Package 'polyBreedR'

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Author Jeffrey B. Endelman
Maintainer Jeffrey Endelman <endelman@wisc.edu></endelman@wisc.edu>
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2 check_ploidy

A_mat

Additive relationship matrix from pedigree

Description

Additive relationship matrix from pedigree

Usage

```
A_mat(ped, ploidy, order.ped = TRUE)
```

Arguments

ped Pedigree in three column format: id, mother, father

ploidy 2 or 4

order.ped TRUE/FALSE does the pedigree need to be ordered so that progeny follow par-

ents

Details

This is a wrapper that prepares the pedigree in the format required for R package AGHmatrix by Amadeu et al. (2016) (cite them if you use this function). A random bivalents model for tetraploid meiosis is assumed.

Value

Additive relationship matrix (dim: indiv x indiv)

References

Amadeu et al. (2016) Plant Genome 9, doi:10.3835/plantgenome2016.01.0009

check_ploidy

Check ploidy

Description

Fraction of simplex or triplex markers

Usage

```
check_ploidy(geno, map)
```

Arguments

geno Genotype matrix (markers x indiv)

map Data frame with marker map (Marker, Chrom, Position)

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Details

For every indiv in the genotype matrix, the fraction of markers per chromosome called as simplex or triplex is calculated, which should be low for diploids. A small amount of missing genotype data can be tolerated.

Value

List containing

mat Matrix (indiv x chrom) of resultsplot ggplot2 barplot

check_trio

Check markers for parent-offspring trio

Description

Check markers for parent-offspring trio

Usage

```
check_trio(parentage, geno, ploidy)
```

Arguments

parentage Data frame with three columns: id, mother, father

geno Matrix of allele dosages: markers x indiv

ploidy 2 or 4

Details

Computes the percentage of markers at which the two parents and offspring have incompatible allele dosages (for tetraploids, the random bivalents model is used). For dihaploid offspring of a single tetraploid parent, use ploidy = 4 and "haploid" for the father in parentage, as well as a diploid (0,1,2) genotype for the offspring. A small amount of missing genotype data can be tolerated.

Value

Data frame with the percentage of incompatible markers for each trio

4 gbs

Description

Convert DArTag to VCF

Usage

```
dart2vcf(counts.file, dosage.file, vcf.file, ploidy, first.data.row = 9)
```

Arguments

counts.file DArTag collapsed counts file
dosage.file DArTag dosage file
vcf.file name of VCF output file (uncompressed)

ver. True manie of ver output me (uncompress

ploidy ploidy

first.data.row default is 9 for DArTag format

Details

Two input files expected. counts.file is the two-row collapsed counts file, whereas dosage.file has one row per target, with chrom and position in columns 4 and 5. DArT reports dosage of REF, whereas VCF standard is based on dosage of ALT. The dosage is exported as GT field in VCF.

Duplicate samples are renamed by appending the "Target ID".

gbs	Genotype calls for GBS

Description

Genotype calls for genotype-by-sequencing (GBS) data

Usage

```
gbs(in.file, out.file, ploidy, prior = "norm", n.core = 1)
```

Arguments

prior model for prior (see Details)

n.core number of cores

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Details

VCF input file must contain AD field. Posterior mode and mean genotypes are added as GT and DS fields. GQ is also added based on probability of posterior mode. Binomial calculation uses R/updog package (Gerard et al. 2018). Previous INFO is discarded; adds NS, AVG.DP, AF.GT, AB, OD, SE, MIN.DP, HWE.P.

Value

marker x indiv matrix of read depths

geno_call

Genotype calls

Description

Genotype calls based on a normal mixture model

Usage

```
geno_call(
  data,
  filename,
  model.ploidy = 4L,
  sample.ploidy = 4L,
  min.posterior = 0,
  transform = TRUE
)
```

Arguments

data matrix (markers x id) of input values for the normal mixture model

filename CSV filename with the model parameters

 $\begin{array}{ll} {\sf model.ploidy} & 2 \text{ or 4 (default)} \\ {\sf sample.ploidy} & 2 \text{ or 4 (default)} \end{array}$

min.posterior minimum posterior probability (default 0) for genotype call

transform TRUE (default) or FALSE whether to apply arcsin square root transformation

Details

The first column of the CSV input file should be the SNP ID, followed by columns for the normal distribution means, standard deviations, and mixture probabilities. Genotype calls are based on the maximum a posteriori (MAP) method. If the posterior probability of the MAP genotype is less than min.posterior, then NA is returned for that sample. By default, an arcsin square root transformation is applied to the input values to match the approach used by R package fitPoly. To use a tetraploid mixture model for diploid samples, set sample.ploidy = 2 and model.ploidy = 4.

