Package 'segtest'

October 21, 2024

Title Tests for Segregation Distortion in Polyploids

Version 1.0.1

```
Description Provides a suite of tests for
      segregation distortion in F1 polyploid populations (for
      now, just tetraploids). This is under different assumptions of
      meiosis. Details of these methods are described in
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```

2 Contents

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Contents

Index

chisq_g4	 	 	 	 	 	 			 	3
chisq_gl4	 	 	 	 	 	 			 	4
em_li	 	 	 	 	 	 			 	5
gcount_to_gvec	 	 	 	 	 	 			 	6
gvec_to_gcount	 	 	 	 	 	 			 	6
is_valid_2	 	 	 	 	 	 			 	7
iter.array	 	 	 	 	 	 			 	8
like_gknown_2	 	 	 	 	 	 			 	9
like_gknown_3	 	 	 	 	 	 			 	10
like_glpknown_2	 	 	 	 	 	 			 	11
like_glpknown_3	 	 	 	 	 	 			 	12
llike_li	 	 	 	 	 	 			 	13
lrt_men_g4	 	 	 	 	 	 			 	14
lrt_men_gl4	 	 	 	 	 	 			 	16
multidog_to_g	 	 	 	 	 	 			 	18
multi_lrt	 	 	 	 	 	 			 	19
nextElem.arrayiter	 	 	 	 	 	 			 	22
offspring_geno	 	 	 	 	 	 			 	23
offspring_gf_2	 	 	 	 	 	 			 	23
offspring_gf_3	 	 	 	 	 	 			 	24
otest_g	 	 	 	 	 	 			 	25
polymapr_test	 	 	 	 	 	 			 	27
po_gl	 	 	 	 	 	 			 	28
pvec_tet_2	 	 	 	 	 	 			 	29
pvec_tet_3	 	 	 	 	 	 			 	30
simf1g	 	 	 	 	 	 			 	30
simf1gl	 	 	 	 	 	 			 	31
simgl	 	 	 	 	 	 			 	32
three_to_two	 	 	 	 	 	 			 	33
 ufit										33

35

chisq_g4

chisq_g4

Chi Square test when genotypes are known

Description

This chi-squared test is run under the assumption of no double reduction and no preferential pairing.

Usage

```
chisq_g4(x, g1, g2)
```

Arguments

Χ	Vector of observed genotype coun	ts
---	----------------------------------	----

g1 Parent 1's genotype

g2 Parent 2's genotype

Value

A list containing the chi-squared statistic, degrees of freedom, and p-value.

Author(s)

Mira Thakkar and David Gerard

```
x <- c(1, 2, 4, 3, 0)
g1 <- 2
g2 <- 2
chisq_g4(x, g1, g2)

x <- c(10, 25, 10, 0, 0)
g1 <- 1
g2 <- 1
chisq_g4(x, g1, g2)</pre>
```

chisq_gl4

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Chi-Sq for GL

Description

Calculates the MLE genotype and runs a chi-squared test assuming no double reduction and no preferential pairing.

Usage

```
chisq_gl4(gl, g1, g2)
```

Arguments

gl	A matrix of offspring genotype log-likelihoods. The rows index the individuals and the columns index the possible genotypes. So gl[i, k] is the offspring genotype log-likelihood for individual i and genotype k-1.
g1	The first parent's genotype.
g2	The second parent's genotype.

Value

A list containing the chi-squared statistic, degrees of freedom, and p-value.

Author(s)

Mira Thakkar and David Gerard

```
## null sim
set.seed(1)
g1 <- 2
g2 <- 2
g1 <- simf1gl(n = 25, g1 = g1, g2 = g2, alpha = 0, xi2 = 1/3)
chisq_g14(g1 = g1, g1 = g1, g2 = g2)</pre>
```

em_li 5

 em_li

EM algorithm from Li (2011)

Description

EM algorithm to estimate prior genotype probabilities from genotype likelihoods.

Usage

```
em_li(B, itermax = 100L, eps = 1e-05)
```

Arguments

B Matrix of genotype log-likelihoods. The rows index the individuals and the

columns index the genotypes.

itermax The maximum number of iterations.

eps The stopping criteria.

Value

A vector of log prior probabilities for each genotype.

Author(s)

David Gerard

References

• Li, H. (2011). A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. *Bioinformatics*, 27(21), 2987-2993. doi:10.1093/bioinformatics/btr509

```
# Simulate some data
set.seed(1)
gl <- simgl(nvec = c(3, 2, 4, 1, 2))
# Run em
lprob <- em_li(B = gl)
# Exponentiate to get probabilities
prob <- exp(c(lprob))
prob</pre>
```

6 gvec_to_gcount

gcount_to_gvec

Converts genotype counts to genotype vectors.

Description

Converts genotype counts to genotype vectors.

Usage

```
gcount_to_gvec(gcount)
```

Arguments

gcount

The vector of genotype counts.

Value

A vector of length sum(gcount), containing gcount[1] copies of 0, gcount[2] copies of 1, gcount[3] copies of 2, etc.

Author(s)

David Gerard

See Also

