Package ‘ribd’

February 3, 2020

Type  Package
Title  Pedigree-based Relatedness Coefficients
Version  1.0.1
Description  Recursive algorithms for computing various relatedness coefficients, including pairwise kinship, kappa and identity coefficients. Both autosomal and X-linked coefficients are computed. Founders are allowed to be inbred. In addition to the standard pairwise coefficients, ribd also computes a range of lesser-known coefficients, including generalised kinship coefficients (Karigl (1981) <doi:10.1111/j.1469-1809.1981.tb00341.x>; Weeks and Lange (1988) <https:www.ncbi.nlm.nih.gov/pmc/articles/PMC1715269>), two-locus coefficients (Thompson (1988) <doi:10.1093/imammb/5.4.261>) and multi-person coefficients. This package is part of the ped suite, a collection of packages for pedigree analysis with 'pedtools' as the core package for creating and handling pedigree objects.
License  GPL-3
URL  https://github.com/magnusdv/ribd
Encoding  UTF-8
Language  en-GB
LazyData  true
Depends  R (>= 3.5.0), pedtools
Imports  slam, kinship2, glue
Suggests  identity, testthat
RoxygenNote  7.0.2
NeedsCompilation  no
Author  Magnus Dehli Vigeland [aut, cre]
        (<https://orcid.org/0000-0002-9134-4962>)
Maintainer  Magnus Dehli Vigeland <m.d.vigeland@medisin.uio.no>
Repository  CRAN
Date/Publication  2020-02-03 15:10:08 UTC
**R topics documented:**

- condensedIdentity .................................................. 2
- condensedIdentityX .................................................. 3
- external_coefs ......................................................... 5
- generalisedKinship ................................................... 6
- generalised_karigl ..................................................... 8
- inbreeding .............................................................. 9
- jicaque ................................................................. 11
- kappaIBD ............................................................... 11
- kinPattern ............................................................... 13
- kinship ................................................................. 14
- minimalPattern .......................................................... 15
- multiPersonIBD ........................................................... 15
- ribd ...................................................................... 17
- twoLocusIBD ............................................................... 17
- twoLocusIdentity ........................................................ 21
- twoLocusKinship ........................................................ 23
- twoLocusPlot .............................................................. 24

**Index**

- condensedIdentity 27

---

**condensedIdentity**  
**Condensed identity coefficients**

**Description**

Computes the 9 condensed identity coefficients of pairwise relationships in a pedigree. Founders of the pedigree may be inbred; use `pedtools::founderInbreeding()` to set this up.

**Usage**

```r
condensedIdentity(x, ids, sparse = NA, verbose = FALSE, checkAnswer = verbose)
```

**Arguments**

- `x`  
  A pedigree in the form of a `pedtools::ped` object

- `ids`  
  A character (or coercible to character) containing ID labels of two or more pedigree members.

- `sparse`  
  A positive integer, indicating the pedigree size limit for using sparse arrays (as implemented by the `slam` package) instead of ordinary arrays.

- `verbose`  
  A logical

- `checkAnswer`  
  A logical. If TRUE, and the `identity` package is installed, the result is checked against the output of `identity::identity.coefs()`. This option is ignored if any of the founders are inbred, or if `ids` has length greater than 2.
Details

The implementation is a modified version of Karigl’s recursive algorithm (1981).

Value

If `ids` has length 2: A vector of length 9, containing the condensed identity coefficients.
If `ids` has length > 2: A data frame with one row for each pair of individuals, and 11 columns. The first two columns contain the ID labels, and columns 3-11 contain the condensed identity coefficients.

References


See Also

kappa(), condensedIdentityX(), pedtools::founderInbreeding()

Examples

# One generation of full sib mating.
# (One of the simplest examples with all 9 coefficients nonzero.)
x = fullSibMating(1)
j1 = condensedIdentity(x, ids = 5:6)

stopifnot(all.equal(j1, c(2, 1,4, 1, 4, 1, 7, 10, 2)/32))

# Recalculate the coefficients when the founders are 100% inbred
founderInbreeding(x, 1:2) = 1
condensedIdentity(x, ids = 5:6)

condensedIdentityX  Identity coefficients on X

Description

Computes the X chromosomal condensed identity coefficients of a pairwise relationship.

