Package: ibdsim2 (via r-universe)

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Type Package

Title Simulation of Chromosomal Regions Shared by Family Members

Version 2.1.2

Description Simulation of segments shared identical-by-descent (IBD) by pedigree members. Using sex specific recombination rates along the human genome (Halldorsson et al. (2019) [<doi:10.1126/science.aau1043>](https://doi.org/10.1126/science.aau1043)), phased chromosomes are simulated for all pedigree members. Applications include calculation of realised relatedness coefficients and IBD segment distributions. 'ibdsim2' is part of the 'pedsuite' collection of packages for pedigree analysis. A detailed presentation of the 'pedsuite', including a separate chapter on 'ibdsim2', is available in the book 'Pedigree analysis in R' (Vigeland, 2021, ISBN:9780128244302). A 'Shiny' app for visualising and comparing IBD distributions is available at <<https://magnusdv.shinyapps.io/ibdsim2-shiny/>>.

License GPL-3

URL <https://github.com/magnusdv/ibdsim2>,

<https://magnusdv.github.io/pedsuite/>,

<https://magnusdv.shinyapps.io/ibdsim2-shiny/>

Depends R ($>= 4.1.0$), pedtools ($>= 2.7.1$)

Imports Rcpp, ggplot2, glue, ribd $(>= 1.6.1)$,

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Repository https://magnusdv.r-universe.dev

RemoteUrl https://github.com/magnusdv/ibdsim2

RemoteRef HEAD

RemoteSha c4c30ab9544770fb809e73a869d786a207997729

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Contents

convertPos *Conversion of genetic map positions*

Description

Convert between physical position (in megabases) and genetic position (centiMorgan) given a chromosome map. Linear extrapolation is used to convert positions between map points.

Usage

```
convertPos(
 chrom = NULL,
 Mb = NULL,cM = NULL,map = "decode19",sex = c("average", "male", "female")
)
```
Arguments

customMap 3

Value

A vector of the same length as the input.

Examples

```
# Chromosome 1 of the built-in recombination map
map = loadMap(chrom = 1)[[1]]
head(map$male)
# Conversion Mb -> cM
phys = 1:5gen = convertPos(Mb = phys, map = map, sex = "male")gen
# Convert back (note the first position, which was outside of map)
convertPos(cM = gen, map = map, sex = "male")
```


customMap *Custom recombination map*

Description

Create custom recombination maps for use in [ibdsim\(\)](#page-10-1).

Usage

customMap(x)

Arguments

x A data frame or matrix. See details for format specifications.

Details

The column names of x must include either

• chrom, mb and cm (sex-averaged map)

or

• chrom, mb, male and female (sex-specific map)

Upper-case letters are allowed in these names. The mb column should contain physical positions in megabases, while cm, male, female give the corresponding genetic position in centiMorgans.

Value

An object of class genomeMap.

See Also

[uniformMap\(\)](#page-26-1), [loadMap\(\)](#page-17-1)

Examples

```
# A map including two chromosomes.
df1 = data frame(chrom = c(1, 1, 2, 2),mb = c(0, 2, 0, 5),cm = c(0, 3, 0, 6)map1 = customMap(df1)
map1
# Use columns "male" and "female" to make sex specific maps
df2 = data frame(chrom = c(1, 1, 2, 2),mb = c(0, 2, 0, 5),male = c(0, 3, 0, 6),female = c(0, 4, 0, 7)map2 = customMap(df2)
map2
```
estimateCoeffs *Estimation of one- and two-locus relatedness coefficients*

Description

Estimate by simulation various relatedness coefficients, and two-locus versions of the same coefficients, for a given recombination rate. The current implementation covers inbreeding coefficients, kinship coefficients, IBD (kappa) coefficients between noninbred individuals, and condensed identity coefficients. These functions are primarily meant as tools for validating exact algorithms, e.g., as implemented in the ribd package.

