provides the level of resolution in values simulated from a triangular distribustep.size tion

logical (TRUE/FALSE) TRUE indicates that the function is simulating values prevalence

of prevalence. This will determine the structure of the output.

#### Value

final.mat A matrix of simulated values. If prevalence is TRUE, final.mat will have the number of columns corresponding to the number of populations sampled else if prevalence is FALSE, final.mat will have number of columns twice the number of reference tests. The columns are sensitivity (or specificity) of the first reference test, the probability of a suspect result as a fraction of the noncorrect test result (i.e. either  $\delta$  or  $\gamma$ ) for the first reference and continues in the same pattern for all reference tests.

#### Author(s)

CVB Statistics < CVB. Data. Help@aphis.usda.gov>

get.values Optimization of Sensitivity and Specificity

## **Description**

Determine final optimized values for the sensitivity and specificity of an experimental test kit (and probability of suspect given disease positive and given disease negative for a 3-state kit).

#### **Usage**

```
get.values(dat, SnR.vec, SpR.vec, prev.vec, N.vec, nstates, tolerance, rep.iter,
  iter.n, parm = NULL)
```

#### **Arguments**

dat vector A vector of counts ordered in a manner consistent with output from the

cellS function.

data. frame Each column corresponds to one reference test. Row 1 contains the SnR.vec

sensitivity for the reference test(s). Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, P(T? | D+) =  $\psi = \delta * (1 - \pi)$  where  $\delta$  is the second row for a given column (reference test).  $\delta = \frac{\psi}{(1-\pi)}$ . Use a zero for a 2-state test (i.e. no

suspect region).

SpR.vec data. frame Each column corresponds to one reference test. Row 1 contains the

> specificity for each reference test. Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, P(T? | D-) =  $\phi = \gamma * (1 - \theta)$  where  $\gamma$  is the second row for a given column (reference test).  $\gamma = \frac{\phi}{(1-\theta)}$ . Use a zero for a 2-state test (i.e. no

suspect region).

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prev.vec	vector A named vector containing the prevalence for each population sampled.
N.vec	$vector\ A\ named\ vector\ containing\ the\ sample\ size\ for\ each\ population\ sampled.$
nstates	vector A vector with length one more than the number of reference tests. The first element is the number of states of the experimental test and the remaining entries are the number of states of each reference test (using the same ordering as SnR.vec and SpR.vec).
tolerance	Setting a limit on the pgtol used in the optim function with the 'L-BFGS-B' method. See also ${\tt optim}$ .
rep.iter	logical (TRUE/FALSE) Indicates if updates should be printed regarding the number of iterations completed.
iter.n	integer indicating the frequency of updates for the number of iterations completed.
parm	vector A vector of starting values to be used for the optimization that is passed to minCell. For a 2-state experimental test, this is a vector of length 2 with entries $(\pi, \theta)$ . For a 3-state experimental test, this is a vector of length 4 with entries $(\pi, \delta, \theta, \gamma)$ . See also estimateSnSp.

#### Value

#### A list:

The following will be returned for both 2 and 3-state experimental tests –

- sens.final vector The optimized values for the sensitivity of the experimental test kit.
- spec.final vector The optimized values for the specificity of the experimental test kit.
- converge vector Each entry is an integer code detailing the convergence of the optimization for each iteration. 0 indicates successful completion. See also optim.
- message vector Each entry includes a character string giving any additional information returned by the optimizer or NULL. See also optim.

If three states -

- $\delta$  vector The optimized values for the probability of a suspect result as a fraction of the non-correct test result for diseased samples.
- $\gamma$  vector The optimized value for the probability of a suspect result as a fraction of the non-correct test result for non-diseased samples.

## Author(s)

CVB Statistics <CVB.Data.Help@aphis.usda.gov>

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|--|

# Description

A function used for optimizing the values of sensitivity and specificity (and  $\delta$  and  $\gamma$  for a 3-state kit). The objective function minimizes the sum of the squared deviations (expected - observed cell counts).

# Usage

```
minCell(parm, SnR, SpR, Prev, xdat, N, nstates)
```

# Arguments

parm	vector A vector of starting values to be used for the optimization that is passed to minCell. For a 2-state experimental test, this is a vector of length 2 with entries $(\pi, \theta)$ For a 3-state experimental test, this is a vector of length 4 with entries $(\pi, \delta, \theta, \gamma)$ . See also estimateSnSp.
SnR	data.frame Each column corresponds to one reference test. Row 1 contains the sensitivity for the reference test(s). Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, P(T?   D+) = $\psi = \delta * (1 - \pi)$ where $\delta$ is the second row for a given column (reference test). $\delta = \frac{\psi}{(1-\pi)}$ . Use a zero for a 2-state test (i.e. no suspect region).
SpR	data. frame Each column corresponds to one reference test. Row 1 contains the specificity for each reference test. Row 2 contains the probability of a suspect result as a fraction of the non-correct test result. This is a value between 0 and 1 (inclusive). Namely, $P(T? \mid D-) = \phi = \gamma * (1 - \theta)$ where $\gamma$ is the second row for a given column (reference test). $\gamma = \frac{\phi}{(1-\theta)}$ . Use a zero for a 2-state test (i.e. no suspect region).
Prev	vector A named vector containing the prevalence for each population sampled.
xdat	vector A vector of the observed cell counts.
N	vector A named vector containing the sample size for each population sampled passed to cells.
nstates	vector A vector with length one more than the number of reference tests. The first element is the number of states of the experimental test and the remaining entries are the number of states of each reference test (using the same ordering as SnR and SpR).

## Value

The sum of the squared deviations between the expected and observed cell counts.

## Author(s)

CVB Statistics <CVB.Data.Help@aphis.usda.gov>

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SampDist	Create Triangular Distribution	

# Description

Creates a discrete step/triangular distribution that can be used to sample values for sensitivity or specificity of a reference test or prevalence of a population.

## Usage

```
SampDist(m, w, h, threestate = FALSE, suspect = 2/3, stepwidth = 0.005,
   sumOne = TRUE)
```

## Arguments

m	This is a point estimate for the parameter in which you are obtaining the distribution, e.g. sensitivity, specificity, or prevalence.
W	vector A vector that provides the half widths of the 3 regions, (w1 closest, w3 farthest).
h	vector A vector of "y" (pseduo-value until scaled to be a probability) corresponding to the height of the shoulder and the height of the plateau.
threestate	logical (TRUE/FALSE) Indicates whether or not there is a "suspect" region (i.e. positive/suspect/negative).
suspect	A fraction that indicates what percentage of the remaining probability would be assigned to the suspect region. For instance, if the function gives sensitivity and then the probability of "suspect" is (1 - sensitivity)*suspect.
stepwidth	distance between the 'x' in the discrete distribution, resolution of possible observations of the created distribution.
sumOne	whether to expresss 'p' as a proportion of its sum.

## Value

```
data.frame of 'x', 'y', and 'p'.
```

## Author(s)

CVB Statistics <CVB.Data.Help@aphis.usda.gov>

updateAlpha	Update alpha values for existing simulation

## **Description**

Report interval estimates with updated alpha values, using a previously evaluated simulation.

## Usage

```
updateAlpha(x, newAlpha)
```

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## **Arguments**

```
x output from estimateSnSp
newAlpha updated alpha value. Must be within [0, 1]
```

## Value

an object of type snsp. See output for estimateSnSp

## Author(s)

CVB Statistics <CVB.Data.Help@aphis.usda.gov>

## See Also

estimateSnSpControl

## **Examples**

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