## Package 'mixIndependR'

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Type Package

Title Genetics and Independence Testing of Mixed Genetic Panels

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**Depends** R (>= 3.6.0)

**Imports** stats (>= 3.3), utils (>= 3.2.3), stringr

**Description** Developed to deal with multi-locus genotype data, this package is especially designed for those panel which include different type of markers. Basic genetic parameters like allele frequency, genotype frequency, heterozygosity and Hardy-Weinberg test of mixed genetic data can be obtained. In addition, a new test for mutual independence which is compatible for mixed genetic data is developed in this package.

License GPL (>= 2)

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AlleleFreq

Calculate Allele Frequency

## Description

Calculate Allele Frequency

#### Usage

AlleleFreq(x)

#### Arguments

х

a dataset of alleles. Type needs to be Homogeneous. Each row denotes each sample. One allele in one cell.In the (2r-1)th column, there is the other allele on the same locus from that in the 2r-th column; noted: no column for ID, make row.names=1 when importing.

## Details

This function calculates the allele frequencies of one dataset.

## Value

a matrix of allele frequencies. Each row denotes each allele; each column denotes each marker. The order of makers follows x.

AlleleShare\_Table Calculate numbers of sharing alleles each pair at each locus

## Description

Calculate numbers of sharing alleles each pair at each locus

#### Usage

AlleleShare\_Table(x,replicate=TRUE)

#### Arguments

x	a dataset of alleles. Each row denotes each individual.One allele in one cell.In the (2r-1)th column, there is the same locus with the 2r-th column; noted: no column for ID, make row.names=1 when importing.
replicate	a logical variable. if replicate is TRUE, the pairs are formed with replicates; if FALSE, the pairs are formed without replicate.

## Details

This function calculates the numbers of shared alleles between each pair of individuals for a dataset. Output a table and Usually followed by write.csv(as.data.frame(y),file = " $\sim$ /\*.csv") to export the results.

## Value

y a matrix of numbers of shared alleles. Each row denotes each pair; Each column denotes each locus.

ComposPare\_K

#### Description

Generate Comparison Observed and Expected No. of Heterozygous Loci.

## Usage

ComposPare\_K(h,Ex,trans)

## Arguments

h	a double made up of "0" and "1" where 1 means heterozygous and 0 means homozygous; Outcome of function "Heterozygous"; Each column denotes each locus and each row denotes each individual.
Ex	a dataframe of expected density, outcome of function "DistHetero", on each possible total number of heterozygous loci.
trans	a logic variable, if True, the outcome is a dataframe of n x 2. n is the number of individuals of original imported database. First column is the observed No. of Heterozygous Loci and the second is the expected one. If False, the dataframe is $2n \times 2$ , where n is the number of individuals of original imported database. The first column is a categorical variable denoting the frequency is observed or expected value; the second column is the frequency of No. of heterozygous loci.

## Details

This function generates a dataframe in which the observed and expected heterozygous loci for each sample are included. The observed ones are calculated from the original dataset. However, the expected ones are simulated according to the expected probability with the same sample size as observed sample.

#### Value

a dataframe of observed and expected No. of heterozygous loci for each individual.

```
h<-matrix(rbinom(20,1,0.5),nrow=5)
Ex <- data.frame(K=c(0:5),Density=rnorm(6,mean = 0.5,sd=0.05))
ComposPare_K(h,Ex,trans = TRUE)</pre>
```

ComposPare\_X

## Description

Generate Comparison Observed and Expected No. of Shared Alleles.

#### Usage

ComposPare\_X(AS,Ex,trans=TRUE)

## Arguments

AS	a double made up of "0", "1" and "2" denoting number of shared alleles; Outcome of function "AlleleShare_Table"; Each column denotes each locus and each row denotes each pair of individuals.
Ex	a dataframe of expected density, outcome of function "DistAlleleShare", on each possible total number of shared Alleles.
trans	a logic variable, if True, the outcome is a dataframe of n x 2. n is the number of individuals of original imported database. First column is the observed No. of Heterozygous Loci and the second is the expected one. If False, the dataframe is $2n \times 2$ , where n is the number of individuals of original imported database. The first column is a categorical variable denoting the frequency is observed or expected value; the second column is the frequency of No. of heterozygous loci.