Usage

```
estimateInbreeding(x, id, Nsim, Xchrom = FALSE, verbose = FALSE, ...)
estimateTwoLocusInbreeding(
  x,
  id,
  rho = NULL,
  cM = NULL,
 Nsim,
  Xchrom = FALSE,
  verbose = FALSE,
```

```
...
\mathcal{L}estimateKinship(x, ids, Nsim, Xchrom = FALSE, verbose = FALSE, ...)
estimateTwoLocusKinship(
 x,
  ids,
 rho = NULL,
 cM = NULL,Nsim,
 Xchrom = FALSE,
 verbose = FALSE,
  ...
\mathcal{L}estimateKappa(x, ids, Nsim, Xchrom = FALSE, verbose = FALSE, ...)
estimateTwoLocusKappa(
 x,
 ids,
 rho = NULL,
 cM = NULL,Nsim,
 Xchrom = FALSE,
 verbose = FALSE,
  ...
\mathcal{L}estimateIdentity(x, ids, Nsim, Xchrom = FALSE, verbose = FALSE, ...)
estimateTwoLocusIdentity(
 x,
 ids,
 rho = NULL,
 cM = NULL,Nsim,
 Xchrom = FALSE,
  verbose = FALSE,
  ...
\mathcal{L}
```
Arguments

Details

In the following, let L1 and L2 denote two arbitrary autosomal loci with recombination rate ρ , and let A and B be members of the pedigree x.

The *two-locus inbreeding coefficient* $f_2(\rho)$ of A is defined as the probability that A is autozygous at both L1 and L2 simultaneously.

The *two-locus kinship coefficient* $\phi_2(\rho)$ of A and B is defined as the probability that a random gamete emitted from A, and a random gamete emitted from B, contain IBD alleles at both L1 and L2.

The *two-locus kappa coefficient* $\kappa_{ij}(\rho)$, for $i, j = 0, 1, 2$, of noninbred A and B, is the probability that A and B share exactly i alleles IBD at L1, and exactly j alleles IBD at L2.

The *two-locus identity coefficient* Δ_{ij} , $i, j = 1, ..., 9$ is defined for any (possibly inbred) A and B, as the probability that A and B are in identity state i at L1, and state j at L2. This uses the conventional ordering of the nine condensed identity states. For details, see for instance the [GitHub](https://github.com/magnusdv/ribd) [page of the](https://github.com/magnusdv/ribd) ribd package.

Value

estimateInbreeding(): a single probability.

estimateTwoLocusInbreeding(): a single probability.

estimateKappa(): a numeric vector of length 3, with the estimated κ coefficients.

estimateTwoLocusKappa(): a symmetric, numerical 3*3 matrix, with the estimated values of κ_{ij} , for $i, j = 0, 1, 2$.

estimateIdentity(): a numeric vector of length 9, with the estimated identity coefficients.

