

Package ‘QTLRel’

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Title Tools for mapping of quantitative traits of genetically related individuals and calculating identity coefficients from a pedigree

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Description This software provides tools for quantitative trait mapping in populations such as advanced intercross lines where relatedness among individuals should not be ignored. It can estimate background genetic variance components, impute missing genotypes, simulate genotypes, perform a genome scan for putative quantitative trait loci (QTL), and plot mapping results. It also has functions to calculate identity coefficients from pedigrees, especially suitable for pedigrees that consist of a large number of generations, or estimate identity coefficients from genotypic data in certain circumstances.

Depends R (>= 2.10)

Imports gdata

Suggests lattice, qtl

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LazyData no

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R topics documented:

aicVC	2
blup	4
cic	5
estVC	7
genMatrix	8
genoImpute	9
genoProb	11
genoSim	13
gls	15
hapSim	15
ibs	17
kinship	18
lodci	18
mAIC	20
miscEx	22
misFct	22
nullSim	22
pedRecode	25
plotit	26
qqPlot	27
qtl2rel	29
qtlVar	30
rel2qtl	31
rem	32
scanOne	33
scanTwo	35
Index	36

aicVC	<i>AIC Model Selection</i>
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Description

Select genetic variance components via Akaike's information criterion (AIC).

Usage

```
aicVC(y,x,v=vector("list",6),initpar,k=2,init=6,keep=6,
      direction=c("forward","backward"),nit=25,verbose=FALSE,
      method=c("Nelder-Mead","BFGS","CG","SANN"),control=list(),
      hessian=FALSE)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
v	a list of variance components (AA, DD, HH, AD, MH, EE, ...) where "AA" and "DD" are respectively additive and dominance genetic matrices, "HH", "AD" and "MH" are other genetic matrices that one may be interested in (see "details"), "EE" is the residual matrix that is usually assumed to be an identity matrix, and "..." are other random components of interest. If a genetic component is not considered, it should be set to NULL.
initpar	optional initial parameter values.
k	penalty on a parameter. The selection criterion is the known "AIC" if $k = 2$ and is "BIC" if $k = \log(n)$ where "n" is the sample size.
init	indicator of the initial model pertaining to genetic variance components. For instance, c(1,2,6) indicates the initial model includes "AA", "DD" and "EE". By default, only "EE" is considered.
keep	indicator of which variance components should be forced into the final model. The default is "EE".
direction	the mode of search. Either "forward" or "backward" with default "forward".
nit	number of iterations to call <code>optim</code> for optimization.
verbose	a logical variable. True if ones wants to track the process for monitoring purpose.
method	the optimization method to be used. See <code>optim</code> for details.
control	a list of control parameters to be passed to <code>optim</code> .
hessian	logical. Should a numerically differentiated Hessian matrix be returned?

Details

In genome-wide association studies (GWAS), random effects are usually added to a model to account for polygenic variation. Abney et al (2000) showed that five variance components including the most interesting additive and dominance variance components are potentially induced by polygenes. The above function is intended for selecting variance components that contribute "most" to a quantitative trait.

Function `estVC` is called by the above function to estimate the parameters and maximum likelihood in each model. Refer to `estVC` for more information.

Value

aic	AIC of the final model.
model	gives parameter estimates, log-likelihood, and other information.
lik	log-likelihood of the model selected at each intermediate step.
trace	indicates which variance components were selected at each intermediate step.

References

Abney, M., M. S. McPeck, and C. Ober (2000). Estimation of variance components of quantitative traits in inbred populations. *Am. J. Hum. Genet.* 141, 629-650.

See Also

[estVC](#) for more information.

Examples

```
data(miscEx)

## Not run:
# forward selection
# any variance component will be selected
# if AIC improve by 1e-5 or larger
v=list(AA=gmF8$AA, DD=gmF8$DD, HH=NULL, AD=NULL,
MH=NULL, EE=diag(length(pdatF8$bwt)))
o<- aicVC(y=pdatF8$bwt, x=pdatF8$sex, k=0, v=v, verbose=TRUE)
o

# forward selection
of<- aicVC(y=pdatF8$bwt, x=pdatF8$sex, v=v, k=1/2, init=6,
direction="for", verbose=TRUE)
of

# backward elimination
ob<- aicVC(y=pdatF8$bwt, x=pdatF8$sex, v=v, k=1/2, keep=6,
direction="back", verbose=TRUE)
ob

## End(Not run)
```

blup

Best Linear Unbiased Prediction

Description

Estimate the best linear unbiased prediction (BLUP) for various effects in the model.

Usage

```
blup(object)
```

Arguments

object an object from [estVC](#) or [aicVC](#).

Value

fixed	BLUP for fixed effects.
AA,DD,...	BLUP for random (genetic) variance components "AA", "DD", ...
EE	BLUP for residual effect.

See Also

[estVC](#) and [aicVC](#).

Examples

```
data(miscEx)

## Not run:
# only consider additive genetic variance component
o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))
b<- blup(o)

## End(Not run)
```

cic

Calculate Jacquard condensed identity coefficients

Description

Calculate Jacquard condensed identity coefficients from a pedigree.

Usage