## Details

This function generates a dataframe in which the observed and expected shared alleles for each pair of individuals. The observed ones are calculated from the original dataset through "Allele-Share\_Table". However, the expected ones are simulated according to the expected probability with the same sample size as the observed sample.

## Value

a dataframe of observed and expected No. of shared alleles for each pair of individuals.

```
AS<-matrix(sample(c(0:2),20,replace=TRUE,prob=c(0.3,0.3,0.4)),nrow=5)
Ex <- data.frame(X=c(0:8),Density=rnorm(9,mean = 0.5,sd=0.05))
ComposPare_X(AS,Ex,trans = TRUE)</pre>
```

counta

#### Description

Simple count including zero###

## Usage

counta(z, y)

## Arguments

Z	a vector you would like to check
у	an element you would like to count.(Even it is not included in z)

## Details

This function counts how many the assigned elements there are in one vector.

## Value

the times that y appears in z

## Examples

```
z <-rbinom(20,1,0.5)
counta(z,0)</pre>
```

DistAlleleShare Build Expected Distribution of Numbers of Shared Alleles

#### Description

Build Expected Distribution of Numbers of Shared Alleles

## Usage

```
DistAlleleShare(e)
```

#### Arguments

a matrix/dataframe of probability of shared alleles; outcome of "ExpProAllele-Share" or "RealProAlleleShare". Each row denotes each locus. The first column is the case of 0 shared alleles, the second column is the case of 1 shared alleles, the third column is the case of 2 shared alleles.

е

## DistHetero

#### Details

This function build the expected distribution of numbers of shared alleles for known shared alleles of each pair of individuals.

#### Value

a dataframe of probabilities of each number of shared alleles(from 0 to 2\*loci); the first column is No. of Shared Alleles; the Second Column is Expected Density

#### References

Chakraborty, R., Stivers, D. N., Su, B., Zhong, Y., & Budowle, B. (1999) <doi:10.1002/(SICI)1522-2683(19990101)20:8<1682::AID-ELPS1682>3.0.CO;2-Z>

## Examples

```
e0<-data.frame("P0"=runif(5,min = 0,max = 0.5),"P1"=runif(5,0,0.5))
e<-data.frame(e0,"P2"=1-rowSums(e0))
DistAlleleShare(e)</pre>
```

DistHetero	Build Expected Distribution of Numbers of Heterozygous Loci	
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#### Description

Build Expected Distribution of Numbers of Heterozygous Loci

#### Usage

```
DistHetero(H)
```

#### Arguments

Н

a vector of average heterozygosity of each locus

#### Details

This function build the expected distribution of numbers of heterozygous loci for known heterozygosity of each loci.

## Value

a dataframe of expected density on each possible total number of heterozygous loci.

#### References

Chakraborty, R. (1981, ISSN:0016-6731)

## Examples

DistHetero(runif(10))

Dist\_SimuChisq Build a simulated distribution for Chi-Square

## Description

Build a simulated distribution for Chi-Square

## Usage

Dist\_SimuChisq(s,prob,b)

## Arguments

S	a matrix of frequencies for each simulated sample. Each row for each sample.
prob	a vector of expected probability for each simulated sample.
b	the times of bootstrapping.

## Details

This function build the distribution of Chi square statistics for simulated samples

## Value

a vector of Chi-square statistics, length is the times of sampling.

```
require(mixIndependR)
h<-runif(10)
s<-Simulate_DistK(h,500,100)
Exp <- DistHetero(h)
Dist_SimuChisq(s,Exp$Density,10)
```

ExpProAlleleShare Calculate the Expected Probability of 0,1 and 2 Shared Alleles###

## Description

Calculate the Expected Probability of 0,1 and 2 Shared Alleles###

## Usage

ExpProAlleleShare(p)

#### Arguments

р

a matrix/double of frequency of alleles; Outcome of "AlleleFreq". Each column denotes each locus. Different alleles is ordered in different rows such as 11,11.3,12,12.2,13... and so on

#### Details

This function Calculates the Expected Probability of 0,1 and 2 Shared Alleles for a set of loci. Usually followed by write.csv(as.data.frame(y),file = "/\*.csv") to export the result of a n x3 matrix.

