

Package ‘hsrecombi’

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Type Package

Title Estimation of Recombination Rate and Maternal LD in Half-Sibs

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Description Paternal recombination rate and maternal linkage disequilibrium (LD) are estimated for pairs of biallelic markers such as single nucleotide polymorphisms (SNPs) from progeny genotypes and sire haplotypes. The implementation relies on paternal half-sib families. If maternal half-sib families are used, the roles of sire/dam are swapped. Multiple families can be considered. For parameter estimation, at least one sire has to be double heterozygous at the investigated pairs of SNPs.

Based on recombination rates, genetic distances between markers can be estimated. Markers with unusually large recombination rate to markers in close proximity (i.e. putatively misplaced markers) shall be discarded in this derivation.

A workflow description is attached as vignette.

A pipeline is available at GitHub

<<https://github.com/wittenburg/hsrecombi>>

Hampel, Teuscher, Gomez-Raya, Doschoris, Wittenburg (2018) ``Estimation of recombination rate and maternal linkage disequilibrium in half-sibs''

<doi:10.3389/fgene.2018.00186>.

Gomez-Raya (2012) ``Maximum likelihood estimation of linkage disequilibrium in half-sib families'' <doi:10.1534/genetics.111.137521>.

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Author Dörte Wittenburg [aut, cre]

Maintainer Dörte Wittenburg <wittenburg@fbn-dummerstorf.de>

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bestmapfun

Best fitting map function

Description

Approximation of mixing parameter of system of map functions

Usage

```
bestmapfun(theta, dist_M)
```

Arguments

theta	vector of recombination rates
dist_M	vector of genetic positions

Details

The genetic mapping function that fits best to the genetic data (recombination rate and genetic distances) is obtained from Rao's system of mapping functions. The corresponding mixing parameter is estimated via 1-dimensional constrained optimisation. See vignette for its application to estimated data.

Value

list (LEN 2)

mixing mixing parameter of system of genetic mapping functions

mse minimum value of target function ($\theta - \text{dist_M}$) 2

References

Rao, D.C., Morton, N.E., Lindsten, J., Hulten, M. & Yee, S (1977) A mapping function for man. Human Heredity 27: 99-104. doi: [10.1159/000152856](https://doi.org/10.1159/000152856)

Examples

```
theta <- seq(0, 0.5, 0.01)
gendist <- -log(1 - 2 * theta) / 2
bestmapfun(theta, gendist)
```

checkCandidates

Candidates for misplacement

Description

Search for SNPs with unusually large estimates of recombination rate

Usage

```
checkCandidates(final, win = 30, quant = 0.99)
```

Arguments

final	table of results produced by editraw with pairwise estimates of recombination rate between p SNPs within chromosome; minimum required data frame with columns SNP1, SNP2 and theta
win	optional value for window size; default value 30
quant	optional value; default value 0.99, see details

Details

Markers with unusually large estimates of recombination rate to close SNPs are candidates for misplacements in the underlying assembly. The mean of recombination rate estimates with `win` subsequent or preceding markers is calculated and those SNPs with mean value exceeding the `quant` quantile are denoted as candidates which have to be manually curated! This can be done, for instance, by visual inspection of a correlation plot containing estimates of recombination rate in a selected region.

Value

vector of SNP indices for further verification

References

Hampel, A., Teuscher, F., Gomez-Raya, L., Doschoris, M. & Wittenburg, D. (2018) Estimation of recombination rate and maternal linkage disequilibrium in half-sibs. *Frontiers in Genetics* 9:186. doi: [10.3389/fgene.2018.00186](https://doi.org/10.3389/fgene.2018.00186)

Examples

```
### test data
data(targetregion)
### make list for paternal half-sib families
hap <- makehaplist(daughterSire, hapSire)
### parameter estimates on a chromosome
res <- hsrecombi(hap, genotype.chr, map.chr$SNP)
### post-processing to achieve final and valid set of estimates
final <- editraw(res, map.chr)
### check for candidates of misplacement
snp <- checkCandidates(final)
```

countNumbers

Count genotype combinations at 2 SNPs

Description

Count genotype combinations at 2 SNPs

Arguments

X	integer matrix of genotypes
---	-----------------------------

Value

count vector of counts of 9 possible genotypes at SNP pair

daughterSire*targetregion: allocation of paternal half-sib families*

Description

Vector of sire ID for each progeny

Usage

```
daughterSire
```

Format

An object of class `integer` of length 265.