Value

matrix of allele dosages (0,1,2,...ploidy) with dimensions markers x individuals

6 GvsA

get_	nedi	gree

Generate pedigree

Description

Generate pedigree for a set of individuals

Usage

```
get_pedigree(id, pedfile, delim = ",", na.string = "NA", trim = TRUE)
```

Arguments

id	Vector of names of individuals
pedfile	Name of pedigree file
delim	Delimiter for the pedigree file (default is "," for CSV)
na.string	String used for NA in the pedigree file (default is "NA")
trim	TRUE/FALSE whether to trim pedigree (see Details)

Details

Finds ancestors of individuals in a three-column pedigree file (id,mother,father). The id column can be the identifier for an individual or cross. String matches must be exact or based on the naming convention crossID-progenyID. The returned pedigree is ordered using R package pedigree so that offspring follow parents. When trim is TRUE (default), the pedigree is trimmed to remove ancestors with only one offspring (which are not needed to compute the pedigree relationship matrix).

Value

Data frame with columns id, mother, father

GvsA

Plot G vs. A

Description

Plot marker-based vs. pedigree-based additive relationship coefficients

Usage

```
GvsA(
  parentage,
  G,
  A,
  filename = NULL,
  thresh.G = Inf,
  thresh.A = 0.5,
  Gmax = NULL,
  Amax = NULL
)
```

G_mat 7

Arguments

parentage Data frame of individuals to plot, with 3 columns: id, mother, father

G Genomic relationship matrix
A Pedigree relationship matrix

filename Name of PDF file to save the results (optional for one individual)

thresh.G Threshold above which names are displayed (default Inf)
thresh.A Threshold above which names are displayed (default 0.5)

Gmax Upper limit for y-axis for plotting. If NULL, maximum value in G is used.

Amax Upper limit for x-axis for plotting. If NULL, maximum value in A is used.

Details

Useful for finding and correcting pedigree errors. If the G or A coefficient for an individual exceeds the threshold, its name is displayed in the figure. If parentage contains one individual, by default a ggplot2 variable will be returned, but the result can also be written to file. If multiple individuals are present, a filename is required.

G_mat

Additive genomic relationships

Description

Relationship matrix for additive effects with bi-allelic markers

Usage

```
G_mat(geno, ploidy)
```

Arguments

geno Matrix of allele dosages (markers x indiv)

ploidy Any even integer (2,4,6,...)

Details

Additive effects are based on the traditional orthogonal decomposition of genetic variance in panmictic populations (Fisher 1918; Kempthorne 1957; Endelman et al. 2018). Missing genotype data is replaced with the population mean.

Value

G matrix

References

Fisher (1918) Trans. Roy. Soc. Edin. 52:399-433.

Kempthorne (1957) An Introduction to Genetic Statistics.

Endelman et al. (2018) Genetics 209:77-87.

8 impute

impute

Impute missing marker data

Description

Impute missing marker data

Usage

```
impute(
  in.file,
  out.file,
  ploidy,
  method,
  geno,
  min.DP = 1,
  max.missing,
  params = NULL,
  n.core = 1
)
```

Arguments

```
VCF input file
in.file
out.file
                  VCF output file
ploidy
                  ploidy
method
                  One of the following: "pop", "EM", "RF"
                  One of the following: "GT", "DS"
geno
                  genotypes below this depth are set to missing (default=1)
min.DP
                  remove markers above this threshold, as proportion of population
max.missing
                  list of method-specific parameters
params
n.core
                  multicore processing
```

Details

Assumes input file is sorted by position. Markers with no genetic variance are removed.

method="pop" imputes with the population mean for geno="DS" and population mode for geno="GT".

method="EM" uses parameter "tol" (default is 0.02, see rrBLUP A.mat documentation). Imputed values are truncated if needed to fall in the interval [0,ploidy].

method="RF" uses parameters "ntree" (default 100) for number of trees and "nflank" (default 100) for the number of flanking markers (on each side) to use as predictors. Because RF first uses EM to generate a complete dataset, parameter "tol" is also recognized.

merge_impute 9

merge_impute	Merge two genotype matrices and impute missing data

Description

Merge two genotype matrices and impute missing data by BLUP

Usage

```
merge_impute(geno1, geno2, ploidy)
```

Arguments

geno1	Genotype matrix (coded 0ploidy) with dimensions markers x indiv
geno2	Genotype matrix (coded 0ploidy) with dimensions markers x indiv
ploidy	Either 2 or 4

Details

Designed to impute from low to high density markers. The BLUP method is equivalent to Eq. 4 of Poland et al. (2012), but this function is not iterative. Additional shrinkage toward the mean is applied if needed to keep the imputed values within the range [0,ploidy]. Missing data in the input matrices are imputed with the population mean for each marker. If an individual appears in both input matrices, it is renamed with suffixes ".1" and ".2" and treated as two different individuals. Monomorphic markers are removed.