Usage

condensedIdentityX(x, ids, sparse = NA, verbose = FALSE)
condensedIdentityX

Arguments

- **x**: A pedigree in the form of a `pedtools::ped` object.
- **ids**: A character (or coercible to character) containing ID labels of two or more pedigree members.
- **sparse**: A positive integer, indicating the pedigree size limit for using sparse arrays (as implemented by the `slam` package) instead of ordinary arrays.
- **verbose**: A logical

Details

The implementation is inspired by Karigl’s recursive algorithm (1981) for the autosomal case, modified to account for X-linked inheritance.

The X chromosomal pairwise identity states depend on the sexes of the two individuals. If both are female, the states are the same as in the autosomal case. When males are involved, the two individuals have less than 4 alleles, hence the states differ from the autosomal ones. However, to avoid drawing (and learning) new pictures we re-use the autosomal states by using the following simple rule: Replace any hemizygous male allele with a pair of autozygous alleles. In this way each X state corresponds to a unique autosomal state.

For simplicity the output always contains 9 coefficients, but with NA’s in the positions of undefined states (depending on the sex combination). The README file on the github home page of ribd has a table illustrating this.

Value

- If `ids` has length 2: A vector of length 9, containing the condensed identity coefficients. If any of the individuals are male, certain states are undefined, and the corresponding coefficients are NA. (See Details.)
- If `ids` has length > 2: A data frame with one row for each pair of individuals, and 11 columns. The first two columns contain the ID labels, and columns 3-11 contain the condensed identity coefficients.

See Also

- `kinshipX()`, `condensedIdentity()`, `pedtools::founderInbreeding()`

Examples

```r
x = fullSibMating(1)
x_sisters = swapSex(x, 5)
x_brothers = swapSex(x, 6)

condensedIdentityX(x, ids = 5:6)
condensedIdentityX(x_sisters, ids = 5:6)
condensedIdentityX(x_brothers, ids = 5:6)
```
external_coefs

---

**Relatedness coefficients by other programs**

---

**Description**

Wrappers for functions in other packages or external programs, computing various relatedness coefficients.

**Usage**

kinship2_kinship(x, ids = NULL)

kinship2_kinshipX(x, ids = NULL)

kinship2_inbreeding(x)

kinship2_inbreedingX(x)

idcoefs(x, ids)

idcoefs2(x, ids, verbose = FALSE, cleanup = TRUE)

**Arguments**

- **x**: A pedigree, in the form of a `pedtools::ped` object.
- **ids**: A integer vector of length 2.
- **verbose**: A logical, indicating if messages from IdCoefs should be printed.
- **cleanup**: A logical: If TRUE, the pedfile and sample file created for the IdCoefs run are deleted automatically.

**Details**

kinship2_inbreeding() and kinship2_kinship() are wrappers of `kinship2::kinship()` with the parameter `chrtype = "autosome"`.

Similarly, `kinship2_inbreedingX()` and `kinship2_kinshipX()` wrap `kinship2::kinship()` using `chrtype = "x"`.

`idcoefs()` wraps `identity::identity.coefs()`, which is an R interface for the C program IdCoefs written by Mark Abney (2009). The `identity.coefs()` function sometimes causes R to crash, hence I have provided an alternative wrapper, `idcoefs2`, which executes an external call to the original C program IdCoefs (version 2.1.1). For this to work, IdCoefs must be installed on the computer (see link in the References section below) and the executable placed in a folder included in the PATH variable. The `idcoefs2()` wrapper works by writing the necessary files to disk and calling IdCoefs via `system()`.
generalisedKinship

Value

For `kinship2_inbreeding()` and `kinship2_inbreedingX()`, a named numerical vector with the inbreeding coefficients and ID labels as names.

For `kinship2_kinship()` and `kinship2_kinshipX()`, either a single numeric (if ids is a pair of pedigree members) or the whole kinship matrix, with the ID labels as dimnames.

For `jaquard()` and `jaquard2()`, a numerical vector of length 9 (in the standard order of Jacquard’s condensed identity coefficients).