editraw*Editing results of hsrecombi*

Description

Process raw results from `hsrecombi`, decide which out of two sets of estimates is more likely and prepare list of final results

Usage

```
editraw(Roh, map1)
```

Arguments

Roh	list of raw results from <code>hsrecombi</code>
map1	data.frame containing information on physical map, at least: SNP SNP ID locus_Mb physical position in Mbp of SNP on chromosomes Chr chromosome of SNP

Value

final table of results

SNP1 index 1. SNP

SNP2 index 2. SNP

D maternal LD

fAA frequency of maternal haplotype 1-1

fAB frequency of maternal haplotype 1-0

fBA frequency of maternal haplotype 0-1
fBB frequency of maternal haplotype 0-0
p1 Maternal allele frequency (allele 1) at SNP1
p2 Maternal allele frequency (allele 1) at SNP2
nfam1 size of genomic family 1
nfam2 size of genomic family 2
error 0 if computations were without error; 1 if EM algorithm did not converge
iteration number of EM iterations
theta paternal recombination rate
r2 r^2 of maternal LD
logL value of log likelihood function
unimodal 1 if likelihood is unimodal; 0 if likelihood is bimodal
critical 0 if parameter estimates were unique; 1 if parameter estimates were obtained via decision process
locus_Mb physical distance between SNPs in Mbp

Examples

```

### test data
data(targetregion)
### make list for paternal half-sib families
hap <- makehaplist(daughterSire, hapSire)
### parameter estimates on a chromosome
res <- hsrecombi(hap, genotype.chr, map.chr$SNP)
### post-processing to achieve final and valid set of estimates
final <- editraw(res, map.chr)

```

geneticPosition *Estimation of genetic position*

Description

Estimation of genetic positions (in centi Morgan)

Usage

```
geneticPosition(final, exclude = NULL, threshold = 0.05)
```

Arguments

final	table of results produced by editraw with pairwise estimates of recombination rate between p SNPs within chromosome; minimum required data frame with columns SNP1, SNP2 and theta
exclude	optional vector (LEN q) of SNPs to be excluded (e.g., candidates of misplaced SNPs)
threshold	optional value; recombination rates \leq threshold are considered for smoothing

Details

Smoothing of recombination rates (θ) ≤ 0.05 via quadratic optimization provides an approximation of genetic distances (in Morgan) between SNPs. The cumulative sum * 100 yields the genetic positions in cM.

The minimization problem $(\theta - D d)^2$ is solved s.t. $d > 0$ where d is the vector of genetic distances between adjacent markers but θ is not restricted to adjacent markers. The incidence matrix D contains 1's for those intervals contributing to the total distance relevant for each θ .

Estimates of $\theta = 1e-6$ are neglected as these values coincide with start values and indicate that (because of a very flat likelihood surface) no meaningful estimate of recombination rate has been obtained.

Value

vector (LEN p) of genetic positions of SNPs (in cM)

References

Qanbari, S. & Wittenburg, D. (2020) Male recombination map of the autosomal genome in German Holstein. Genetics Selection Evolution 52:73. doi: [10.1186/s1271102000593z](https://doi.org/10.1186/s1271102000593z)

Examples

```
### test data
data(targetregion)
### make list for paternal half-sib families
hap <- makehaplist(daughterSire, hapSire)
### parameter estimates on a chromosome
res <- hsrecombi(hap, genotype.chr, map.chr$SNP)
### post-processing to achieve final and valid set of estimates
final <- editraw(res, map.chr)
### approximation of genetic positions
pos <- geneticPosition(final)
```

genotype.chr

targetregion: progeny genotypes

Description

matrix of progeny genotypes in target region on chromosome BTA1

Usage

genotype.chr

Format

An object of class `matrix` (inherits from `array`) with 265 rows and 300 columns.

<code>hapSire</code>	<i>targetregion: sire haplotypes</i>
----------------------	--------------------------------------

Description

matrix of sire haplotypes in target region on chromosome BTA1

Usage

```
hapSire
```

Format

An object of class `matrix` (inherits from `array`) with 10 rows and 301 columns.