```
gvec_to_gcount()
```

Examples

```
gcount <- c(1, 2, 3, 0, 5)
gcount_to_gvec(gcount = gcount)</pre>
```

gvec_to_gcount

Inverse function of gcount_to_gvec().

Description

Inverse function of gcount_to_gvec().

Usage

```
gvec_to_gcount(gvec, ploidy = 4)
```

is_valid_2

Arguments

gvec The vector of genotypes. gvec[i] is the genotype for individual i.

ploidy The ploidy of the species.

Value

A vector of counts. Element k is the number of individuals with genotype k-1.

Author(s)

David Gerard

See Also

```
gcount_to_gvec()
```

Examples

```
gvec <- c(1, 2, 3, 2, 3, 1, 4, 0, 1, 0, 0, 1, 0, 0)
gvec_to_gcount(gvec = gvec)</pre>
```

is_valid_2

Tests if the two parameter model is valid

Description

There is a dependence on the bounds of two-parameter model. This function returns TRUE if those bounds are satisfied and FALSE otherwise.

Usage

```
is_valid_2(dr, pp, drbound = 1/6)
```

Arguments

dr The double reduction rate.

pp The preferential pairing parameter.

drbound The maximum double reduction rate possible.

Value

TRUE if the model is valid, FALSE otherwise.

Author(s)

David Gerard

iter.array

Examples

```
TOL <- 1e-6
is_valid_2(dr = 1/6, pp = 1/3, drbound = 1/6) # Valid
is_valid_2(dr = 1/6, pp = 1/3 - TOL, drbound = 1/6) # Not valid
is_valid_2(dr = 1/6, pp = 1/3 + TOL, drbound = 1/6) # Not valid
```

iter.array

Iterator over array

Description

Iterator over array

Usage

```
## S3 method for class 'array'
iter(obj, by = 1, recycle = FALSE, ...)
```

Arguments

obj An array.

by The dimension to iterate over. recycle Should the iterator reset?

. . . not used

Value

An iterator. This is an S3 arrayiter object, used in conjunction with nextElem to iterate over one index of an array.

Author(s)

David Gerard

See Also

```
nextElem.arrayiter()
```

```
glist <- multidog_to_g(mout = ufit, type = "all_gl", p1 = "indigocrisp", p2 = "sweetcrisp")
g <- iterators::iter(glist$g, by = 3)
head(iterators::nextElem(g))
head(iterators::nextElem(g))
head(iterators::nextElem(g))</pre>
```

like_gknown_2

 $like_gknown_2$

Likelihood under three parameter model when genotypes are known

Description

This is under the two parameter model.

Usage

```
like_gknown_2(x, alpha, xi1, xi2, g1, g2, log_p = TRUE, pen = 0)
```

Arguments

x	A vector of length 5. x[i] is the count of individuals with genotype i-1.
alpha	The double reduction rate.
xi1	The preferential pairing parameter of parent 1.
xi2	The preferential pairing parameter of parent 2.
g1	Parent 1's genotype.
g2	Parent 2's genotype.
log_p	A logical. Should we return the log likelihood or not?
pen	A tiny penalty to help with numerical stability

Value

The (log) likelihood.

Author(s)

David Gerard

```
x <- c(1, 4, 5, 3, 1)
alpha <- 0.01
xi1 <- 0.5
xi2 <- 0.3
g1 <- 1
g2 <- 2
like_gknown_2(
    x = x,
    alpha = alpha,
    xi1 = xi1,
    xi2 = xi2,
    g1 = g1,
    g2 = g2)</pre>
```

10 like_gknown_3

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Likelihood under three parameter model when genotypes are known

Description

This is under the three parameter model.

Usage

```
like_gknown_3(x, tau, beta, gamma1, gamma2, g1, g2, log_p = TRUE, pen = 0)
```

Arguments

x	A vector of length 5. $x[i]$ is the count of individuals with genotype i-1.
tau	The probability of quadrivalent formation.
beta	The probability of double reduction given quadrivalent formation.
gamma1	The probability of AA_aa pairing for parent 1.
gamma2	The probability of AA_aa pairing for parent 2.
g1	Parent 1's genotype.
g2	Parent 2's genotype.
log_p	A logical. Should we return the log likelihood or not?
pen	A tiny penalty to help with numerical stability

Value

The (log) likelihood.

Author(s)

David Gerard

```
x <- c(1, 4, 5, 3, 1)
tau <- 0.5
beta <- 0.1
gamma1 <- 0.5
gamma2 <- 0.3
g1 <- 1
g2 <- 2
like_gknown_3(
    x = x,
    tau = tau,
    beta = beta,
    gamma1 = gamma1,
    gamma2 = gamma2,</pre>
```

like_glpknown_2

```
g1 = g1,
g2 = g2)
```

like_glpknown_2

Likelihood under three parameter model when using offspring genotypes likelihoods but parent genotypes are known.

Description

This is under the two parameter model.

Usage