estimateTwoLocusIdentity(): a symmetric, numerical 9*9 matrix, with the estimated values of Δ_{ij} , for $i, j = 1, ..., 9$.

```
############################
### Two-locus inbreeding ###
############################
x = \text{cousinPed}(0, \text{ child} = \text{TRUE})rho = 0.25Nsim = 10 # Increase!
estimateTwoLocusInbreeding(x, id = 5, rho = rho, Nsim = Nsim, seed = 123)
```

```
extractIds 7
```

```
# Exact:
right::twoLocusInbreeding(x, id = 5, rho = rho)########################################
### Two-locus kappa: ###
### Grandparent vs half sib vs uncle ###
########################################
# These are indistinguishable with unlinked loci, see e.g.
# pages 182-183 in Egeland, Kling and Mostad (2016).
# In the following, each simulation approximation is followed
# by its exact counterpart.
rho = 0.25; R = .5 * (rho^2 + (1-rho)^2)
Nsim = 10 # Should be increased to at least 10000
# Grandparent/grandchild
G = linearPed(2); G.ids = c(1,5); # plot(G, hatched = G.ids)estimateTwoLocusKappa(G, G.ids, rho = rho, Nsim = Nsim, seed = 123)[2,2]
.5*(1-rho) # exact
# Half sibs
H = halfSibPed(); H.ids = c(4,5); # plot(H, hatched = H.ids)
estimateTwoLocusKappa(H, H.ids, rho = rho, Nsim = Nsim, seed = 123)[2,2]
R # exact
# Uncle
U = avuncularPed(); U.ids = c(3,6); # plot(U, 4) hatched = U.ids)estimateTwoLocusKappa(U, U.ids, rho = rho, Nsim = Nsim, seed = 123)[2,2]
(1-rho) * R + rho/4 # exact# Exact calculations by ribd:
# ribd::twoLocusIBD(G, G.ids, rho = rho, coefs = "k11")
# ribd::twoLocusIBD(H, H.ids, rho = rho, coefs = "k11")
# ribd::twoLocusIBD(U, U.ids, rho = rho, coefs = "k11")
##########################
### Two-locus Jacquard ###
##########################
x = fullSibMatrix(1)rho = 0.25Nsim = 10 # (increase to at least 10000)
estimateTwoLocusIdentity(x, ids = 5:6, rho = rho, Nsim = Nsim, seed = 123)
# Exact by ribd:
# ribd::twoLocusIdentity(x, ids = 5:6, rho = rho)
```
Description

Extract ID labels from simulation output

Usage

```
extractIds(sim)
```
Arguments

sim Output from [ibdsim\(\)](#page-10-1)

Value

A character vector

Examples

```
s = ibdsim(nuclearPed(2), N=1, ids = 3:4)stopifnot(all(extractIds(s) == c("3", "4")))
```
findPattern *Find specific IBD patterns*

Description

Find segments satisfying a particular pattern of IBD sharing, in a list of IBD simulations.

Usage

```
findPattern(sims, pattern, merge = TRUE, cutoff = 0, unit = "mb")
```
Arguments

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Details

For each simulation, this function extracts the subset of rows satisfying the allele sharing specified by pattern. That is, segments where, for some allele,

- all of pattern\$autozygous are autozygous
- all of pattern\$heterozygous have exactly one copy
- all of pattern\$carriers have at least one copy
- none of pattern\$noncarriers carry the allele.

Value

A matrix (if sims is a single genomeSim object), or a list of matrices.

See Also

[segmentStats\(\)](#page-25-1)

Examples

```
x = nuclearPed(3)s = ibdsim(x, N = 1, map = uniformMap(M = 1), seed = 1729)# Segments where some allele is shared by 3 and 4, but not 5
pattern = list(carriers = 3:4, noncarriers = 5)findPattern(s, pattern)
# Exclude segments less than 7 cM
findPattern(s, pattern, cutoff = 7)
# Visual confirmation:
haploDraw(x, s)
```
haploDraw *Draw haplotypes onto a pedigree plot*

Description

Visualise the IBD pattern of a single chromosome, by drawing haplotypes onto the pedigree.

Usage

```
haploDraw(
 x,
  ibd,
 chrom = NULL,
  ids = NULL,unit = "mb",
```
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```
L = NULL,pos = 1,
 cols = NULL,
 height = 4,
 width = 0.75,
 sep = 0.75,dist = 1,
 keep.par = FALSE,
  ...
\mathcal{L}
```
Arguments

Value

None.

Examples

############################### # Example 1: A family quartet #

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```
###############################
x = nuclearPed(2)map = uniformMap(M = 1)s = ibdsim(x, map = map, seed = 4276)haploDraw(x, s)
# Custom colours and placements
haploDraw(x, s, cols = c(3,7,2,4), pos = c(2,4,2,4))
# Standard plot options apply
haploDraw(x, s, margins = 3, cex = 1.5, title = "Full sibs")
###########################
# Example 2: Autozygosity #
###########################
x = \text{halfCousinPed}(0, \text{ child} = \text{TRUE})map = uniformMap(M = 1)s = ibdsim(x, map = map, skipRecomb = c(1,3), seed = 2)# Only include relevant individuals (skip 1 and 3)
haploDraw(x, s, ids = c(2,4,5,6), pos = c(1,2,4,4))
###############################
# Example 3: X-chromosomal sims
###############################
x = nuclearPed(2, sex = 2:1)s = ibdsim(x, N = 1, map = uniformMap(M = 1, chrom = "X"), seed = 123)
haploDraw(x, s)
```
ibdsim *IBD simulation*