<code>hsrecombi</code>	<i>Estimation of recombination rate and maternal LD</i>
------------------------	---

Description

Wrapper function for estimating recombination rate and maternal linkage disequilibrium between intra-chromosomal SNP pairs by calling EM algorithm

Usage

```
hsrecombi(hap, genotype.chr, snp.chr, only.adj = FALSE, prec = 1e-06)
```

Arguments

<code>hap</code>	list (LEN 2) of lists
<code>famID</code>	list (LEN number of sires) of vectors (LEN n.progeny) of progeny indices relating to lines in genotype matrix
<code>sireHap</code>	list (LEN number of sires) of matrices (DIM 2 x p) of sire haplotypes (0, 1) on investigated chromosome
<code>genotype.chr</code>	matrix (DIM n x p) of all progeny genotypes (0, 1, 2) on a chromosome with p SNPs
<code>snp.chr</code>	vector(LEN p) of SNP indices as in physical map
<code>only.adj</code>	logical; if TRUE, recombination rate is calculated only between neighbouring markers
<code>prec</code>	scalar; precision of estimation

Details

Paternal recombination rate and maternal linkage disequilibrium (LD) are estimated for pairs of biallelic markers (such as single nucleotide polymorphisms; SNPs) from progeny genotypes and sire haplotypes. At least one sire has to be double heterozygous at the investigated pairs of SNPs. All progeny are merged in two genomic families: (1) coupling phase family if sires are double heterozygous 0-0/1-1 and (2) repulsion phase family if sires are double heterozygous 0-1/1-0. So far it is recommended processing the chromosomes separately. If maternal half-sib families are used, the roles of sire/dam are swapped. Multiple families can be considered.

Value

list (LEN p - 1) of data.frames; for each SNP, parameters are estimated with all following SNPs; two solutions (prefix sln1 and sln2) are obtained for two runs of the EM algorithm

SNP1 index 1. SNP

SNP2 index 2. SNP

D maternal LD

fAA frequency of maternal haplotype 1-1

fAB frequency of maternal haplotype 1-0

fBA frequency of maternal haplotype 0-1

fBB frequency of maternal haplotype 0-0

p1 Maternal allele frequency (allele 1) at SNP1

p2 Maternal allele frequency (allele 1) at SNP2

nfam1 size of genomic family 1

nfam2 size of genomic family 2

error 0 if computations were without error; 1 if EM algorithm did not converge

iteration number of EM iterations

theta paternal recombination rate

r2 r^2 of maternal LD

logL value of log likelihood function

unimodal 1 if likelihood is unimodal; 0 if likelihood is bimodal

critical 0 if parameter estimates are unique; 1 if parameter estimates at both solutions are valid, then decision process follows in post-processing function "editraw"

Afterwards, solutions are compared and processed with function editraw, yielding the final estimates for each valid pair of SNPs.

References

Hampel, A., Teuscher, F., Gomez-Raya, L., Doschoris, M. & Wittenburg, D. (2018) Estimation of recombination rate and maternal linkage disequilibrium in half-sibs. Frontiers in Genetics 9:186. doi: [10.3389/fgene.2018.00186](https://doi.org/10.3389/fgene.2018.00186)

Gomez-Raya, L. (2012) Maximum likelihood estimation of linkage disequilibrium in half-sib families. Genetics 191:195-213.

Examples

```
### test data
data(targetregion)
### make list for paternal half-sib families
hap <- makehaplist(daughterSire, hapSire)
### parameter estimates on a chromosome
res <- hsrecombi(hap, genotype.chr, map.chr$SNP)
### post-processing to achieve final and valid set of estimates
final <- editraw(res, map.chr)
```

LDHScpp

Expectation Maximisation (EM) algorithm

Description

Expectation Maximisation (EM) algorithm

Usage

```
LDHScpp(XGF1, XGF2, fAA, fAB, fBA, theta, display, threshold)
```

Arguments

XGF1	integer matrix of progeny genotypes in genomic family 1
XGF2	integer matrix of progeny genotypes in genomic family 2
fAA	frequency of maternal haplotype 1-1
fAB	frequency of maternal haplotype 1-0
fBA	frequency of maternal haplotype 0-1
theta	paternal recombination rate
display	logical for displaying additional information
threshold	convergence criterion

Value

- list of parameter estimates
- D maternal LD
- fAA frequency of maternal haplotype 1-1
- fAB frequency of maternal haplotype 1-0
- fBA frequency of maternal haplotype 0-1
- fBB frequency of maternal haplotype 0-0
- p1 Maternal allele frequency (allele 1) at 1. SNP
- p2 Maternal allele frequency (allele 1) at 2. SNP
- nfam1 size of genomic family 1

```

nfam2 size of genomic family 2
error 0 if computations were without error; 1 if EM algorithm did not converge
iteration number of EM iterations
theta paternal recombination rate
r2  $r^2$  of maternal LD
logL value of log likelihood function

```

loglikfun*Calculate log-likelihood function***Description**

Calculate log-likelihood function

Arguments

counts	integer vector of observed 2-locus genotype
fAA	frequency of maternal haplotype 1-1
fAB	frequency of maternal haplotype 1-0
fBA	frequency of maternal haplotype 0-1
fBB	frequency of maternal haplotype 0-0
theta	paternal recombination rate

Value

lik value of log likelihood at parameter estimates

makehap*Make list of imputed sire haplotypes***Description**

List of sire haplotypes is set up in the format required for hsrecombi. Sire haplotypes are imputed from progeny genotypes using R package hspase.