```
like_glpknown_2(gl, alpha, xi1, xi2, g1, g2, log_p = TRUE)
```

Arguments

gl	The matrix of genotype likelihoods of the offspring. Rows index The individuals, columns index the genotypes.
alpha	The double reduction rate.
xi1	The preferential pairing parameter of parent 1.
xi2	The preferential pairing parameter of parent 2.
g1	Parent 1's genotype.
g2	Parent 2's genotype.
log_p	A logical. Should we return the log likelihood or not?

Value

The (log) likelihood of the two parameter model when using genotype likelihoods.

Author(s)

David Gerard

```
g1 <- 1
g2 <- 0
g1 <- simf1g1(
n = 25,
g1 = g1,
g2 = g2,
rd = 10,
alpha = 0,
xi1 = 1/3,
xi2 = 1/3)
```

12 like_glpknown_3

```
like_glpknown_2(
  gl = gl,
  alpha = 0.01,
  xi1 = 0.5,
  xi2 = 0.3,
  g1 = g1,
  g2 = g2,
  log_p = TRUE)
```

like_glpknown_3

Likelihood under three parameter model when using offspring genotypes likelihoods but parent genotypes are known.

Description

This is under the three parameter model.

Usage

```
like_glpknown_3(gl, tau, beta, gamma1, gamma2, g1, g2, log_p = TRUE)
```

Arguments

gl	The matrix of genotype likelihoods of the offspring. Rows index The individuals, columns index the genotypes.
tau	The probability of quadrivalent formation.
beta	The probability of double reduction given quadrivalent formation.
gamma1	The probability of AA_aa pairing for parent 1.
gamma2	The probability of AA_aa pairing for parent 2.
g1	Parent 1's genotype.
g2	Parent 2's genotype.
log_p	A logical. Should we return the log likelihood or not?

Value

The (log) likelihood of the three parameter model when using genotype likelihoods.

Author(s)

David Gerard

llike_li

Examples

```
g1 <- 1
g2 <- 0
gl <- simf1gl(</pre>
  n = 25,
  g1 = g1,
  g2 = g2,
  rd = 10,
  alpha = 0,
  xi1 = 1/3,
  xi2 = 1/3)
like_glpknown_3(
  gl = gl,
  tau = 1/2,
  beta = 1/12,
  gamma1 = 1/3,
  gamma2 = 1/3,
  g1 = g1,
  g2 = g2,
  log_p = TRUE)
```

llike_li

Objective function for em_li()

Description

Objective function for em_li()

Usage

```
llike_li(B, lpivec)
```

Arguments

B The log-likelihood matrix. Rows are individuals columns are genotypes.

lpivec The log prior vector.

Value

The log-likelihood of a vector of genotype frequencies when using genotype likelihoods. This is from Li (2011).

Author(s)

David Gerard

14 lrt_men_g4

References

• Li, H. (2011). A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. *Bioinformatics*, 27(21), 2987-2993. doi:10.1093/bioinformatics/btr509

Examples

```
# Simulate some data
set.seed(1)
gl <- simgl(nvec = c(3, 2, 4, 1, 2))
# Log-likelihood at given log-priors
prob <- c(0.1, 0.2, 0.4, 0.2, 0.1)
lprob <- log(prob)
llike_li(B = gl, lpivec = lprob)</pre>
```

1rt_men_g4

Likelihood ratio test for segregation distortion with known genotypes

Description

This will run a likelihood ratio test using the genotypes of an F1 population of tetraploids for the null of Mendelian segregation (accounting for double reduction and preferential pairing) against the alternative of segregation distortion. This is when the genotypes are assumed known.

Usage

```
lrt_men_g4(
    x,
    g1,
    g2,
    drbound = 1/6,
    pp = TRUE,
    dr = TRUE,
    alpha = 0,
    xi1 = 1/3,
    xi2 = 1/3
)
```

Arguments

drbound

A vector of genotype counts. x[i] is the number of offspring with genotype i-1.
 g1 The genotype of parent 1.

g2 The genotype of parent 2.

The maximum rate of double reduction. A default of 1/6 is provided, which is the rate under the complete equational segregation model of meiosis.

Irt_men_g4 15

pp	A logical. Should we account for preferential pairing (TRUE) or not (FALSE)?
dr	A logical. Should we account for double reduction (TRUE) or not (FALSE)?
alpha	If dr = FALSE, this is the known rate of double reduction.
xi1	If pp = FALSE, this is the known preferential pairing parameter of parent 1.
xi2	If pp = FALSE, this is the known preferential pairing parameter of parent 2.

Value

A list with the following elements

```
statistic The log-likelihood ratio test statistic.
```

df The degrees of freedom.

p_value The p-value.

alpha The estimated double reduction rate.

xi1 The estimated preferential pairing parameter of parent 1.

xi2 The estimated preferential pairing parameter of parent 2.

Impossible genotypes

Some offspring genotype combinations are impossible given the parental genotypes. If these impossible genotypes combinations show up, we return a p-value of 0, a log-likelihood ratio statistic of Infinity, and missing values for all other return items. The impossible genotypes are:

```
g1 = 0 \&\& g2 = 0 Only offspring genotypes of 0 are possible.
```

g1 = 4 && g2 = 4 Only offspring genotypes of 4 are possible.

 $g1 = 0 \&\& g2 = 4 \mid \mid g1 == 4 \&\& g2 == 0$ Only offspring genotypes of 2 are possible.

g1 = 0 && g2 %in% c(1, 2, 3) || g1 = %in% c(1, 2, 3) && g2 == 0 Only offspring genotypes of 0, 1, and 2 are possible.

g1 = 4 && g2 %in% c(1, 2, 3) || g1 = %in% c(1, 2, 3) && g2 == 4 Only offspring genotypes of 2, 3, and 4 are possible.

Unidentified parameters

When g1 = 2 or g2 = 2 (or both), the model is not identified and those estimates (alpha, xi1, and xi2) are meaningless. Do NOT interpret them.

The estimate of alpha (double reduction rate) IS identified as long as at least one parent is simplex, and no parent is duplex. However, the estimates of the double reduction rate have extremely high variance.

Author(s)

David Gerard

lrt_men_gl4

Examples

```
set.seed(100)
gf <- offspring_gf_2(alpha = 1/12, xi1 = 0.2, xi2 = 0.6, p1 = 1, p2 = 0)
x <- offspring_geno(gf = gf, n = 100)
lrt_men_g4(x = x, g1 = 1, g2 = 0)</pre>
```

1rt_men_gl4

Likelihood ratio test using genotype likelihoods.

Description

This will run a likelihood ratio test using the genotypes of an F1 population of tetraploids for the null of Mendelian segregation (accounting for double reduction and preferential pairing) against the alternative of segregation distortion. This is when genotype uncertainty is accounted for through genotype likelihoods.

Usage

```
lrt_men_gl4(
   gl,
   g1 = NULL,
   g2 = NULL,
   drbound = 1/6,
   pp = TRUE,
   dr = TRUE,
   alpha = 0,
   xi1 = 1/3,
   xi2 = 1/3
)
```

Arguments

gl	The genotype log-likelihoods. The rows index the individuals and the columns index the genotypes.
g1	Either parent 1's genotype, or parent 1's genotype log-likelihoods.
g2	Either parent 2's genotype, or parent 2's genotype log-likelihoods.
drbound	The upper bound on the double reduction rate.
рр	Is (partial) preferential pairing possible (TRUE) or not (FALSE)?
dr	Is double reduction possible (TRUE) or not (FALSE)?
alpha	If dr = FALSE, this is the known rate of double reduction.
xi1	If pp = FALSE, this is the known preferential pairing parameter of parent 1.
xi2	If pp = FALSE, this is the known preferential pairing parameter of parent 2.

lrt_men_gl4

Value

```
A list with the following elements

statistic The log-likelihood ratio test statistic.

df The degrees of freedom.

p_value The p-value.

alpha The estimated double reduction rate.

xi1 The estimated preferential pairing parameter of parent 1.
```

xi2 The estimated preferential pairing parameter of parent 2.

Unidentified parameters

When g1 = 2 or g2 = 2 (or both), the model is not identified and those estimates (alpha, xi1, and xi2) are meaningless. Do NOT interpret them.

The estimate of alpha (double reduction rate) IS identified as long as at least one parent is simplex, and no parent is duplex. However, the estimates of the double reduction rate have extremely high variance.

Author(s)

David Gerard

```
## null simulation
set.seed(1)
g1 <- 2
g2 <- 2
g1 <- simf1g1(n = 25, g1 = g1, g2 = g2, alpha = 1/12, xi2 = 1/2)
## LRT when parent genotypes are known.
lrt_men_g14(g1 = g1, g1 = g1, g2 = g2)
## LRT when parent genotypes are not known
lrt_men_g14(g1 = g1)
## Alternative simulation
g1 <- simg1(nvec = rep(5, 5))
lrt_men_g14(g1 = g1, g1 = g1, g2 = g2)</pre>
```

18 multidog_to_g

multidog_to_g

Converts multidog output to a format usable for multi_lrt()

Description

Converts multidog output to a format usable for multi_lrt()

Usage

```
multidog_to_g(
  mout,
  type = c("off_gl", "all_gl", "all_g", "off_g"),
  p1 = NULL,
  p2 = NULL,
  ploidy = 4
)
```

Arguments

mout	The output of multidog().
type	"off_gl" Genotype likelihoods of offspring but not parents. This is the typical choice if you used the "fl" or "flpp" options when genotyping.
	"all_gl" Genotype likelihoods of offspring and parents. This is only done if you did <i>not</i> use the "f1" or "f1pp" options when genotyping. If this is the case, then you need to specify which individuals are the parents.
	"off_g" Genotypes, assuming that they are known. You used the "f1" or "f1pp" option when genotyping.
	"all_g" Genotypes, assuming that they are known. You did <i>not</i> use the "f1" or "f1pp" option when genotyping. If this is the case, then you need to specify which individuals are the parents.
p1	The first parent name if using type = "all_gl" or type = "all_g".
p2	The second parent name if using type = "all_gl" or type = "all_g".
ploidy	The ploidy. Note that most methods in this package (including those in multi_lrt()) assume that the ploidy is 4. But we allow for arbitrary ploidy in this function since it might be useful in the future.

Value

A list with the following elements

g Either a matrix of counts, where the columns index the genotype and the rows index the loci (type = "all_g" or type = "off_g"). Or an array of genotype (natural) log-likelihoods where the rows index the loci, the columns index the individuals, and the slices index the genotypes (type = "all_gl" or type = "off_gl").

multi_lrt 19

p1 Either a vector of known parental genotypes (type = "off_g1", type = "all_g" or type = "off_g"). Or a matrix of genotype (natural) log-likelihoods where the rows index the loci and the columns index the genotypes (type = "all_g1").

p2 Either a vector of known parental genotypes (type = "off_g1", type = "all_g" or type = "off_g"). Or a matrix of genotype (natural) log-likelihoods where the rows index the loci and the columns index the genotypes (type = "all_g1").

Author(s)

David Gerard

Examples