Author(s)

Magnus Dehli Vigeland

References


See Also

`kinship2::kinship()`, `identity::identity.coefs()`

Examples

```r
# A random pedigree with 2 founders and 5 matings
p = pedtools::randomPed(g = 5, founders = 2, seed = 111)

# Check that ribd agrees with kinship2 package
kinship_k2 = kinship2_kinship(p)
kinship_ribd = kinship(p)
stopifnot(identical(kinship_k2, kinship_ribd))

# Check on X also
kinshipX_k2 = kinship2_kinshipX(p)
kinshipX_ribd = kinshipX(p)
stopifnot(identical(kinshipX_k2, kinshipX_ribd))
```

---

generalisedKinship Generalised kinship coefficients

Description

Usage

generalisedKinship(x, pattern, mem = NULL, verbose = FALSE, debug = FALSE)

Arguments

x
A ped object.
pattern
A kinPattern object.
mem
An environment (for internal use).
verbose
A logical.
debug
A logical.

Value

A single probability.

References


Examples

x = nuclearPed(3)
kp = kinPattern(x, list(c(1,1,1)))
generalisedKinship(x, kp)

##### IBD coefficients via generalised kinship ####
#(Clearly not the simplest way; serves as a check)
IBD_from_gk = function(x, ids) {
  fa1 = father(x, ids[1])
  fa2 = father(x, ids[2])
  mo1 = mother(x, ids[1])
  mo2 = mother(x, ids[2])
  GK = function(...) generalisedKinship(x, list(...))
  k0 = GK(fa1, fa2, mo1, mo2)
  k1 = GK(c(fa1, fa2), mo1, mo2) + GK(c(fa1, mo2), fa2, mo1) +
      GK(c(mo1, fa2), fa1, mo2) + GK(c(mo1, mo2), fa1, fa2)
  k2 = GK(c(fa1, fa2), c(mo1, mo2)) + GK(c(fa1, mo2), c(mo1, fa2))
  c(k0, k1, k2)
}
y1 = nuclearPed(2); ids = 3:4
stopifnot(IBD_from_gk(y1, ids) == kappaIBD(y1, ids))
y2 = quadHalfFirstCousins()
ids = 9:10
stopifnot(IBD_from_gk(y2, ids) == kappaIBD(y2, ids))
### Triple/quad kinship (compare with karigl)

```r
x = fullSibMating(1)
ids = c(1, 5, 6)
stopifnot(generalisedKinship(x, list(ids)) == generalisedKinship3(x, ids))
ids = c(1, 5, 6, 5)
stopifnot(generalisedKinship(x, list(ids)) == generalisedKinship4(x, ids))
```

---

**generalised_karigl**  
Karigl's generalised kinship coefficients

---

**Description**

Compute generalised kinship coefficients, as defined by Karigl (1981), involving up to 4 pedigree members. The founders may be inbred; see Examples.

**Usage**

```r
generalisedKinship3(
  x,
  ids,
  sparse = NA,
  chromType = "autosomal",
  verbose = FALSE
)
```

```r
generalisedKinship4(
  x,
  ids,
  sparse = NA,
  chromType = "autosomal",
  verbose = FALSE
)
```

```r
generalisedKinship22(
  x,
  ids,
  sparse = NA,
  chromType = "autosomal",
  verbose = FALSE
)
```

**Arguments**

- `x` A pedigree, in the form of a `pedtools::ped` object.
- `ids` A vector of ID labels, of length 3 for `generalisedKinship3()` and 4 for `generalisedKinship4()` and `generalisedKinship22()`.
inbreeding

### sparse
A positive integer, indicating the pedigree size limit for using sparse arrays. If NA, a default limit of 50 is used.

### chromType
Either "autosomal" or "x".

### verbose
A logical.

### Details
The function `generalisedKinship3()` computes the generalised kinship coefficient of three (not necessarily distinct) members a, b and c, defined as the probability that if a random allele is chosen from each of them, they are all identical by descent.

The function `generalisedKinship4()` computes the generalised kinship coefficient of four individuals, defined similarly to the above.