```
cic(ped,ids,inter,df=3,ask=FALSE,verbose=FALSE)
```

Arguments

ped	a pedigree, which is a data frame (id, sire, dam, ...). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", ... If "sex" is included, male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). Note that 0 is reserved for missing values.
ids	IDs of the individuals for which to calculate the Jacquard condensed identity coefficients. If missing, all individuals in the pedigree ped will be considered.
inter	if given, it should indicate intermediate generations.
df	if inter is missing, df is used to derive (optimal) inter. If df = 0, then there will no intermediate generations. If df is large, then all generations will be used as intermediate generations.
ask	if true, users will be asked whether to proceed.
verbose	if true, will print out some messages.

Details

The coefficients will be calculated for individuals with IDs specified by `ids`. All individuals will be considered if `ids` is missing. This is not recommended if the total number of individuals in the pedigree is large. Instead, it is recommended that `ids` is specified for interested individuals only

`df` is a tuning parameter. It should not be 0 (or smaller than 1) if the pedigree is large in depth (many generations) but the number of individuals is not small; otherwise, it can take forever to finish. It should not be `Inf` (or a large number) if the number of individuals in certain intermediate generation is very large.

Value

A matrix `G` with $G_{[,j]}$ being the j -th Jacquard identity coefficients.

Note

You may need the administrative privilege to run this function on systems such as Windows 7. It may require your operating system support "long long" integer type in C++. If you run this function in a windows system, make sure the working directory is under system volume C and you have the write privilege.

It is better to remove the working directory if the program is interrupted by external forces (e.g. killed by users).

Warning: you may need to run this program on a 64-bit machine in case of seeing such a message!

References

Abney, M., M. S. McPeck, and C. Ober (2000). Estimation of variance components of quantitative traits in inbred populations. *Am. J. Hum. Genet.* 141, 629-650.

Examples

```
data(miscEx)

ids<- sample(pedF8$id[300:500],20)

## Not run:
# run 'cic' for the sampled individuals
# top-down
oo<- cic(pedF8, ids=ids, df=Inf, verbose=TRUE)
# bottom-up
o0<- cic(pedF8, ids=ids, df=0, verbose=TRUE)
# hybrid of top-down and bottom-up
o2<- cic(pedF8, ids=ids, ask=TRUE, verbose=TRUE)
# same results
c(sum(abs(oo-o0) >1e-7),sum(abs(o2-o0) >1e-7))

## End(Not run)
```

estVC

*Estimate Variance Component Parameters***Description**

Estimate model parameters for covariates, genetic variance components and residual effect.

Usage

```
estVC(y,x,v=vector("list",6),initpar,nit=25,
      method=c("Nelder-Mead","BFGS","CG","SANN"),
      control=list(),hessian=FALSE)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
v	a list of variance components (AA, DD, HH, AD, MH, EE,...), where "AA" and "DD" are respectively additive and dominance genetic matrices, "HH", "AD" and "MH" are other genetic matrices that one may be interested in (see aicVC), "EE" is the residual matrix that is usually assumed to be an identity matrix, and "..." are other random components of interest. If a genetic component is not considered, it should be set to NULL.
initpar	optional initial parameter values.
nit	number of iterations to call optim for optimization.
method	the optimization method to be used. See optim for details.
control	a list of control parameters to be passed to optim .
hessian	logical. Should a numerically differentiated Hessian matrix be returned?

Details

The optimization function [optim](#) is adopted in the above function to estimate the parameters and maximum likelihood. Several optimization methods are available for the optimization algorithm in [optim](#), but we recommend "Nelder-Mead" for the sake of stability. Alternatively, one may choose other options, e.g., "BFGS" to initialize and speed up the estimation procedure and then the procedure will automatically turn to "Nelder-Mead" for final results.

Normality is assumed for the random effects. Input data should be free of missing values.

Value

par	estimates of the model parameters.
value	log-likelihood of the model.
y	y used.

x associated with x used.
 v variance component matrices v used.
 ... other information.

See Also

[optim](#) and [rem](#).

Examples

```
data(miscEx)

## Not run:
# no sex effect
o<- estVC(y=pdatF8$bwt, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))
o

# sex as fixed effect
fo<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))
fo
2*(fo$value-o$value) # log-likelihood test statistic

# sex as random effect
SM<- rem(~sex, data=pdatF8)
ro<- estVC(y=pdatF8$bwt, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, SE=SM$sex, EE=diag(length(pdatF8$bwt))))
ro
2*(ro$value-o$value) # log-likelihood test statistic

## End(Not run)
```

genMatrix

Derive genetic matrices

Description

Derive genetic matrices from Jacquard condensed identity coefficients or genotypic data.

Usage

```
genMatrix(x)
```

Arguments

x an object of [cic](#) or [ibs](#), or genotypic data in a matrix or a data frame with each row representing an observation and each column a marker locus and entry being "AA", "AB", "BB" (or 1, 2, 3) without missing genotypes.

Value

AA	additive genetic matrix.
DD	dominance genetic matrix.
AD, HH, MH	other three genetic matrices (see Abney et. al. 2000).
ib	inbreeding coefficients.

References

Abney, M., M. S. McPeck, and C. Ober (2000). Estimation of variance components of quantitative traits in inbred populations. *Am. J. Hum. Genet.* 141, 629-650.

See Also

[cic](#)

Examples

```
data(miscEx)

ids<- sample(pedF8$id[300:500],20)

## Not run:
# get condensed identity coefficients
oo<- cic(pedF8, ids=ids, df=0)
ksp<- kinship(pedF8, ids=ids) # kinship coefficients only
# extract genetic matrices
gm<- genMatrix(oo)
sum((gm$AA-2*ksp)>1e-7) # same results

## End(Not run)
```

genoImpute

Impute Genotypic Data

Description

Impute missing genotypic data in advance intercross lines (AIL).

Usage

```
genoImpute(gdat, gmap, prd=NULL, step=Inf, gr=2, pos=NULL,
  method=c("Haldane", "Kosambi"), na.str="NA", verbose=FALSE)
```

Arguments

gdat	genotype data. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The column names should be marker names. Genotypes can be 1, 2 and 3, or "AA", "AB" and "BB". Optional if an object prd from genoProb is used as an argument.
gmap	a genetic map. Should be data frame (snp, chr, dist,...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome.
prd	an object from genoProb if not NULL. See "details" for more information.
step	the maximum distance (in cM) between two adjacent loci for which the probabilities are calculated. The distance corresponds to the "cumulative" recombination rate at gr-th generation.
gr	the generation under consideration.
pos	data frame (chr, dist, snp, ...). If given, step will be ignored.
method	whether "Haldane" or "Kosambi" mapping function should be used.
na.str	string for missing values.
verbose	a logical variable. If TRUE, certain information will be printed out during calculation.