```
multidog_to_g(mout = ufit, type = "all_g", p1 = "indigocrisp", p2 = "sweetcrisp")
multidog_to_g(mout = ufit, type = "all_gl", p1 = "indigocrisp", p2 = "sweetcrisp")
multidog_to_g(mout = ufit2, type = "off_g")
multidog_to_g(mout = ufit2, type = "off_gl")
multidog_to_g(mout = ufit3, type = "off_g")
multidog_to_g(mout = ufit3, type = "off_gl")
```

multi_lrt

Parallelized likelihood ratio test for segregation distortion.

Description

Uses the future package to implement parallelization support for the likelihood ratio tests for segregation distortion. Right now, this is only supported for tetraploids (allo, auto, or segmental). This function is only somewhat tested (the single-locus LRT functions in the "See Also" section are very well tested). So please send any bugs you notice to https://github.com/dcgerard/segtest/issues.

Usage

```
multi_lrt(
    g,
    p1,
    p2,
    drbound = 1/6,
    pp = TRUE,
    dr = TRUE,
    alpha = 0,
    xi1 = 1/3,
    xi2 = 1/3,
    nullprop = FALSE
)
```

20 multi_lrt

Arguments

g One of two inputs

• A matrix of genotype counts. The rows index the loci and the columns index the genotypes.

 An array of genotype log-likelihoods. The rows index the loci, the columns index the individuals, and the slices index the genotypes. Log-likelihoods are base e (natural log).

p1 One of three inputs

- A vector of parent 1's genotypes.
- A matrix of parent 1's genotype log-likelihoods. The rows index the loci and the columns index the genotypes. Logs are in base e (natural log).
- NULL (only supported when using genotype likelihoods for the offspring)

p2 One of three inputs

- A vector of parent 1's genotypes.
- A matrix of parent 1's genotype log-likelihoods. The rows index the loci and the columns index the genotypes. Logs are in base e (natural log).
- NULL (only supported when using genotype likelihoods for the offspring)

drbound The upper bound on the double reduction rate.

pp Is (partial) preferential pairing possible (TRUE) or not (FALSE)?

dr Is double reduction possible (TRUE) or not (FALSE)?

alpha If dr = FALSE, this is the known rate of double reduction.

xi1 If pp = FALSE, this is the known preferential pairing parameter of parent 1. xi2 If pp = FALSE, this is the known preferential pairing parameter of parent 2.

nullprop Should we return the null proportions (TRUE) or not (FALSE)?

Value

A data frame with the following elements:

statistic The likelihood ratio test statistic

p_value The p-value of the likelihood ratio test.

df The degrees of freedom of the test.

alpha The MLE of the double reduction rate. Do not use for real work.

- xi1 The MLE of the first parent's partial preferential pairing parameter. Do not use for real work.
- xi2 The MLE of the second parent's partial preferential pairing parameter. Do not use for real work.
- p1 (Estimate of) the first parent's genotype.
- p2 (Estimate of) the second parent's genotype.
- snp The name of the SNP.

multi_Irt 21

Parallel Computation

The multi_lrt() function supports parallel computing. It does so through the future package.

You first specify the evaluation plan with plan() from the future package. On a local machine, this is typically just future::plan(future::multisession, workers = nc) where nc is the number of workers you want. You can find the maximum number of possible workers with availableCores(). You then run multi_lrt(), then shut down the workers with future::plan(future::sequential).

Author(s)

David Gerard

See Also

- lrt_men_g4() Single locus LRT for segregation distortion when genotypes are known.
- 1rt_men_g14() Single locus LRT for segregation distortion when using genotype likelihoods.