The function `generalisedKinship22()` computes the generalised kinship coefficient of two pairs of members, defined as the probability that in both pairs simultaneously, random alleles chosen from the two individuals are IBD.

### Value
A numeric of length 1.

### See Also
`kinship()`, `kinshipX()`, `condensedIdentity()`, `condensedIdentityX()`

### Examples
```r
# Generalised kinship between three siblings
x = nuclearPed(3)
phi3 = generalisedKinship3(x, ids = 3:5)

# Recalculate if the father is 100% inbred
founderInbreeding(x, 1) = 1
phi3_inbred = generalisedKinship3(x, ids = 3:5)

stopifnot(phi3 == 1/16, phi3_inbred == 1/8 + 1/32)
```

### inbreeding

<table>
<thead>
<tr>
<th>Inbreeding coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td>inbreeding</td>
</tr>
</tbody>
</table>

### Description
Compute the inbreeding coefficients of all members of a pedigree. These are simple wrappers of `kinship()` and `kinshipX()`. The founders may be inbred; see pedtools::founderInbreeding() for how to set this up.
Usage

inbreeding(x)

inbreedingX(x)

Arguments

x A pedigree, in the form of a pedtools::ped object.

Details

The autosomal inbreeding coefficient of a pedigree member is defined as the probability that, at a random autosomal locus, the two alleles carried by the member are identical by descent relative to the pedigree. It follows from the definition that the inbreeding coefficient of a member equals the kinship coefficient of the parents.

The X chromosomal inbreeding coefficient of an female member is defined similarly to the autosomal case above. For males is it always 1.

The inbreeding coefficients are computed from the kinship coefficients by the formula

\[ f_a = 2 \times \phi_{aa} - 1. \]

Value

A numeric vector of length pedsize(x).

See Also

kinship()

Examples

# Child of half siblings: f = 1/8
x = halfCousinPed(0, child = TRUE)
inbreeding(x)

# If the father is 100% inbred, the inbreeding coeff of the child doubles
founderInbreeding(x, 1) = 1
inbreeding(x)

# The X inbreeding coefficients depend on the genders in the pedigree.
# To exemplify this, we look at a child of half siblings.
# To exemplify this, we look at a child of half siblings.
x.pat = halfSibPed(sex2 = 2) # paternal half sibs
x.pat = addChildren(x.pat, father = 4, mother = 5, nch = 1, sex = 2)
stopifnot(inbreedingX(x.pat)[6] == 0)

# Change to maternal half sibs => coeff becomes 1/4.
x.mat = swapSex(x.pat, 1)
stopifnot(inbreedingX(x.mat)[6] == 0.25)
**jicaque**

**Jicaque pedigree**

*Description*

A data frame describing a pedigree from the Jicaque tribe, studied by Chapman and Jacquard (1971).

*Usage*

```r
jicaque
```

*Format*

A data frame with 22 rows and four columns:

- **id**: individual ID
- **fid**: father’s ID (or 0 if not included)
- **mid**: mother’s ID (or 0 if not included)
- **sex**: Gender codes, where 1 = male and 2 = female

*References*


**kappaIBD**

*IBD (kappa) coefficients*

*Description*

Computes the three IBD coefficients summarising the relationship between two non-inbred individuals. Both autosomal and X chromosomal versions are implemented.

*Usage*

```r
kappaIBD(x, ids = labels(x), inbredAction = 1)
kappaIbdX(x, ids, sparse = NA, verbose = FALSE)
```
Arguments

- **x**: A pedigree in the form of a `pedtools::ped` object.
- **ids**: A character (or coercible to character) containing ID labels of two or more pedigree members.
- **inbredAction**: An integer telling the program what to do if either of the ids individuals are inbred. Possible values are: 0 = do nothing; 1 = print a warning message (default); 2 = raise an error. In the first two cases the coefficients are reported as NA.
- **sparse**: A positive integer, indicating the pedigree size limit for using sparse arrays (as implemented by the slam package) instead of ordinary arrays.
- **verbose**: A logical.