Details

The missing genotypic value is randomly assigned with a probability conditional on the genotypes of the flanking SNPs (makers).

An object, prd, from [genoProb](#) alone can be used for the purpose of imputation. Then, the output (especially the putative loci) will be determined by prd. Optionally, it can be used together with gdat so that missing values in gdat will be imputed if possible, depending on whether loci in the columns of gdat can be identified in the third dimension of prd; this won't change the original genotypic data. See examples.

Value

A matrix with the number of rows being the same as gdat and with the number of columns depending on the SNP set in both gdat and gmap and the step length.

Note

Currently only suitable for advanced intercross lines.

See Also

[genoProb](#)

Examples

```

data(miscEx)

# briefly look at genotype data
sum(is.na(gdatF8))
gdatF8[1:5,1:5]

## Not run:
# run 'genoProb'
gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)

# imputation based on 'genoProb' object
tmp<- genoImpute(prd=prDat)
sum(is.na(tmp))
tmp[1:5,1:5]

# imputation based on both genotype data and 'genoProb' object
tmp<- genoImpute(gdatF8, prd=prDat)
sum(is.na(tmp))
tmp[1:5,1:5]

# imputation based on genotype data
tmp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA)
sum(is.na(tmp))
tmp[1:5, 1:5]
# set "verbose=TRUE" for more information
tmp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA, verbose=TRUE)
sum(is.na(tmp))
tmp[1:5, 1:5]

## End(Not run)

```

genoProb

Probability of a Genotype.

Description

Calculate the probability of a genotype at a locus conditional on the genotypes of its flanking markers in advance intercross lines (AIL).

Usage

```

genoProb(gdat, gmap, step=Inf, gr=2, pos=NULL, method=c("Haldane",
  "Kosambi"), verbose = FALSE)

```

Arguments

<code>gdat</code>	genotype data. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The column names should be marker names. Each entry should be 1, 2, 3 or 0, corresponding to "AA", "AB", "BB" or missing genotype.
<code>gmap</code>	a genetic map. Should be data frame (snp, chr, dist,...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome.
<code>step</code>	the maximum "cumulative" distance (in cM) between two adjacent loci for which the probabilities are calculated. The distance corresponds to the "cumulative" recombination rate at <code>gr</code> -th generation.
<code>gr</code>	the generation under consideration.
<code>pos</code>	data frame (chr, dist, snp, ...). If given, <code>step</code> will be ignored.
<code>method</code>	whether "Haldane" or "Kosambi" mapping function should be used.
<code>verbose</code>	a logical variable. If TRUE, certain information will be printed out during calculation.

Details

The "cumulative" genetic distance between any two adjacent loci for which probabilities are calculated is not larger than `step`. If `step = Inf`, probabilities will only be calculated at loci in both the columns of `gdat` and the rows of `gmap`. If `step` is small, a large set of putative loci will be considered, including all loci defined by the columns of `gdat` and the rows of `gmap`.

Value

<code>pr</code>	a 3-D array with the first dimension corresponding to that of <code>gdat</code> , the second to three genotype and the third to the putative loci. The probabilities will be -1 if not imputable, which happens when the genotype data is missing at all loci on the chromosome.
<code>chr</code>	chromosome where the locus is located.
<code>dist</code>	genetic distance (in cM) of the locus from the first locus on the chromosome.
<code>snp</code>	SNP (marker) that the locus represents.

Note

Currently only suitable for advanced intercross lines.

Examples

```
data(miscEx)

## Not run:
# briefly look at genotype data
sum(is.na(gdatF8))
```

```

gdatF8[1:5,1:5]

gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
# In case an individual is not imputable, then
# one needs to assign genotypes manually
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)
prDat$pr[1:5,,1:5]

## End(Not run)

```

genoSim *Generate Genotypic Data*

Description

Simulate genotypic data from a pedigree in advanced intercross lines (AIL).

Usage

```

genoSim(ped,gmap,ids,hap,method=c("Haldane","Kosambi"),
  recode.pedigree=FALSE)

```

Arguments

ped	a pedigree, which is a data frame (id, sex, sire, dam, ...). In "sex", male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", ... Note that 0 is reserved for missing values.
gmap	a genetic map. Should be data frame (snp, chr, dist, ...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome. If gmap is missing but hap not, all but the first two columns of hap are ignored.
ids	genotypic data are extracted only for individuals with IDs specified by ids. If missing, genotypic data are extracted for all individuals in the pedigree. If ped is an object of pedRecode , ids should be referred to "old" IDs.
hap	founders' haplotype data if not missing. Rows correspond to all founders, which should be in the first places in the pedigree ped, in the exact order and columns correspond to loci in the genetic map gmap in the exact order. For an individual, the haplotype should be (f1 m1 f2 m2 ...) where fi is the allele from father at the i-th locus and mi is the allele from mother at the i-th locus. Elements should be non-negative integers that are not larger than 16384. If missing, two founders with alleles 1 and 2 are assumed.
method	whether "Haldane" or "Kosambi" mapping function should be used. This will be ignored if the recombination rate recRate is a component of gmap.

```
recode.pedigree
```

a logical variable. True if the pedigree needs to be recoded.

Details

The pedigree should be in the same format as an output of [pedRecode](#). Note that two and only two founders are allowed; Otherwise, errors will occur without notice.

Value

a matrix, with entry value $s-1$ where s is the summation of the numbers representing two alleles at a locus. For instance, 1, 2, and 3 representing genotypes "AA", "AB" and "BB" respectively if hap is not specified. Each row represent an observation, and each column corresponds to SNP in gmap.