```
## Assuming genotypes are known (typically a bad idea)
glist <- multidog_to_g(mout = ufit, type = "all_g", p1 = "indigocrisp", p2 = "sweetcrisp")</pre>
p1_1 <- glist$p1
p2_1 <- glist$p2
g_1 <- glist$g
multi_lrt(g = g_1, p1 = p1_1, p2 = p2_1)
## Using genotype likelihoods (typically a good idea)
glist <- multidog_to_g(mout = ufit, type = "all_gl", p1 = "indigocrisp", p2 = "sweetcrisp")</pre>
p1_2 \leftarrow glist p1
p2_2 \leftarrow glistp2
g_2 <- glist$g
multi_lrt(g = g_2, p1 = p1_2, p2 = p2_2)
## Offspring genotype likelihoods and parent genotypes known
multi_lrt(g = g_2, p1 = p1_1, p2 = p2_1)
## Offspring genotype likelihoods and no information on parent genotypes
multi_lrt(g = g_2, p1 = NULL, p2 = NULL)
## Parallel computing is supported through the future package
future::plan(future::multisession, workers = 2)
multi_lrt(g = g_2, p1 = p1_2, p2 = p2_2)
future::plan(future::sequential)
```

22 nextElem.arrayiter

nextElem.arrayiter

Next element in an array

Description

This is applied to an arrayiter object to obtain the next sub-array along one of the dimensions.

Usage

```
## S3 method for class 'arrayiter'
nextElem(obj, ...)
```

Arguments

obj An arrayiter object

... not used

Value

The next sub-array.

Author(s)

David Gerard

See Also

```
iter.array()
```

```
glist <- multidog_to_g(mout = ufit, type = "all_gl", p1 = "indigocrisp", p2 = "sweetcrisp")
g <- iterators::iter(glist$g, by = 3)
head(iterators::nextElem(g))
head(iterators::nextElem(g))
head(iterators::nextElem(g))</pre>
```

offspring_geno 23

offspring_geno

Simulates genotypes given genotype frequencies.

Description

Takes as input the offspring genotype frequencies and a sample size and returns simulated genotypes.

Usage

```
offspring_geno(gf, n)
```

Arguments

gf Vector of offspring genotype frequencies

n Sample size

Value

Simulated genotypes

Author(s)

Mira Thakkar

Examples

```
set.seed(1) gf \leftarrow offspring_gf_2(alpha = 1/6, xi1 = 1/3, xi2 = 1/3, p1 = 2, p2 = 3) offspring_geno(gf = gf, n = 10)
```

offspring_gf_2

Calculates offspring genotype frequencies under the two-parameter model.

Description

Calculates offspring genotype frequencies under the two-parameter model.

Usage

```
offspring_gf_2(alpha, xi1, xi2 = xi1, p1, p2)
```

24 offspring_gf_3

Arguments

alpha	The double reduction rate
xi1	The preferential pairing parameter of the first parent.
xi2	The preferential pairing parameter of the second parent.
p1	The first parent's genotype
p2	The second parent's genotype

Value

Offspring genotype frequencies

Author(s)

Mira Thakkar

Examples

```
alpha <- 1/6
xi1 <- 1/3
xi2 <- 1/3
p1 <- 2
p2 <- 3
offspring_gf_2(alpha = alpha, xi1 = xi1, xi2 = xi2, p1 = p1, p2 = p2)</pre>
```

offspring_gf_3 Calculates offspring genotype frequencies under the three-parameter model.

Description

Calculates offspring genotype frequencies under the three-parameter model.

Usage

```
offspring_gf_3(tau, beta, gamma1, gamma2 = gamma1, p1, p2)
```

Arguments

tau	Probability of quadrivalent formation
beta	Probability of double reduction given quadrivalent formation
gamma1	Probability of AA_aa pairing in parent 1
gamma2	Probability of AA_aa pairing in parent 2
p1	The first parent's genotype
p2	The second parent's genotype

otest_g 25

Value

Offspring genotype frequencies

Author(s)

David Gerard

Examples

```
offspring_gf_3(
tau = 1/2,
beta = 1/6,
gamma1 = 1/3,
gamma2 = 1/3,
p1 = 1,
p2 = 2)
```

 $otest_g$

Jointly tests for segregation distortion and number of incompatible genotypes

Description

This is experimental. I haven't tested it out in lots of scenarios yet.

Usage

```
otest_g(
    x,
    g1,
    g2,
    pbad = 0.03,
    drbound = 1/6,
    pp = TRUE,
    dr = TRUE,
    alpha = 0,
    xi1 = 1/3,
    xi2 = 1/3
)
```

Arguments

```
x A vector of genotype counts. x[i] is the number of offspring with genotype i-1.
```

g1 The genotype of parent 1.

g2 The genotype of parent 2.

26 otest_g

pbad	The upper bound on the number of bad genotypes
drbound	The maximum rate of double reduction. A default of 1/6 is provided, which is the rate under the complete equational segregation model of meiosis.
рр	A logical. Should we account for preferential pairing (TRUE) or not (FALSE)?
dr	A logical. Should we account for double reduction (TRUE) or not (FALSE)?
alpha	If dr = FALSE, this is the known rate of double reduction.
xi1	If pp = FALSE, this is the known preferential pairing parameter of parent 1.
xi2	If pp = FALSE, this is the known preferential pairing parameter of parent 2.

Details

Here, we test if the compatible genotypes are consistent with F1 populations and separately test that the number of incompatible genotypes isn't too large (less than 3 percent by default). This is the strategy the polymapR software uses. But we use a Bonferroni correction to combine these tests (minimum of two times the p-values), while they just multiply the p-values together. So our approach accounts for double reduction and preferential pairing, while also controlling the family-wise error rate.

Value