Description

This is the main function of the package, simulating the recombination process in each meioses of a pedigree. The output summarises the IBD segments between all or a subset of individuals.

Usage

ibdsim(x, $N = 1$, 12 ibdsim and the state of the state of

```
ids = labels(x),
map = "decode",model = c("chi", "haldane"),skipRecomb = NULL,
simplify1 = TRUE,
seed = NULL,
verbose = TRUE
```
λ

Arguments

Details

Each simulation starts by unique alleles (labelled 1, 2, ...) being distributed to the pedigree founders. In each meiosis, homologue chromosomes are made to recombine according to the value of model:

- model = "haldane": In this model, crossover events are modelled as a Poisson process along each chromosome.
- model = "chi" (default): This uses a renewal process along the four-strand bundle, with waiting times following a chi square distribution.

Recombination rates along each chromosome are determined by the map parameter. The default value ("decode19") loads a thinned version of the recombination map of the human genome published by Halldorsson et al (2019).

In many applications, the fine-scale default map is not necessary, and should be replaced by simpler maps with constant recombination rates. See [uniformMap\(\)](#page-26-1) and [loadMap\(\)](#page-17-1) for ways to produce such maps.

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Value

A list of N objects of class genomeSim.

If $N = 1$ the outer list layer is removed by default, which is typically desired in interactive use (especially when piping). To enforce a list output, add simplify1 = FALSE.

A genomeSim object is essentially a numerical matrix describing the allele flow through the pedigree in a single simulated. Each row corresponds to a chromosomal segment. The first 3 columns (chrom, startMB, endMB) describe the physical location of the segment. Next, the genetic coordinates (startCM, endCM), which are computed from map by averaging the male and female values. Then follow the allele columns, two for each individual in ids, suffixed by ":p" and ":m" signifying the paternal and maternal alleles, respectively.

If ids has length 1, a column named Aut is added, whose entries are 1 for autozygous segments and 0 otherwise.

If ids has length 2, two columns are added:

- IBD : The IBD status of each segment (= number of alleles shared identical by descent). For a given segment, the IBD status is either 0, 1, 2 or NA. If either individual is autozygous in a segment, the IBD status is reported as NA. With inbred individuals the Sigma column (see below) is more informative than the IBD column.
- Sigma : The condensed identity ("Jacquard") state of each segment, given as an integer in the range 1-9. The numbers correspond to the standard ordering of the condensed states. In particular, for non-inbred individuals, the states 9, 8, 7 correspond to IBD status 0, 1, 2 respectively.

References

Halldorsson et al. *Characterizing mutagenic effects of recombination through a sequence-level genetic map.* Science 363, no. 6425 (2019).

```
### Example 1: Half siblings ###
hs = halfSibPed()
sim = ibdsim(hs, map = uniformMap(M = 1), seed = 10)sim
# Plot haplotypes
haploDraw(hs, sim)
#' ### Example 2: Full sib mating ###
x = fullSibMating(1)sim = ibdsim(x, ids = 5:6, map = uniformMap(M = 10), seed = 1)head(sim)
# All 9 identity states are present
stopifnot(setequal(sim[, 'Sigma'], 1:9))
```


Description

Show chromosomal segments in a diploid karyogram

Usage

```
karyoDiploid(
 paternal,
 maternal,
  chrom = 1:22,
  col = c(paternal = "lightblue", material = "orange"),alpha = 1,
 bgcol = "gray95",
  title = NULL
)
```
Arguments