Note

Sex may be used as a covariate if significance on x-chromosome is assessed by gene dropping through this function.

See Also

[pedRecode](#) for more information.

Examples

```
data(miscEx)

## Not run:
# prepare pedigree in desired format
pedR<- pedRecode(pedF8)

# simulate genotype data for all individuals
gd1<- genoSim(pedR, gmapF8)
dim(gd1)

# simulate genotype data for all individuals
# no need to recode pedigree ahead
gd2<- genoSim(pedF8, gmapF8, recode.pedigree=TRUE)
dim(gd2)

# simulate genotypes for F8 individuals
gd3<- genoSim(pedR, gmapF8, ids=pedR$id[pedR$gen=="F8"],
  recode.pedigree=TRUE)
dim(gd3)
gd3[1:5,1:5]

## End(Not run)
```

gls	<i>Generalized Least Squares Estimates</i>
-----	--

Description

Obtain estimates using generalized least squares (gls).

Usage

```
gls(formula, data, vc=NULL)
```

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an data frame containing the variables in the model.
vc	an object from estVC or aicVC or an estimated variance-covariance matrix induced by relatedness and environment if not NULL.

Value

A matrix with columns: "Estimate", "Std. Error", "t value" and "Pr(>|t|)".

See Also

[lm](#).

hapSim	<i>Generate Genotypic Data</i>
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Description

Simulate gametic data from a pedigree.

Usage

```
hapSim(ped, gmap, ids, hap, method=c("Haldane", "Kosambi"),  
       recode.pedigree=FALSE)
```

Arguments

ped	a pedigree, which is a data frame (id, sex, sire, dam, ...). In "sex", male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", ... Note that 0 is reserved for missing values.
gmap	a genetic map. Should be data frame (snp, chr, dist, ...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome. If gmap is missing but hap not, all but the first two columns of hap are ignored.
ids	genotypic data are extracted only for individuals with IDs specified by ids. If missing, genotypic data are extracted for all individuals in the pedigree. If ped is an object of pedRecode , ids should be referred to "old" IDs.
hap	founders' haplotype data if not missing. Rows correspond to all founders, which should be in the first places in the pedigree ped, in the exact order and columns correspond to loci in the genetic map gmap in the exact order. For an individual, the haplotype should be (f1 m1 f2 m2 ...) where fi is the allele from father at the i-th locus and mi is the allele from mother at the i-th locus. Elements should be non-negative integers that are not larger than 16384. If missing, two founders with alleles 1 and 2 are assumed.
method	whether "Haldane" or "Kosambi" mapping function should be used. This will be ignored if the recombination rate recRate is a component of gmap.
recode.pedigree	a logical variable. True if the pedigree needs to be recoded.

Details

The pedigree should be in the same format as an output of [pedRecode](#).

Value

a matrix giving haplotypes.

See Also

[pedRecode](#) for more information.

Examples

```
data(miscEx)

## Not run:
# prepare pedigree in desired format
pedR<- pedRecode(pedF8)
# fake founder haplotypes
hapDat<- rbind(rep(1:2,nrow(gmapF8)),rep(3:4,nrow(gmapF8)))
# simulate hyplotypes for F8 individuals
hd<- hapSim(pedR, gmapF8, ids=pedR$id[pedR$gen=="F8"],
```



```
hap=hapDat, recode.pedigree=TRUE)
dim(hd)
hd[1:5,1:10]

## End(Not run)
```

ibs*Estimate Jacquard condensed identity coefficients*

Description

Estimate Jacquard condensed identity coefficients by identity-by-state (IBS) from genotypic data.

Usage

```
ibs(x)
```

Arguments

x genotype data with genotypes ("AA", "AB", "BB", or, 1, 2, 3) and without missing data, or probabilities for these genotypes (e.g., obtained by using [genoProb](#)). In case of genotype data, rows represent individuals and columns represent SNPs.

Value

A matrix G with $G[,j]$ being the j -th Jacquard identity coefficients.

Note

Currently only support the two-allele data.

See Also

[genMatrix](#)

kinship	<i>Calculate kinship coefficients</i>
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Description

Calculate kinship coefficients from a pedigree.

Usage

```
kinship(ped, ids)
```

Arguments

ped	a pedigree, which a data frame (id, sire, dam, ...). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", ... If "sex" is included, male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). Note that 0 is reserved for missing values.
ids	IDs of the individuals. If given, kinship coefficients are extracted for individuals with ID ids; otherwise, kinship coefficients are provided for all individuals in the pedigree.

Value

A matrix giving kinship coefficients.

Examples

```
data(miscEx)

ids<- sample(pedF8$id,10)
## Not run:
ksp<- kinship(pedF8,ids=ids)

## End(Not run)
```

lodci	<i>Estimate LOD Support Intervals</i>
-------	---------------------------------------

Description

Estimate LOD support intervals.

Usage

```
lodci(l1k, cv=0, lod=1.5, drop=3)
```

Arguments

llk	a data frame with components (chr, dist, y, ...), where "chr" is the chromosome on which the scanning locus is located, "dist" is the genetic or physical position of the scanning locus, and "y" is the test statistic.
cv	threshold. Reported support intervals cover at least one scanning locus where $llk\$y > cv$.
lod	lod (1.5 by default) LOD support intervals are reported when llk\$y is converted to LOD score.
drop	3 by default. See "details".

Details

In case of multiple peaks on a chromosome, a peak has to satisfy: a) above the threshold cv; b) drops, e.g., 3 LOD on both sides except chromosome ends. So if two peaks close to each other but LOD between them doesn't drop, e.g., 3 LOD, only one of them is considered.