```
A list with the following elements
```

statistic The log-likelihood ratio test statistic.

df The degrees of freedom.

p_value The Bonferroni corrected p-value.

p_lrt The p-value of the LRT.

p_binom The p-value of the one-sided binomial test.

alpha The estimated double reduction rate.

xi1 The estimated preferential pairing parameter of parent 1.

xi2 The estimated preferential pairing parameter of parent 2.

Author(s)

David Gerard

```
# Run a test where genotypes 0, 1, and 2 are possible
x <- c(10, 10, 4, 0, 5)
otest_g(x = x, g1 = 1, g2 = 0)

# polymapR's multiplication and the Bonferroni differ
df <- expand.grid(p1 = seq(0, 1, length.out = 20), p2 = seq(0, 1, length.out = 20))
df$polymapr <- NA
df$bonferroni <- NA
for (i in seq_len(nrow(df))) {
    df$polymapr[[i]] <- df$p1[[i]] * df$p2[[i]]</pre>
```

polymapr_test 27

polymapr_test

Run segregation distortion tests as implemented in the polymapR package.

Description

The polymapR package tests for segregation distortion by iterating through all possible forms of disomic or polysomic inheritance from either parent, tests for concordance of the offspring genotypes using a chi-squared test, and returns the largest p-value. It sometimes chooses a different p-value based on other heuristics. They also sometimes return NA. When type = "segtest", we only look at patterns of the given parent genotypes, choosing the largest p-value. When type = "polymapR", we return what they use via their heuristics.

Usage

```
polymapr_test(x, g1 = NULL, g2 = NULL, type = c("segtest", "polymapR"))
```

Arguments

X	Either a vector of genotype counts, or a matrix of genotype posteriors where the rows index the individuals and the columns index the genotypes.
g1	Parent 1's genotype.
g2	Parent 2's genotype.
type	Either my implementation which approximates that of polymapR ("segtest") or the implementation through polymapR ("polymapR"). Note that polymapR needs to be installed for type = "polymapR".

Value

A list with the following elements:

```
p_value The p-value of the test.bestfit The best fit model, using the same notation as in checkF1().frq_invalid The frequency of invalid genotypes.
```

Author(s)

David Gerard

See Also

```
checkF1().
```

28 po_gl

Examples

```
g1 <- 0
g2 <- 1
x <- c(4, 16, 0, 0, 0)
polymapr_test(x = x, g1 = g1, g2 = g2, type = "segtest")
```

po_gl

Generate genotype likelihoods from offspring genotypes.

Description

Takes as input (i) the parent genotypes, (ii) the offspring genotype frequency, (iii) sequencing error rate, (iv) read depth, (v) bias, and (vi) overdispersion. It returns genotype likelihoods.

Usage

```
po_gl(
   genovec,
   ploidy,
   p1_geno = NULL,
   p2_geno = NULL,
   rd = 10,
   seq = 0.01,
   bias = 1,
   od = 0.01
)
```

Arguments

```
genovec
                  Offspring genotypes. genovec[i] is the dosage for individual i.
ploidy
                  Ploidy
p1_geno
                  Parent 1 genotype
                  Parent 2 genotype
p2_geno
rd
                  Read depth. Lower is more uncertain.
                  Sequencing error rate. Higher means more uncertain.
seq
bias
                  Bias. 1 means no bias.
                  Overdispersion. Typical value is like 0.01. Higher means more uncertain.
od
```

Value

Genotype likelihoods

pvec_tet_2 29

Author(s)

Mira Thakkar

Examples

```
set.seed(1)
po_gl(genovec = c(1, 2, 1, 1, 3), p1_geno = 2, p2_geno = 2, ploidy = 4)
```

 $pvec_tet_2$

Tetraploid gamete frequencies of gametes when one parent's genotype is known

Description

This is under the two parameter model.

Usage

```
pvec_tet_2(alpha, xi, ell)
```

Arguments

alpha	The double reduction rate
xi	The preferential pairing parameter
ell	The parental genotype

Value

The gamete genotype frequencies

Author(s)

Mira Thakkar and David Gerard

```
alpha <- 1/6
xi <- 1/3
pvec_tet_2(alpha = alpha, xi = xi, ell = 0)
pvec_tet_2(alpha = alpha, xi = xi, ell = 1)
pvec_tet_2(alpha = alpha, xi = xi, ell = 2)
pvec_tet_2(alpha = alpha, xi = xi, ell = 3)
pvec_tet_2(alpha = alpha, xi = xi, ell = 4)</pre>
```

30 simf1g

pvec_tet_3	Tetraploid gamete frequencies of gametes when one parent's genotype is known

Description

This is under the three parameter model.

Usage

```
pvec_tet_3(tau, beta, gamma, ell)
```

Arguments

tau	Probability of quadrivalent formation
-----	---------------------------------------

beta Probability of double reduction given quadrivalent formation

gamma Probability of AA/aa pairing given bivalent formation

ell The parent genotype

Value

The gamete genotype frequencies

Author(s)

David Gerard

Examples