Value

The plot object is returned invisibly, so that additional ggplot layers may be added if needed.

```
## Not run:
pat = data.frame(chrom = c(1,4,5,5,10,10), start = c(100,50,20,80,10,80),
                 end = c(120,100,25,100,70,120))
mat = data.frame(chrom = c(2,4,5,5,10), start = c(80,50,10,80,50),
```
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```
end = c(120,100,35,100,120))
karyoDiploid(pat, mat)
## End(Not run)
```
karyogram2 *Karyogram plots*

Description

Functions for visualising IBD segments in karyograms. The karyogram1() and karyogram2() functions produces karyograms illustrating the output of [ibdsim\(\)](#page-10-1) for one or two specified individuals. The actual plotting is done by functions karyoHaploid() and karyoDiploid().

Usage

 $karyogram2(sim, ids = NULL, verbose = TRUE, ...)$

Arguments

Value

A plot object returned invisibly.

```
x = quadHalfFirstCousins()
s = ibdsim(x, seed = 1729)# karyogram2(s, ids = leaves(x), title = "QHFC")
```


Description

Show chromosomal segments in a haploid karyogram

Usage

```
karyoHaploid(
 segments,
 chrom = 1:22,
 colBy = NULL,
 col = NULL,separate = TRUE,
 alpha = 1,
  bgcol = "gray92",title = NULL,
 legendTitle = NULL,
 base_size = 16
)
```
Arguments

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Value

The plot object is returned invisibly, so that additional ggplot layers may be added if needed.

Examples

```
## Not run:
segs = data. frame(chrom = c(1, 4, 5, 5, 10, 10),
                      start = c(100,50,20,80,10,50),
                      end = c(120,100,25,100,70,120),
                      IBD = c("paternal","maternal"))
\text{cols} = \text{c}(\text{patternal} = \text{"blue", material} = \text{"red"})karyoHaploid(segs, col = "cyan")
karyoHaploid(segs, colBy = "IBD", col = cols)
karyoHaploid(segs, col = "cyan")<br>karyoHaploid(segs, colBy = "IBD", col =<br># Note difference if `separate = FALSE`
karyoHaploid(segs, colBy = "IBD", col = cols, separate = FALSE)
# Reduce alpha to see the overlaps:
karyoHaploid(segs, colBy = "IBD", col = cols, separate = FALSE, alpha = 0.7)
```
End(Not run)

launchApp *Launch the ibdsim2 app*

Description

This launches the Shiny app for simulating IBD segment distributions.

Usage

launchApp()

Value

No return value, called for side effects.

Examples

Not run: launchApp()

End(Not run)

Description

This function loads one of the built-in genetic maps. Currently, the only option is a detailed human recombination map, based on the publication by Halldorsson et al. (2019).

Usage

```
loadMap(map = "decode19", chrom = 1:22, uniform = FALSE, sexAverage = FALSE)
```
Arguments

Details

For reasons of speed and efficiency, the map published by map Halldorsson et al. (2019) has been thinned down to around 60 000 data points.

By setting uniform = TRUE, a uniform version of the map is returned, in which each chromosome has the same genetic lengths as in the original, but with constant recombination rates. This gives much faster simulations and may be preferable in some applications.

Value

An object of class genomeMap, which is a list of chromMap objects. A chromMap is a list of two matrices, named "male" and "female", with various attributes:

- physStart: The first physical position (Mb) on the chromosome covered by the map
- physEnd: The last physical position (Mb) on the chromosome covered by the map
- physRange: The physical map length (Mb), equal to physEnd physStart
- mapLen: A vector of length 2, containing the centiMorgan lengths of the male and female strands
- chrom: A chromosome label
- Xchrom: A logical. This is checked by ibdsim() and other function, to select mode of inheritance

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References

Halldorsson et al. *Characterizing mutagenic effects of recombination through a sequence-level genetic map.* Science 363, no. 6425 (2019).

See Also

[uniformMap\(\)](#page-26-1), [customMap\(\)](#page-2-1)

Examples