Value

A data frame with the following components:

chr	the chromosome
lower	the lower bound
upper	the upper bound
index	indicates which scanning loci

Examples

```
data(miscEx)

## Not run:
# impute missing genotypes
gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
# run 'genoProb'
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)
# estimate variance components
o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))

# genome scan
llk.hk<- scanOne(y=pdatF8$bwt, x=pdatF8$sex, vc=o, prdat=prDat)

# extract LOD support intervals
tmp<- data.frame(y=llk.hk$p, chr=llk.hk$chr, dist=llk.hk$dist)
lodci(tmp, cv=10, lod=1.5, drop=3)

## End(Not run)
```

mAIC

*Multiple QTL AIC***Description**

Multiple QTL model selection by AIC criterion.

Usage

```
mAIC(y, x, gdat, prdat=NULL, vc=NULL, chrIdx, xin, k=2,
      direction=c("both", "backward", "forward"), ext=FALSE, verbose=FALSE)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
gdat	genotype data. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The column names should be marker names. Numeric coding of genotype is treated as numeric. Ignored if prdat is an object from genoProb .
vc	an object from estVC or aicVC , or an estimated variance-covariance matrix induced by relatedness. The scan will assume no polygenic variation if vc is NULL.
prdat	an object from genoProb .
chrIdx	chromosome index of markers in columns of gdat if given. Ignored if prdat is an object from genoProb .
xin	vector indicating whether a locus is already in the model.
k	penalty on a parameter. The selection criterion is the known "AIC" if $k = 2$ and is "BIC" if $k = \log(n)$ where "n" is the sample size.
direction	the mode of search: "both", "forward" or "backward" with default "both".
ext	a logical variable. True if ones wants more exhaustive search.
verbose	a logical variable. True if ones wants to track the process for monitoring purpose.

Details

Makes use of "Haley-Knott" method (Haley and Knott 1992) if prdat is an object from [genoProb](#).

Value

a list with the following components:

model: the resulting model;

aic: AIC of the model;

snp: selected SNPs.

xin: vector indicating whether a SNP is selected.

Note

Currently only suitable for advanced intercross lines (or diallelic data).

References

Haley, C. S., and S. A. Knott (1992). A simple regression method for mapping quantitative trait loci in line crosses using flanking markers. *Heredity* 69: 315-324.

See Also

[optim](#), [genoProb](#) and [aicVC](#).

Examples

```
data(miscEx)

## Not run:
# impute missing genotypes
gdat.imp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA)
# estimate variance components
o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
  HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))

# run 'genoProb'
gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)

# genome scan
llk.hk<- scanOne(y=pdatF8$bwt, x=pdatF8$sex, prdat=prDat, vc=o)
xin<- llk.hk$p > 10

# run 'mAIC' based on genome scan results
mg<- mAIC(y=pdatF8$bwt, x=pdatF8$sex, prdat=prDat, vc=o, xin=xin,
  k=5, direction="back", verbose=TRUE)
mg$model$value # likelihood of the final model

## End(Not run)
```

miscEx	<i>Genotype data, phenotype data, genetic map and pedigree.</i>
--------	---

Description

AIL F8 data include the following:

"gmF8": a list with elements inbreeding coefficients "ib", additive genetic matrix "AA", dominance genetic matrix "DD" and other genetic matrices.

"pedF8": pedigree data.

"gmapF8": genetic map.

"gdatF8": genotype data.

"pdatF8": phenotype data.

Usage

```
data(miscEx)
```

misFct	<i>A collection of other functions.</i>
--------	---

Description

A collection of other functions that are not needed by users.

nullSim	<i>Simulate null distribution</i>
---------	-----------------------------------

Description

Simulate distribution under the hypothesis of no QTL by permutation (of genotypic data) or gene dropping.

Usage

```
nullSim(y, x, gdat, prdat, ped, gmap, hap,
method=c("permutation", "gene dropping"), vc=NULL, intcovar=NULL,
test = c("None", "F", "Chisq"), minorGenoFreq=0.05, rmv=TRUE,
recode.pedigree=FALSE, gr=2, ntimes=10)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
gdat	genotype data without missing values. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. Ignored in the case of gene dropping.
prdat	an object from genoProb , or in the same form.
ped	a pedigree, which is a data frame (id, sex, sire, dam, ...). In "sex", male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", ... Note that 0 is reserved for missing values. Ignored in the case of permutation.
gmap	a genetic map. Should be data frame (snp, chr, dist, ...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome. Ignored in the case of permutation.
hap	founders' haplotype data if not missing. Rows correspond to all founders, which should be in the first places in the pedigree ped, in the exact order and columns correspond to loci in the genetic map gmap in the exact order. For an individual, the haplotype should be (f1 m1 f2 m2 ...) where fi is the allele from father at the i-th locus and mi is the allele from mother at the i-th locus. Elements should be non-negative integers that are not larger than 16384. If missing, two founders with alleles 1 and 2 are assumed.
method	permutation or gene dropping.
vc	an object from estVC or aicVC , or an estimated variance-covariance matrix induced by relatedness. The scan will assume no polygenic variation if vc is NULL.
intcovar	Covariates that interact with QTL.
test	"None", "F" or "Chisq".
minorGenoFreq	specify the minimum tolerable minor genotype frequency at a scanning locus if gdat is used.
rmv	a logical variable. If true, then the scanning locus will be skipped if the minor genotype frequency at the locus is smaller than <code>minorGenoFreq</code> . Otherwise, the scanning process will stop and return with NULL.
recode.pedigree	a logical variable. True if the pedigree needs to be recoded.
gr	the generation under consideration.
ntimes	number of simulations.

Details

Two methods considered here are permutation test and gene dropping test as described as follows.

Permutation test. Depending on the genome-scan, one can provide either `gdat` or `prdat` respectively corresponding to single-marker analysis or interval mapping. Then only arguments in `scanOne` are needed in addition to `method` and `ntimes`.