#### Value

a matrix/double of expected probabilities of 0,1 and 2 shared alleles for each locus. Each row denotes each locus. The first column denotes the probability of 0 shared alleles, the second denotes 1 shared allele, the third denotes 2 shared alleles.

#### References

Weir, B. S. (2004, ISSN:0022-1198)

```
a0<-matrix(runif(20),nrow=5)
a1<-colSums(a0)
a<-data.frame(STR1=a0[,1]/a1[1],STR2=a0[,2]/a1[2],STR3=a0[,3]/a1[3],STR4=a0[,4]/a1[4])
ExpProAlleleShare(a)
```

FreqAlleleShare

#### Description

Build Observed Distribution of No. of Shared Alleles

## Usage

```
FreqAlleleShare(AS)
```

#### Arguments

AS

a matrix of number of shared alleles, made up with 0, 1 and 2, outcome of function "AlleleShare\_Table". Rows for individuals, and columns for markers.

## Details

This function build the observed distributions from observed Allele Share table, made up of 0,1 and 2.

#### Value

a dataframe of frequencies of each number of shared alleles(from 0 to 2\*N0. of loci)

#### Examples

```
AS<-matrix(sample(c(0:2),20,replace=TRUE,prob=c(0.3,0.3,0.4)),nrow=5)
FreqAlleleShare(AS)</pre>
```

FreqHetero

Build Observed Distribution of No. of Heterozygous loci

#### Description

Build Observed Distribution of No. of Heterozygous loci

## Usage

```
FreqHetero(h)
```

#### Arguments

h

a dataframe of heterozygosity, made up with 0 and 1, outcome of function "Heterozygous" Rows for individuals, and columns for markers.

#### GenotypeFreq

## Details

This function build the observed distributions from observed heterozygosity table, made up of 0,1.

#### Value

a dataframe of frequencies of each number of heterozygous loci(from 0 to No. of loci)

## Examples

```
h<-matrix(rbinom(20,1,0.5),nrow=5)
FreqHetero(h)</pre>
```

GenotypeFreq Calculate Genotype Frequency###

## Description

Calculate Genotype Frequency###

## Usage

GenotypeFreq(x,p,expect=TRUE)

#### Arguments

x	a dataset of alleles. Each row denotes each sample. One allele in one cell.In the (2r-1)th column, there is the other allele on the same locus from that in the 2r-th column; noted: no column for ID, make row.names=1 when importing.
р	a matrix of allele frequencies. Each row denotes each allele; each column de- notes each marker. The order of markers follows x.
expect	a logic variable. If expect is true, the function will calculate the expected geno- type probabilities. If false, calculate the observed genotype frequencies.

#### Details

This function calculates the observed or expected genotype frequency from dataset and allele frequency.######

## Value

y a matrix of genotype frequencies. Each row denotes each genotype; each column denotes each loci. The order of markers follows x; the genotypes are ordered by: from 1:1-th column, the genotypes are homozygous in order as : p1p1, p2p2,p3p3,...,plp1;from ll-th to u-th column, the genotypes are heterozygous in order as:choose(1,2) like: p1p2,p1p3,...,p1p1,p2p3,p2p4,...p2p1,...p(l-1)pl

## References

Chakraborty, R., Srinivasan, M. R., & Daiger, S. P. (1993, ISSN:0002-9297).

## Examples

Heterozygous Test heterozygosity at each locus

## Description

Test heterozygosity at each locus

#### Usage

```
Heterozygous(x)
```

#### Arguments

Х

a dataset of alleles. Each row denotes each individual.One allele in one cell.In the (2r-1)th column, there is the same locus with the 2r-th column; noted: no column for ID, make row.names=1 when importing.

## Details

This function test the heterozygosity of each individuals at each locus. Output a table and Usually followed by write.csv(as.data.frame(y),file = "/\*.csv") to export the results.