```
# By default, the complete map of all 22 autosomes is returned
loadMap()
```

```
# Uniform version
m = loadMap(uniform = TRUE)
m
# Check chromosome 1
m1 = m[[1]]m1
m1$male
m1$female
```
The X chromosome loadMap(chrom = "X")[[1]]

maplengths *Physical and genetic map lengths*

Description

Utility functions for extracting the physical or genetic length of chromosome maps and genome maps.

Usage

```
mapLen(x, \ldots)## S3 method for class 'chromMap'
mapLen(x, sex = c("male", "female"), ...)## S3 method for class 'genomeMap'
mapLen(x, sex = c("male", "female"), ...)physRange(x, ...)
## S3 method for class 'chromMap'
```
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```
physRange(x, ...)
```
S3 method for class 'genomeMap' physRange(x, ...)

Arguments

Value

mapLen() returns a numeric of the same length as sex, with the genetic length(s) in centiMorgan.

physRange() returns the physical length (in Mb) of the chromosome/genome covered by the map. For example, for a chromosome map starting at 2 Mb and ending at 8 Mb, the output is 6.

See Also

[loadMap\(\)](#page-17-1), [uniformMap\(\)](#page-26-1)

Examples

```
m = loadMap(chrom = 1:2)
m
# Applied to `genomeMap` object:
physRange(m)
mapLen(m)
# Applied to `chromMap` object:
```
plotSegmentDistribution

physRange(m[[1]]) mapLen(m[[1]])

Scatter plots of IBD segment distributions

Description

Visualise and compare count/length distributions of IBD segments. Two types are currently implemented: Segments of autozygosity (for a single person) and segments with (pairwise) IBD state 1.

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Usage

```
plotSegmentDistribution(
```

```
...,
type = c("autozygosity", "ibd1"),
ids = NULL,unit = "cm",labels = NULL,
col = NULL,shape = 1,
alpha = 1,
ellipses = TRUE,
title = NULL,
xlab = NULL,
ylab = NULL,
legendInside = TRUE
```
Arguments

)

Details

This function takes as input one or several complete outputs from the [ibdsim\(\)](#page-10-1), and produces a scatter plot of the number and average length of IBD segments from each.

Contour curves are added to plot, corresponding to the theoretical/pedigree-based values: either inbreeding coefficients (if type = "autozygosity") or κ_1 (if type = "ibd1").

```
# Simulation parameters used in the below examples.
map = uniformMap(M = 10) # recombination map
N = 5 # number of sims
# For more realistic results, replace with e.g.:
# map = loadMap("decode19")
# N = 1000
#################################################################
# EXAMPLE 1
# Comparison of IBD segment distributions
# between paternal and maternal half siblings.
#################################################################
# Define the pedigrees
xPat = halfSibPed()
xMat = swapSex(xPat, 1)simPat = ibdsim(xPat, N = N, map = map)simMat = ibdsim(xMat, N = N, map = map)# By default, the IBD segments of the "leaves" are computed and plotted
plotSegmentDistribution(simPat, simMat, type = "ibd1", ids = 4:5,
                       labels = c("HSpat", "HSmat"))
#################################################################
# EXAMPLE 2
# Half siblings vs half uncle vs grandparent/grandchild
#################################################################
# Only one pedigree needed here
x = addSon(halfSibPed(), 5)s = ibdsim(x, N = N, map = map)# Indicate the pairs explicitly this time.
ids = list(GR = c(2,7), HS = 4:5, HU = c(4,7))
# List names are used as labels in the plot
plotSegmentDistribution(s, type = "ibd1", ids = ids, shape = 1:3)
#################################################################
# EXAMPLE 3
# Comparison of autozygosity distributions in various individuals
# with the same expected inbreeding coefficient (f = 1/8)
#################################################################
G = linearPed(2) |> swapSex(5) |> addSon(c(1,5)) # grandfath/granddaughter
HSpat = halfSibPed(sex2 = 2) |> addSon(4:5) # paternal half sibs
```
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```
HSmat = swapSex(HSpat, 2) # maternal half sibs
QHFC = quadHalfFirstCousins() # quad half first cousins
QHFC = addSon(QHFC, 9:10)
peds = list(G = G, HSpat = HSpat, HSmat = HSmat, QHFC = QHFC)
plotPedList(peds, newdev = TRUE)
dev.off()
# Simulations
s = lapply(peds, function(p)
 ibdsim(p, N = N, ids = leaves(p), verbose = FALSE, map = map))# Plot distributions
plotSegmentDistribution(s, type = "autoz", title = "Autozygous segments")
```
profileSimIBD *Simulate markers conditional on a given IBD pattern*

Description

This function simulates genotypes for a set of markers conditional on a specific underlying IBD pattern (typically produced with [ibdsim\(\)](#page-10-1)).

Usage