Usage

```
makehap(sireID, daughterSire, genotype.chr, nmin = 30)
```

Arguments

<code>sireID</code>	vector (LEN N) of IDs of all sires
<code>daughterSire</code>	vector (LEN n) of sire ID for each progeny
<code>genotype.chr</code>	matrix (DIM n x p) of progeny genotypes on a single chromosome with p SNPs
<code>nmin</code>	scalar, minimum required number of progeny for proper imputation, default 30

Value

list (LEN 2) of lists. For each sire:

`famID` list (LEN N) of vectors (LEN n.progeny) of progeny indices relating to lines in genotype matrix
`sireHap` list (LEN N) of matrices (DIM 2 x p) of sire haplotypes (0, 1) on investigated chromosome

References

Ferdosi, M., Kinghorn, B., van der Werf, J., Lee, S. & Gondro, C. (2014) hsphase: an R package for pedigree reconstruction, detection of recombination events, phasing and imputation of half-sib family groups BMC Bioinformatics 15:172. <https://CRAN.R-project.org/package=hsphase>

Examples

```
data(targetregion)
hap <- makehap(unique(daughterSire), daughterSire, genotype.chr)
```

makehaplist *Make list of sire haplotypes*

Description

List of sire haplotypes is set up in the format required for hsrecombi. Haplotypes (obtained by external software) are provided.

Usage

```
makehaplist(daughterSire, hapSire, nmin = 1)
```

Arguments

<code>daughterSire</code>	vector (LEN n) of sire ID for each progeny
<code>hapSire</code>	matrix (DIM 2N x p + 1) of sire haplotype at p SNPs; 2 lines per sire, 1. column contains sire ID
<code>nmin</code>	scalar, minimum number of progeny required, default 1

Value

hap list (LEN 2) of lists. For each sire:

famID list (LEN N) of vectors (LEN n.progeny) of progeny indices relating to lines in genotype matrix

sireHap list (LEN N) of matrices (DIM 2 x p) of sire haplotypes (0, 1) on investigated chromosome

Examples

```
data(targetregion)
hap <- makehaplist(daughterSire, hapSire)
```

makehappm

Make list of imputed haplotypes and recombination rate

Description

List of sire haplotypes is set up in the format required for hsrecombi. Sire haplotypes are imputed from progeny genotypes using R package hspphase. Furthermore, recombination rate estimates between adjacent SNPs from hspphase are reported.

Usage

```
makehappm(sireID, daughterSire, genotype.chr, nmin = 30)
```

Arguments

sireID	vector (LEN N) of IDs of all sires
daughterSire	vector (LEN n) of sire ID for each progeny
genotype.chr	matrix (DIM n x p) of progeny genotypes on a single chromosome with p SNPs
nmin	scalar, minimum number of progeny required, default 1

Value

hap list (LEN 2) of lists. For each sire:

famID list (LEN N) of vectors (LEN n.progeny) of progeny indices relating to lines in genotype matrix

sireHap list (LEN N) of matrices (DIM 2 x p) of sire haplotypes (0, 1) on investigated chromosome

probRec vector (LEN p - 1) of proportion of recombinant progeny over all families between adjacent SNPs

numberRec list (LEN N) of vectors (LEN n.progeny) of number of recombination events per animal

References

Ferdosi, M., Kinghorn, B., van der Werf, J., Lee, S. & Gondro, C. (2014) hsphase: an R package for pedigree reconstruction, detection of recombination events, phasing and imputation of half-sib family groups BMC Bioinformatics 15:172. <https://CRAN.R-project.org/package=hsphase>

Examples

```
data(targetregion)
hap <- makehappm(unique(daughterSire), daughterSire, genotype.chr)
```

map.chr

targetregion: physical map

Description

SNP marker map in target region on chromosome BTA1 according to ARS-UCD1.2

Usage

map.chr

Arguments

map.chr	data frame
SNP	SNP index
Chr	chromosome of SNP
locus_bp	physical position of SNP in bp
locus_Mb	physical position of SNP in Mbp
markername	official SNP name