#### Value

a dataframe of heterozygosity.0 is homozygous;1 is heterozygous. Each row denotes each individual; Each column denotes each locus.

#### Examples

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HWE.Chisq

## Description

Test the Hardy Weinberg Equilibrium with Chi-square test####

## Usage

HWE.Chisq(x,x0,rescale.p=FALSE,simulate.p.value=FALSE,B)

#### Arguments

x	a matrix of observed genotype frequencies. Each row denotes each genotype; each column denotes each loci. The order of markers follows x; the genotypes are ordered by: from 1:1-th column, the genotypes are homozygous in order as : p1p1, p2p2,p3p3,,plp1;from ll-th to u-th column, the genotypes are heterozy- gous in order as:choose(1,2) like: p1p2,p1p3,,p1p1,p2p3,p2p4,,p2p1,,p(1-1)p1
x0	a matrix of expected Probabilities;each row denotes each genotype; each col- umn denotes each loci. The order of markers follows x; the genotypes are or- dered by: from 1:1-th column, the genotypes are homozygous in order as : p1p1, p2p2,p3p3,,plp1;from ll-th to u-th column, the genotypes are heterozygous in order as:choose(1,2) like: p1p2,p1p3,,p1p1,p2p3,p2p4,p2p1,,p(l-1)pl
rescale.p	a logical scalar; if TRUE then p is rescaled (if necessary) to sum to 1. If rescale.p is FALSE, and p does not sum to 1, an error is given.
simulate.p.value	
	a logical indicating whether to compute p-values by Monte Carlo simulation.
В	an integer specifying the number of replicates used in the Monte Carlo test.

## Details

This function check the Hardy Weinberg Equilibrium from observed and expected distribution with Chi-square test######

## Value

y a list of result of chi-square test, \$chi \$pvalue; chi and pvalue are vectors of chi square statistics/ p values. Orders follows x.

```
g <- GenotypeFreq(x,p,expect=FALSE)
g0 <- GenotypeFreq(x,p,expect=TRUE)
HWE.Chisq(g,g0,rescale.p=FALSE,simulate.p.value=TRUE,2000)</pre>
```

HWE.Fisher

Test the Hardy Weinberg Equilibrium with Fisher's exact test###

## Description

Test the Hardy Weinberg Equilibrium with Fisher's exact test###

## Usage

HWE.Fisher(p,H,y)

## Arguments

р	a matrix of allele frequency;each row denotes allele; each column denotes each loci;
Н	a vector of number of Heterozygotes on each loci; length is number of loci.
у	a matrix of observed genotype Densities(Not count). Each row denotes each genotype; each column denotes each loci. The order of markers follows x; the genotypes are ordered by: from 1:1-th column, the genotypes are homozygous in order as : p1p1, p2p2,p3p3,,plp1;from ll-th to u-th column, the genotypes are heterozygous in order as:choose(1,2) like: p1p2,p1p3,,p1p1,p2p3,p2p4,p2p1,p(l-1)pl

## Details

This function check the Hardy Weinberg Equilibrium with Fisher's exact Test.#####

## Value

a vector of p-values of Fisher's test; ordered by the order of loci in p or x

## References

Weir, B. S. (1996, ISBN:9780878939022)

## Examples

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```
H <- RxpHetero(h,p,HWE = FALSE)
HWE.Fisher(p,H,G/colSums(G))</pre>
```

Prop\_Pvalue

#### Loop and Find distribution of P values

## Description

Loop and Find distribution of P values

#### Usage

Prop\_Pvalue(x,N,B,t,m=NA,Part=FALSE,alpha=0.05)

#### Arguments

x	a dataset of alleles. Each row denotes each individual.One allele in one cell.In the (2r-1)th column, there is the same locus with the 2r-th column; noted: no column for ID, make row.names=1 when importing.
Ν	times of running this loop, also the number of p-values in the bundle.
В	times of bootstrapping in Chi Squares Test.
t	times of simulation in "Simulate_DistK" and "Simulate_DistX".
m	when Part is TRUE, sub-samples are chosen from x. m is the number of loci in the subsample
Part	a logical variable. If TRUE, this function will calculate p-values for sub-samples with a given sample size m.
alpha	1- confidence level; if the confidence level is $95\%$ , alpha =0.05

## Details

This function can generate a bundle of p-values for one sample or for fix-sized sub-samples. The bundle of cumulative probabilities and the proportion of p-values(1- cumulative probability, please refer to the description of Dist\_SimuChisq.R) smaller than Alpha are exported.

## Value

a list of bundle of cumulative probabilities for number of heterozygous loci and bundle for number of shared alleles; and the proportions of p values smaller than alpha.

#### Examples

RealProAlleleShare Calculate the Real Probability of 0,1 and 2 Shared Alleles###

#### Description

Calculate the Real Probability of 0,1 and 2 Shared Alleles###

#### Usage