Gene dropping test. If `prdat` is provided, then `gdat` will be ignored. The procedure will first call `genoSim` to generate new genotype data and then call `genoProb` to generate data for Haley-Knott interval mapping. If `prdat` is not provided, then `gdat` should be provided. The procedure will generate new genotype data and scan the genome using these generated genotype data. Haldane mapping function is used to generate data.

Value

A vector of numbers of length `ntimes` if `minorGenoFreq > 0` and `rmv = TRUE`, each element of which is maximum of the test statistic over the genome scan (so test should be "None"), or a matrix of `ntimes` rows, each row of which records a genome scan.

See Also

[genoSim](#), [genoProb](#) and [scanOne](#).

Examples

```
data(miscEx)

## Not run:
# recode the pedigree for later use
pedR<- pedRecode(pedF8)

# impute missing genotypes
gdatTmp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA)
# estimate variance components
o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
  HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))

# scan marker loci & permutation
ex1<- nullSim(y=pdatF8$bwt, x=pdatF8$sex, gdat=gdatTmp,
  method="permutation", vc=o, ntimes=10)
dim(ex1)

# scan marker loci & gene dropping
ex2<- nullSim(y=pdatF8$bwt, x=pdatF8$sex, gdat=gdatTmp, ped=pedR,
  gmap=gmapF8, method="gene", vc=o, ntimes=10)
dim(ex2)

# Haley-Knott method & permutation
gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)
ex3<- nullSim(y=pdatF8$bwt, x=pdatF8$sex, prdat=prDat,
  method="permutation", vc=o, ntimes=10)
```



```

dim(ex3)

# Haley-Knott method & gene dropping
ex4<- nullSim(y=pdatF8$bwt, x=pdatF8$sex, prdat=prDat, ped=pedR,
gmap=gmapF8, method="gene", vc=o, gr=8, ntimes=10)
dim(ex4)

## End(Not run)

```

pedRecode

Recode a Pedigree

Description

Prepare a pedigree in a format that is suitable for certain functions

Usage

```
pedRecode(ped, ids)
```

Arguments

ped	a pedigree, which is a data frame (id, sire, dam, ...). If "sex" is a component, male should be "M", "Male" or 1, and female should be "F", "Female" or 2 (other than 0 and 1). If given, "generation" can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2", etc. 0 is reserved for unknown sire or dam.
ids	If given, only individuals with ids and their ancestors are kept in the recoded pedigree.

Value

A recoded pedigree.

Examples

```

data(miscEx)

pedF8[1:10,]
pedR<- pedRecode(pedF8)
pedR[1:10,]
dim(pedR)
pedR<- pedRecode(pedF8, ids=pedF8$id[pedF8$gener=="F8"])
dim(pedR)

```

plotit

*Plotting***Description**

Plot mapping results.

Usage

```
## S3 method for class 'scanOne'
plot(x, ...)

plotit(lrt, cv, bychr=FALSE, chr.labels=TRUE, type="p", lty=NULL,
       col=NULL, pch=NULL, cex=NULL, ...)
```

Arguments

x	object from scanOne or scanTwo .
lrt	a data frame with (chr, dist, y,...) or (chr, dist, y, group,...), where "chr" represents chromosome, "dist" position on the chromosome, "y" the test statistic.
cv	threshold to be drawn on the plot.
cex	see par .
bychr	a logical variable. If true, the plot will be displayed per chromosomes.
chr.labels	a logical variable. If true, the chromosome names will be drawn.
type, lty, col, pch	See plot.default .
...	other options passed to R plot function. To call plot to plot results of scanOne , one may need to provide a genetic map gmap that should be data frame (snp, chr, dist, ...), where "snp" is the SNP (marker) name, "chr" is the chromosome where the "snp" is, and "dist" is the genetic distance in centi-Morgan (cM) from the leftmost SNP (marker) on the chromosome.

Note

A genetic map 'gmap' may be needed to plot an object of [scanOne](#) or [scanTwo](#). The color option may not give what is expected.

Examples

```
data(miscEx)

## Not run:
# impute missing genotypes
gdat.imp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA)
# estimate variance components
```

```

o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))

# genome scan
llk<- scanOne(y=pdatF8$bwt, x=pdatF8$sex, vc=o, gdat=gdat.imp)

# plotting
plot(llk, gmap=gmapF8) # gmap is needed

# plotting in another way
idx<- match(colnames(gdat.imp), gmapF8$snp)
tmp<- data.frame(chr=gmapF8$chr[idx],dist=gmapF8$dist[idx],y=llk$p)
plotit(tmp, main="Mapping Plot", xlab="Chromosome", ylab="LRT",
       col=as.integer(tmp$ch)%2+2,type="p")

## End(Not run)

```

qqPlot

*Quantile-Quantile Plots***Description**

Quantile-Quantile Plots With the Ability to Draw Confidence Bands.

Usage

```

qqPlot(y,x="norm", ...,
       xlab=if(is.numeric(x)) deparse(substitute(x)) else x,
       ylab=deparse(substitute(y)),main="Q-Q Plot",
       type="p",col=1,lty=2,lwd=1,pch=1,cex=0.7,plot.it=TRUE,
       confidence=.95,qqline=c("observed", "expected", "none"))

```

Arguments

y	a numeric vector of data values.
x	either a numeric vector of data values, or a character string naming a distribution function such as "norm".
...	parameters passed to the distribution specified by x (if non-numerical).
xlab	a label for the x axis.
ylab	a label for the y axis.
main	a main title for the plot.
type	1-character string giving the type of plot desired.
col	color for points and lines.
lty	line type.
lwd	line width.

pch	plotting character for points.
cex	factor for expanding the size of plotted symbols.
plot.it	whether or not to draw a plot. if plotting, points outside the confidence bands will be indicated by different a color.
confidence	confidence level for the confidence band, or FALSE for no band.
qqline	whether or not to draw a reference line. if "observed", the line passes through the first and third observed quartiles; if "expected", the point (x,y) is expected to fall on the line if x and y follow the same distribution; if "none", no reference line is drawn.