```
profileSimIBD(
  x,
  ibdpattern,
  ids = NULL,
  markers = NULL,
  seed = NULL,
  verbose = TRUE
```
)

Arguments

Details

It should be noted that the only *random* part of this function is the sampling of founder alleles for each marker. Given those, all other genotypes in the pedigree are determined by the underlying IBD pattern.

A copy of x where marker genotypes have been simulated conditional on ibdpattern.

See Also

[ibdsim\(\)](#page-10-1), forrel::profileSim().

```
# Brother-sister pedigree
ped = nuclearPed(2, sex = 1:2)# Alleles
als = letters[1:10]
### Autosomal simulation
x = ped |>addMarket(alleles = als, chrom = 1, posMb = 20) |>
  addMarker(alleles = als, chrom = 1, posMb = 50) |>
  addMarker(alleles = als, chrom = 1, posMb = 70)
# Simulate the underlying IBD pattern in the pedigree
sim = ibdsim(x, map = uniformMap(M = 1, chrom = 1), seed = 123)# Simulate genotypes for the sibs conditional on the given IBD pattern
profilesimIBD(x, sim, ids = 3:4, seed = 123)# With a different seed
profileSimIBD(x, sim, ids = 3:4, seed = 124)### X chromosomal simulation
y = ped |>addMarker(alleles = als, chrom = "X", posMb = 1) |>
  addMarker(alleles = als, chrom = "X", posMb = 50) |>
  addMarker(alleles = als, chrom = "X", posMb = 100)
simy = ibdsim(y, map = loadMap("decode19", chrom = 23), seed = 11)
profileSimIBD(y, simy, seed = 12)
```
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Description

Compute the realised values of various pedigree coefficients, from simulated data. The current implementation covers inbreeding coefficients for single pedigree members, and kinship, kappa and condensed identity coefficients for pairwise relationships.

Usage

```
realisedInbreeding(sims, id = NULL, unit = "cm")
realisedKinship(sims, ids = NULL, unit = "cm")
realisedKappa(sims, ids = NULL, unit = "cm")
realisedIdentity(sims, ids = NULL, unit = "cm")
```
Arguments

Details

The inbreeding coefficient f of a pedigree member is defined as the probability of autozygosity (homozygous for alleles that are identical by descent) in a random autosomal locus. Equivalently, the inbreeding coefficient is the *expected* autozygous proportion of the autosomal chromosomes.

The *realised* inbreeding coefficient f_R in a given individual is the actual fraction of the autosomes covered by autozygous segments. Because of the stochastic nature of meiotic recombination, this may deviate substantially from the pedigree-based expectation.

Similarly, the pedigree-based IBD coefficients $\kappa_0, \kappa_1, \kappa_2$ of noninbred pairs of individuals have realised counterparts. For any given pair of individuals we define k_i to be the actual fraction of the autosome where the individuals share exactly i alleles IBD, where $i = 0, 1, 2$.

Finally, we can do the same thing for each of the nine condensed identity coefficients of Jacquard. For each $i = 1, ..., 9$ we define D_i the be the fraction of the autosome where a given pair of individuals are in identity state i . This uses the conventional ordering of the nine condensed identity states; see for instance the ribd [GitHub page.](https://github.com/magnusdv/ribd)

Examples