```
pvec_tet_3(tau = 0.5, beta = 0.1, gamma = 0.5, ell = 2)
```

simf1g

Simulate genotype counts from F1 individuals

Description

Simulate genotype counts from F1 individuals

Usage

```
simf1g(n, g1, g2, alpha = 0, xi1 = 1/3, xi2 = 1/3)
```

simf1g1 31

Arguments

n	Sample size.
g1	The first parent's genotype.
g2	The second parent's genotype.
alpha	The double reduction rate.
xi1	The first parent's preferential pairing parameter.
xi2	The second parent's preferential pairing parameter.

Value

A vector of counts, where element i is the number of simulated individuals with genotype i-1.

Author(s)

David Gerard

Examples

```
set.seed(1)
simf1g(n = 10, g1 = 1, g2 = 2)
```

simf1gl

Simulate genotype likelihoods of F1 individuals.

Description

Simulate genotype likelihoods of F1 individuals.

Usage

```
simf1g1(n, g1, g2, rd = 10, alpha = 0, xi1 = 1/3, xi2 = 1/3)
```

Arguments

n	Sample size.
g1	The first parent's genotype.
g2	The second parent's genotype.
rd	The read depth.
alpha	The double reduction rate.
xi1	The first parent's preferential pairing parameter.
xi2	The second parent's preferential pairing parameter.

32 simgl

Value

The matrix of offspring genotype log-likelihoods.

Author(s)

David Gerard

Examples

```
set.seed(1)
simf1gl(n = 10, g1 = 1, g2 = 2)
```

simgl

Simulate genotype likelihoods from genotype counts

Description

Provide a vector of genotype counts and this will return a matrix of genotype log-likelihoods.

Usage

```
simgl(nvec, rd = 10, seq = 0.01, bias = 1, od = 0.01)
```

Arguments

nvec	A vector of counts. $nvec[k]$ is the number of folks with a genotype of k-1.
rd	Read depth. Lower is more uncertain.
seq	Sequencing error rate. Higher means more uncertain.
bias	Bias. 1 means no bias.
od	Overdispersion. Typical value is like 0.01. Higher means more uncertain.

Value

A matrix of genotype log-likelihoods. The rows index the individuals and the columns index the genotypes. This is natural log (base e).

Author(s)

David Gerard

```
set.seed(1)
simgl(nvec = c(1, 2, 1, 1, 3))
```

three_to_two 33

three_to_two

Convert from three parameters to two parameters

Description

Convert from three parameters to two parameters

Usage

```
three_to_two(tau, beta, gamma)
```

Arguments

tau Probability of quadrivalent formation

beta Probability of double reduction given quadrivalent formation

gamma Probability of AA/aa pairing given bivalent formation

Value

A vector of length two. The first is the double reduction rate (alpha), and the second is the preferential pairing parameter (xi).

Author(s)

David Gerard

Examples

```
three_to_two(tau = 0.1, beta = 1/6, gamma = 1/4)
```

ufit

Genotype data from Cappai et al. (2020)

Description

A subset of data from Cappai et al. (2020), fit using multidog(). This just contains a random set of 10 loci.

Usage

ufit

ufit2

ufit3

34 ufit

Format

```
An object of type multidog output from multidog().

ufit Uses the model = "norm" option.

ufit2 Uses the model = "f1pp" option.

ufit3 Uses the model = "f1" option.

An object of class multidog of length 2.

An object of class multidog of length 2.
```

Source

doi:10.5281/zenodo.13715703

References

Cappai, F., Amadeu, R. R., Benevenuto, J., Cullen, R., Garcia, A., Grossman, A., Ferrão, L., & Munoz, P. (2020). High-resolution linkage map and QTL analyses of fruit firmness in autotetraploid blueberry. *Frontiers in plant science*, 11, 562171. doi:10.3389/fpls.2020.562171.

Index

* datasets ufit, 33 availableCores, 21	plan, 21 po_gl, 28 polymapr_test, 27 pvec_tet_2, 29 pvec_tet_3, 30
<pre>checkF1, 27 chisq_g4, 3 chisq_g14, 4 em_li, 5, 13</pre>	simf1g, 30 simf1g1, 31 simg1, 32
<pre>gcount_to_gvec, 6, 6 gcount_to_gvec(), 7 gvec_to_gcount, 6 gvec_to_gcount(), 6</pre>	ufit, 33 ufit2 (ufit), 33 ufit3 (ufit), 33
<pre>is_valid_2,7 iter.array, 8 iter.array(), 22</pre>	
like_gknown_2, 9 like_gknown_3, 10 like_glpknown_2, 11 like_glpknown_3, 12 llike_li, 13 lrt_men_g4, 14 lrt_men_g4(), 21 lrt_men_g14, 16 lrt_men_g14(), 21	
multi_lrt, 18, 19 multidog, 18, 33, 34 multidog_to_g, 18	
<pre>nextElem.arrayiter, 22 nextElem.arrayiter(), 8</pre>	
offspring_geno, 23 offspring_gf_2, 23 offspring_gf_3, 24 otest_g, 25	