Value

Imputed genotype matrix (markers x indiv)

References

Poland et al. (2012) Plant Genome 5:103-113.

readXY	Read SNP arra	v intensity data

Description

Read SNP array intensity data

Usage

```
readXY(filename, skip = 9, output = "ratio")
```

Arguments

filename	filename
TITELIAIILE	mename

skip number of lines to skip before the header line with the column names

output One of three options: "ratio", "theta", "AD"

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Details

The first two columns of the tab-delimited input file should be the SNP and Sample ID. Columns labeled "X" and "Y" contain the signal intensities for the two alleles. Use output to specify whether to return the ratio = Y/(X+Y) or theta = atan(Y/X)*2/pi. Option "AD" exports the XY data in the allele depth format for a VCF file ("X,Y"), with the X and Y values multiplied by 100 and rounded to the nearest integer.

Value

matrix with dimensions markers x individuals

update_alias

Update names based on alias

Description

Update names based on data frame with alias and preferred name

Usage

```
update_alias(x, alias, remove.space = TRUE, filename = NULL)
```

Arguments

x Vector of names to update

alias Data frame with two columns: first is the preferred name and second is the alias

remove.space TRUE/FALSE

filename update names in file (variable "id")

Details

Parameter remove. space indicates whether blank spaces should be removed before string matching

Value

Vector with updated names

write_vcf

write_vcf Create VCFv4.3 file	
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Description

Create VCFv4.3 file

Usage

```
write_vcf(filename, fixed, geno, other.meta = NULL)
```

Arguments

filename VCF file name

fixed character matrix with 8 columns: CHROM, POS, ID, REF, ALT, QUAL, FIL-

TER, INFO

geno named list of genotype matrices, see Details

other .meta optional, other metadata (without ##) besides INFO and FORMAT keys

Details

Several standard INFO key are recognized: ##INFO=<ID=NS,Number=1,Type=Integer,Description="Number of samples with data"> ##INFO=<ID=AVG.DP,Number=1,Type=Float,Description="Average Sample Depth"> ##INFO=<ID=DP,Number=1,Type=Integer,Description="Total Depth"> ##INFO=<ID=AB,Number=1,Type=Integer,Description="Sequencing Error (PHRED)"> ##INFO=<ID=AB,Number=1,Type=Integer,Description="Sequencing Error (PHRED)"> ##INFO=<ID=AB,Number=1,Type=Integer,Description="Allele Frequency"> ##INFO=<ID=AB,Number=1,Type=Integer,Description="Sequencing Error (PHRED)"> ##INFO=<ID=AF,GT,Number=1,Type=Integer,Description="Sequency"> ##INFO=<ID=AF,GT,Number=1,Type=Integer,Description="Sequency"> ##INFO=<ID=AF,GT,Number=1,Type=Integer,Description="Sequency"> ##INFO=<ID=AF,GT,Number=1,Type=Integer,Description="Sequency"> ##INFO=<ID=AF,GT,Number=1,Type=Integer,Description="Posterion="Posterion="Allele count in genotypes"> ##INFO=<ID=AN,Number=1,Type=Integer,Description="Total number of alleles"> ##INFO=<ID=AN,Number=1,Type=Integer,Description="Total number of geno. Metadata for geno is generated from the names of geno. Sample names taken from colnames of geno. Metadata for geno is generated from the names of the list: ##FORMAT=<ID=GT,Number=1,Type=String,Description="Allele Depth"> ##FORMAT=<ID=DP,Number=1,Type=Depth"> ##FORMAT=<ID=DP,Number=1,Type=Depth"> ##FORMAT=<ID=DP,Number=1,Type=Integer,Description="Posterior Mean Dosage"> ##FORMAT=<ID=DP,Number=1,Type=Depth"> ##FORMAT=<ID=DP,Number=1,Type=Integer,Description="Genotype Quality"> ##FORMAT=<ID=DP,Number=1,Type=Integer,Description="Genotype Qu

Any additional metadata should be included without the ## prefix.

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