```
# Realised IBD coefficients between full siblings
x = nuclearPed(2)s = ibdsim(x, N = 2) # increase NrealisedKappa(s, ids = 3:4)
```
###########

Realised inbreeding coefficients, child of first cousins

```
x = \text{cousinPed}(1, \text{ child} = \text{TRUE})s = ibdsim(x, N = 2) # increase N
realisedInbreeding(s, id = 9)
# Same data: realised kinship coefficients between the parents
realisedKinship(s,ids = parents(x, 9))###########
# Realised identity coefficients after full sib mating
x = fullSibMating(1)
s = ibdsim(x, N = 2) # increase N
realisedIdentity(s, ids = 5:6)
```
segmentStats *Summary statistics for identified segments*

Description

Compute summary statistics for segments identified by [findPattern\(\)](#page-7-1).

Usage

```
segmentStats(
  x,
 quantiles = c(0.025, 0.5, 0.975),
 returnAll = FALSE,unit = "mb"\lambda
```
Arguments

Value

A list containing a data frame perSim, a matrix summary and (if returnAll is TRUE) a vector allSegs.

Variables used in the output:

• Count: The total number of segments in a simulation

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- Total: The total sum of the segment lengths in a simulation
- Average: The average segment lengths in a simulation
- Shortest: The length of the shortest segment in a simulation
- Longest: The length of the longest segment in a simulation
- Overall (only in summary): A summary of all segments from all simulations

See Also

[findPattern\(\)](#page-7-1)

Examples

```
x = nuclearPed(3)sims = ibdsim(x, N = 2, map = uniformMap(M = 2), model = "haldane", seed = 1729)
# Segments where all siblings carry the same allele
segs = findPattern(sims, pattern = list(carriers = 3:5))
# Summarise
segmentStats(segs, unit = "mb")
# The unit does not matter in this case (since the map is trivial)
segmentStats(segs, unit = "cm")
```
uniformMap *Uniform recombination maps*

Description

Create a uniform recombination map of a given length.

Usage

```
uniformMap(Mb = NULL, cM = NULL, M = NULL, cmPerMb = 1, chrom = 1)
```
Arguments

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Value

An object of class chromMap. See [loadMap\(\)](#page-17-1) for details.

See Also

[loadMap\(\)](#page-17-1), [customMap\(\)](#page-2-1)

Examples

```
m = uniformMap(Mb = 1, cM = 2:3)m
m$male
m$female
mx = uniformMap(M = 1, chrom = "X")mx
mx$male
mx$female
```
zeroIBD *Probability of zero IBD*

Description

Estimate the probability of no IBD sharing in a pairwise relationship.

Usage

```
zeroIBD(sims,ids = NULL, threshold = 0, unit = "cm")
```
Arguments

Value

A list with the following two entries:

- zeroprob: The fraction of sims in which ids have no IBD sharing
- stErr: The standard error of zeroprob

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```
###
# The following example computes the probability of
# no IBD sharing between a pair of fourth cousins.
# We also show how the probability is affected by
# truncation, i.e., ignoring short segments.
###
# Define the pedigree
x = \text{cousinPed}(4)cous = leaves(x)# Simulate (increase N!)
s = ibdsim(x, N = 10)# Probability of zero ibd segments. (By default all segs are used)
zeroIBD(s, ids = cous)
# Re-compute with nonzero threshold
zeroIBD(s, is = cous, threshold = 1, unit = "cm")zeroIBD(s, is = cous, threshold = 1, unit = "mb")
```
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