Details

If `x` is numeric, a two-sample test of the null hypothesis that `x` and `y` were drawn from the same continuous distribution is performed. Alternatively, `x` can be a character string naming a continuous distribution function. In such a case, a one-sample test is carried out of the null that `y` was draw from distribution `x` with parameters specified by "...".

Value

<code>x</code>	quantiles of <code>x</code>
<code>y</code>	quantiles of <code>y</code>
<code>lower</code> , <code>upper</code>	lower and upper limits if confidence is specified

References

George Marsaglia, Wai Wan Tsang and Jingbo Wang (2003), Evaluating Kolmogorov's distribution. *Journal of Statistical Software* 8 (18): 1-4.

Vijayan N. Nair (1982). Q-Q plots with confidence bands for comparing several populations.

William J. Conover (1971). *Practical Nonparametric Statistics*. New York: John Wiley & Sons.

See Also

[ks.test](#).

Examples

```
## Not run:
par(mfrow=c(1,2))
x<- rnorm(200, mean=0.7,sd=2); y<- rnorm(200, sd=2)
qqPlot(y,x,qqline="exp")
qqPlot(y=y,x="norm",sd=2)
ks.test(x,y)

## End(Not run)
```

`qt12rel`*Convert data from R/qt1 to QTLRel format*

Description

Convert the data for a QTL mapping experiment from the R/qt1 format (see <http://www.rqt1.org>) to that used by QTLRel.

Usage

```
qt12rel(cross)
```

Arguments

`cross` An object of class "cross", as defined by the R/qt1 package

Details

The input cross must be an intercross (class "f2").

Simple pedigree information is created, assuming the data are from a standard intercross.

Value

A list with four components: "ped" (pedigree information), "gdat" (genotype data), "pdat" (phenotype data), and "gmap" (genetic map), in the formats used by QTLRel.

Author(s)

Karl W Broman, <kbroman@biostat.wisc.edu>

See Also

[rel2qt1](#)

Examples

```
library(qt1)
data(listeria)
listeria <- listeria[as.character(1:19),]
reldat <- qt12rel(listeria)
```

qtlVar

*QTL Variance***Description**

Estimate variance in a quantitative trait induced by QTL.

Usage

```
qtlVar(lrt,prdat,simulation=FALSE,nsim=25)
```

Arguments

lrt	a data frame (a, d, ...), where 'a' and 'd' are respectively additive and dominance effects.
prdat	a 3-D array that provides probabilities of genotypes "AA", "AB" and "BB". If prDat is an object of genoProb , then prdat can be prDat\$pr.
simulation	whether to use simulations to estimate the variance explained by QTL.
nsim	number of simulations to perform if simulation is TRUE.

Value

A vector displaying the estimated variance at each loci.

Note

Correlations among observations are ignored, and this function should be used with caution.

See Also

[scanOne](#) and [genoProb](#)

Examples

```
data(miscEx)

## Not run:
# impute missing genotypes
gdtmp<- gdatF8
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
# rung 'genoProb'
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)
# estimate variance components
o<- estVC(y=pdatF8$bwt, x=pdatF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
  HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdatF8$bwt))))

# genome scan
```

```
pv.hk<- scanOne(y=pdatF8$bwt, x=pdatF8$sex, prdat=prDat, vc=o)

# run 'qtlVar'
qef<- NULL
for(n in 1:length(11k.hk$par))
  qef<- rbind(qef,11k.hk$par[[n]][c("a","d")])
  qef<- as.data.frame(qef)
qtlVar(qef,prDat$pr)[1:3]

## End(Not run)
```

rel2qtl	<i>Convert data from QTLRel to R/qtl format</i>
---------	---

Description

Convert the data for a QTL mapping experiment from the QTLRel format to that used by R/qtl (<http://www.rqtl.org>).

Usage

```
rel2qtl(gdat, pdat, gmap)
```

Arguments

gdat	Genotype data
pdat	Phenotype data
gmap	Genetic map

Details

Pedigree information is ignored, and X chromosome data is omitted.
The data are treated as an intercross.

Value

A cross object for the R/qtl package (<http://www.rqtl.org>).

Author(s)

Karl W Broman, <kbroman@biostat.wisc.edu>

See Also

[qtl2rel](#)

Examples

```
data(miscEx)
f8 <- rel2qt1(gdatF8, pdatF8, gmapF8)
summary(f8)
```

 rem

Random effect matrices

Description

Construct matrices associated with random effects.

Usage

```
rem(formula,data)
```

Arguments

formula	a formula of the form: $\sim Z \mid G1/.../Gk + \dots$, corresponding to random effects $Z * G_i + Z * G_{\{ij\}} + \dots$ in a mixed effect model. If $Z=1$ as in most cases, then it can be $\sim G1/.../Gk + \dots$
data	a data frame that contains all the variables in the formula.

Value

a list of matrices that are associated with random effects.

Examples

```
## Not run:
# make-up example
dat<- data.frame(
  group=c("A", "A", "A", "A", "A", "A", "B", "B", "B", "B"),
  sex=c("F", "F", "F", "M", "M", "M", "F", "F", "M", "M"),
  pass=c("Y", "N", "N", "Y", "Y", "Y", "Y", "N", "N", "Y"),
  z=1:10)

# random effect pass, group and sex, where sex is nested
# within group:
# y_{ijk} = x_{ij}*b + group_i + sex_{ij} + z*pass_{ij}
#           + e_{ijk}
rem(~ group/sex + z|pass,data=dat)

## End(Not run)
```

scanOne	<i>Genome Scan for QTL</i>
---------	----------------------------

Description

Evaluate log-likelihood ratio test statistics or P-values at scanning loci along the genome.