```
RealProAlleleShare(AS)
```

#### Arguments

AS

a matrix/double of no. of Shared alleles, made up with 0,1 and 2; Outcome of "AlleleShare\_Table". Each column denotes each locus. Each row denotes each individual.

## Details

This function Calculates the density of 0,1 and 2 Shared Alleles for a set of loci. Usually followed by write.csv(as.data.frame(y),file = "/\*.csv") to export the result of a n x3 matrix.

## Value

a matrix/double of real density of 0,1 and 2 shared alleles for each locus. Each row denotes each locus. The first column denotes the probability of 0 shared alleles, the second denotes 1 shared allele, the third denotes 2 shared alleles.

#### Examples

```
AS<-matrix(sample(c(0:2),20,replace=TRUE,prob=c(0.3,0.3,0.4)),nrow=5)
RealProAlleleShare(AS)</pre>
```

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RxpHetero

## Description

Calculate Real or Expected Average Heterozygosity at each locus

## Usage

RxpHetero(h,p,HWE)

#### Arguments

h	a dataset of heterozygosity, made up with 0 and 1. Output of function "Heterozygous". Each row denotes each individual. Each row denotes each locus.
р	a dataset of allele frequency, Output of function "AlleleFreq". Each row denotes each allele, and each column denotes each locus.
HWE	a logic variable. When TRUE, this function will calculate the expected heterozygosity under Hardy-Weinberg Equilibrium: $H= 1$ -sum(q_i^2); q_i is the allele frequency; If FALSE, this function calculate the average heterozygosity from real heterozygosity table.

#### Details

This function calculate average heterozygosity at each locus.Output a vector of number of loci.

## Value

a vector of average heterozygosity on each loci.

#### References

Chakraborty, R., & Jin, L. (1992, ISSN:1432-1203) <doi:10.1007/BF00197257>

Simulate\_DistK

#### Description

Generate a Bundle of Simulated distributions for No. of heterozygous loci with known heterozygosites

## Usage

Simulate\_DistK(H,m,t)

#### Arguments

#### Details

This function generates multinomial distribution for loci known the heterozygosity and build the simulated distribution for no. of heterozygous loci.

#### Value

a matrix of frequencies of No. of Heterozygous Loci. Each row denotes each simulated sample; Each column denotes each No. of Heterozygous loci, from 0 to length of H.

#### Examples

Simulate\_DistK(runif(10),500,100)

Simulate\_DistX Build a simulated distribution for No. of Shared Alleles

## Description

Build a simulated distribution for No. of Shared Alleles

#### Usage

Simulate\_DistX(e,m,t)

#### Arguments

е	a matrix of Probability of Sharing 2,1 or 0 alleles at each loci. Each row denotes each locus. Three columns denote sharing 0,1 or 2 alleles.
m	the sample size you want, usually similar to the real sample size.
t	the number of samples you want to build/ the times to generate a sample

## Details

This function generates multinomial distribution for loci known the Allele Frequency and Expected Probability of Shared 2,1 or 0 alleles

## Value

y a matrix of frequencies of No. of shared alleles. Each row denotes each simulated sample; Each column denotes each No. of shared alleles, from 0 to 2e length of e.

```
e0<-data.frame("P0"=runif(5,min = 0,max = 0.5),"P1"=runif(5,0,0.5))
e<-data.frame(e0,"P2"=1-rowSums(e0))
Simulate_DistX(e,500,10)</pre>
```

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