Format

An object of class `data.frame` with 300 rows and 5 columns.

rao*System of genetic mapping functions*

Description

Calculation of genetic distances from recombination rates given a mixing parameter

Usage

```
rao(p, x)
```

Arguments

p	mixing parameter (see details); $0 \leq p \leq 1$
x	vector of recombination rates

Details

Mixing parameter $p=0$ would match to Morgan, $p=0.25$ to Carter, $p=0.5$ to Kosambi and $p=1$ to Haldane map function

Value

vector of genetic positions in Morgan units

References

Rao, D.C., Morton, N.E., Lindsten, J., Hulten, M. & Yee, S (1977) A mapping function for man. Human Heredity 27: 99-104. doi: [10.1159/000152856](https://doi.org/10.1159/000152856)

Examples

```
rao(0.25, seq(0, 0.5, 0.01))
```

startvalue*Start value for maternal allele and haplotype frequencies*

Description

Determine default start values for Expectation Maximisation (EM) algorithm that is used to estimate paternal recombination rate and maternal haplotype frequencies

Usage

```
startvalue(Fam1, Fam2, Dd = 0, prec = 1e-06)
```

Arguments

Fam1	matrix (DIM n.progeny x 2) of progeny genotypes of genomic family with coupling phase sires (1) at SNP pair
Fam2	matrix (DIM n.progeny x 2) of progeny genotypes of genomic family with repulsion phase sires (2) at SNP pair
Dd	maternal LD, default 0
prec	minimum accepted start value for fAA, fAB, fBA; default 1e-6

Value

list (LEN 8)	
fAA.start	frequency of maternal haplotype 1-1
fAB.start	frequency of maternal haplotype 1-0
fBA.start	frequency of maternal haplotype 0-1
p1	estimate of maternal allele frequency (allele 1) when sire is heterozygous at SNP1
p2	estimate of maternal allele frequency (allele 1) when sire is heterozygous at SNP2
L1	lower bound of maternal LD
L2	upper bound for maternal LD
critical	0 if parameter estimates are unique; 1 if parameter estimates at both solutions are valid

Examples

```
n1 <- 100
n2 <- 20
G1 <- matrix(ncol = 2, nrow = n1, sample(c(0:2), replace = TRUE,
  size = 2 * n1))
G2 <- matrix(ncol = 2, nrow = n2, sample(c(0:2), replace = TRUE,
  size = 2 * n2))
startvalue(G1, G2)
```

targetregion	<i>Description of the targetregion data set</i>
--------------	---

Description

The data set contains sire haplotypes, assignment of progeny to sire, progeny genotypes and physical map information in a target region

The raw data can be downloaded at the source given below. Then, executing the following R code leads to the data provided in `targetregion.RData`.

hapSire	matrix of sire haplotypes of each sire; 2 lines per sire; 1. column contains sireID
daughterSire	vector of sire ID for each progeny
genotype.chr	matrix of progeny genotypes
map.chr	SNP marker map in target region

Source

The data are available at RADAR doi: [10.22000/280](https://doi.org/10.22000/280)

Examples

```
## Not run:  
# download data from RADAR (requires about 1.4 GB)  
url <- "https://www.radar-service.eu/radar-backend/archives/fqSPQoIvjt0GJlav/versions/1/content"  
curl_download(url = url, 'tmp.tar')  
untar('tmp.tar')  
file.remove('tmp.tar')  
path <- '10.22000-280/data/dataset'  
## list of haplotypes of sires for each chromosome  
load(file.path(path, 'sire_haplotypes.RData'))  
## assign progeny to sire  
daughterSire <- read.table(file.path(path, 'assign_to_family.txt'))[, 1]  
## progeny genotypes  
X <- as.matrix(read.table(file.path(path, 'XFam-ARS.txt')))  
## physical and approximated genetic map  
map <- read.table(file.path(path, 'map50K_ARS_reordered.txt'), header = T)  
## select target region  
chr <- 1  
window <- 301:600  
## map information of target region  
map.chr <- map[map$Chr == chr, ][window, 1:5]  
## matrix of sire haplotypes in target region  
hapSire <- rlist::list.rbind(haps[[chr]])  
sireID <- 1:length(unique(daughterSire))  
hapSire <- cbind(rep(sireID, each = 2), hapSire[, window])  
## matrix of progeny genotypes  
genotype.chr <- X[, map.chr$SNP]  
  
## End(Not run)
```

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