Usage

```
scanOne(y, x, gdat, prdat=NULL, vc=NULL, intcovar=NULL,
        numGeno=FALSE, test=c("None", "F", "Chisq"),
        minorGenoFreq=0, rmv=TRUE)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
gdat	genotype data. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The column names should be marker names. Optional if an object prdat from genoProb is used as an argument.
prdat	an object from genoProb , or in the same form.
vc	an object from estVC or aicVC , or an estimated variance-covariance matrix induced by relatedness and environment.
intcovar	Covariates that interact with QTL.
numGeno	whether to treat numeric coding of genotypes as numeric.
test	"None", "F" or "Chisq".
minorGenoFreq	specify the minimum tolerable minor genotype frequency at a scanning locus if gdat is used.
rmv	a logical variable. If true, then the scanning locus will be skipped if the minor genotype frequency at the locus is smaller than minorGenoFreq. Otherwise, the scanning process will stop and return with NULL.

Details

The test at a scanning locus under the assumption of no QTL effect versus the assumption of QTL effect is performed by conditioning on the estimated polygenic genetic variance-covariance matrix. Normality is assumed for the random effects.

It is possible to extend the Haley-Knott approach to multiple-allelic cases under the assumption that allele effects are all additive. Then, prdat should be provided and be of class "addEff".

Value

A list with at least the following components:

p	P-value at the snp (marker) if test is "F" or "Chisq", or the log-likelihood ratio statistic at the SNP (marker) if "test" is "None"
v	Percentage of variation explained by QTL related effects at the snp (marker)
parameters	estimated parameters at all scanning loci, including additive effect a and dominance effect d if prdat is not NULL

References

Haley, C. S., and S. A. Knott (1992). A simple regression method for mapping quantitative trait loci in line crosses using flanking markers. *Heredity* 69: 315-324.

See Also

[genoImpute](#) and [genoProb](#).

Examples

```
data(miscEx)

## Not run:
# impute missing genotypes
gdatTmp<- genoImpute(gdatF8, gmap=gmapF8, step=Inf,
  gr=8, na.str=NA)

# estimate variance components
o<- estVC(y=pdafF8$bwt, x=pdafF8$sex, v=list(AA=gmF8$AA,DD=gmF8$DD,
HH=NULL, AD=NULL, MH=NULL, EE=diag(length(pdafF8$bwt))))

# genome scan and plotting
pv<- scanOne(y=pdafF8$bwt, x=pdafF8$sex, gdat=gdatTmp, vc=o)
pv
plot(pv,gmap=gmapF8)

# Haley-Knott method
gdtmp<- gdatF8; unique(unlist(gdtmp))
  gdtmp<- replace(gdtmp,is.na(gdtmp),0)
prDat<- genoProb(gdat=gdtmp, gmap=gmapF8, step=Inf,
  gr=8, method="Haldane", verbose=TRUE)
pv.hk<- scanOne(y=pdafF8$bwt, x=pdafF8$sex, prdat=prDat, vc=o)
pv.hk
plot(pv.hk)

## End(Not run)
```

 scanTwo

Genome Scan for Epistasis

Description

Evaluate log-likelihood ratio test statistic for epistasis (QTL by QTL interaction).

Usage

```
scanTwo(y, x, gdat, prdat=NULL, vc=NULL, numGeno=FALSE,
        minorGenoFreq=0, rmv=TRUE)
```

Arguments

y	a numeric vector or a numeric matrix of one column (representing a phenotype for instance).
x	a data frame or matrix, representing covariates if not missing.
gdat	genotype data. Should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The column names should be marker names. Optional if an object prdat from genoProb is used as an argument.
prdat	an object from genoProb .
vc	an object from estVC or aicVC , or an estimated variance-covariance matrix induced by relatedness and environment.
numGeno	whether to treat numeric coding of genotypes as numeric.
minorGenoFreq	specify the minimum tolerable minor genotype frequency at a scanning locus if gdat is used.
rmv	a logical variable. If true, then the scanning locus will be skipped if the minor genotype frequency at the locus is smaller than minorGenoFreq. Otherwise, the scanning process will stop and return with NULL.

Value

A matrix whose entry in the upper triangle is the log-likelihood test statistic for epistatic effect.

See Also

[scanOne](#).

Index

*Topic **datasets**
miscEx, 22

*Topic **manip**
qtl2rel, 29
rel2qtl, 31

aicVC, 2, 4, 5, 7, 15, 20, 21, 23, 33, 35

blup, 4

cic, 5, 8, 9

estVC, 3–5, 7, 15, 20, 23, 33, 35

fv (misFct), 22

gdatF8 (miscEx), 22
genMatrix, 8, 17
genoImpute, 9, 34
genoPos (misFct), 22
genoProb, 10, 11, 17, 20, 21, 23, 24, 30, 33–35
genoSim, 13, 24
gls, 15
gmapF8 (miscEx), 22
gmF8 (miscEx), 22
gvar (misFct), 22

hapSim, 15

ibs, 8, 17

kinship, 18
ks.test, 28

lm, 15
lodci, 18

mAIC, 20
miscEx, 22
misFct, 22

nullSim, 22

optim, 3, 7, 8, 21

par, 26
pdatF8 (miscEx), 22
pedF8 (miscEx), 22
pedRecode, 13, 14, 16, 25
pkolm (misFct), 22
plot, 26
plot (plotit), 26
plot.default, 26
plotit, 26

qkolm (misFct), 22
qqPlot, 27
qtl2rel, 29, 31
qtlVar, 30

rel2qtl, 29, 31
rem, 8, 32

scanOne, 24, 26, 30, 33, 35
scanTwo, 26, 35