Details

For non-inbred individuals a and b, their autosomal IBD coefficients ($\kappa_0, \kappa_1, \kappa_2$) are defined as follows:

$$
\kappa_i = P(a and b share IBD at random autosomal locus)
$$

The autosomal kappa coefficients are computed from the kinship coefficients. When a and b are both nonfounders, the following formulas are well-known:

- $\kappa_2 = \phi_M M * \phi_F F + \phi_M F * \phi_F M$
- $\kappa_1 = 4 * \phi_a b - 2 * \kappa_2$
- $\kappa_0 = 1 - \kappa_1 - \kappa_2$

Here $\phi_M M$ denotes the kinship coefficient between the mothers of a and b, and so on. If either a or b is a founder, then $\kappa_2 = 0$, while the other two formulas remain as above.

The X chromosomal IBD coefficients are defined as in the autosomal case, with the exception that $\kappa_2$ is undefined when at least one of the two individuals is male. Hence the computation is greatly simplified when males are involved. Denoting the standard kinship coefficient by $\phi$, the formulas are:

- Both male: $(\kappa_0, \kappa_1, \kappa_2) = (1 - \phi, \phi, NA)$
- One male, one female: $(\kappa_0, \kappa_1, \kappa_2) = (1 - 2 * \phi, 2 * \phi, NA)$
- Two females: As in the autosomal case.

Value

If `ids` has length 2: A numeric vector of length 3: $(\kappa_0, \kappa_1, \kappa_2)$.

If `ids` has length > 2: A data frame with one row for each pair of individuals, and 5 columns. The first two columns contain the ID labels, and columns 3-5 contain the IBD coefficients.

Unless `inbredAction = 2`, the coefficients of pairs involving inbred individuals (inbred females in the X version) are reported as NA. Furthermore, the X chromosomal $\kappa_2$ is NA whenever at least one of the two individuals is male.

See Also

`kinship()`, `condensedIdentity()`
Examples

```r
# Siblings
x = nuclearPed(2)
k = kappaIBD(x, 3:4)
stopifnot(identical(k, c(.25, .5, .25)))

# Quad half first cousins
x = quadHalfFirstCousins()
k = kappaIBD(x, leaves(x))
stopifnot(identical(k, c(17/32, 14/32, 1/32)))

# Paternal half brothers with 100% inbred father
# Genetically indistinguishable from an (outbred) father-son relationship
x = halfSibPed()
founderInbreeding(x, 1) = 1

k = kappaIBD(x, 4:5)
stopifnot(identical(k, c(0, 1, 0)))
```

---

**kinPattern**

*Generalised kinship pattern*

Description

Generalised kinship pattern

Usage

```r
kinPattern(x, pattern, internal = FALSE)
```

Arguments

- `x`: A `ped` object
- `pattern`: A list of vectors of ID labels.
- `internal`: A logical

Value

An object of class `kinPattern`.

Examples

```r
kinPattern(nuclearPed(2), list(1, 3:4))
```
## kinship

### Kinship coefficients

**Description**

Compute the matrix of kinship coefficients (autosomal or X) of all members of a pedigree. The founders may be inbred; see `pedtools::founderInbreeding()` for how to set this up.

**Usage**

```r
kinship(x)
kinshipX(x)
```

**Arguments**

- `x` A pedigree, in the form of a `pedtools::ped` object.

**Details**

For two (not necessarily distinct) members A, B of a pedigree, their autosomal (resp. X) kinship coefficient is defined as the probability that random alleles sampled from A and B at the same autosomal (resp. X) locus, are identical by descent relative to the pedigree.

**Value**

A symmetric matrix containing all pairwise kinship coefficients in `x`.

**See Also**

`inbreeding()`, `kappa()`

**Examples**

```r
# Kinship coefficients in a nuclear family with two children
x = nuclearPed(2)
kinship(x)

# X chromosomal kinship coefficients in the same family
kinshipX(x)

# Recalculate the autosomal kinships if the father is 100% inbred
founderInbreeding(x, 1) = 1
kinship(x)
```
**minimalPattern**

**Minimal IBD pattern**

**Description**

Compute the minimal form of given multiperson IBD pattern.

**Usage**

```r
minimalPattern(x)
```

**Arguments**

- `x` An integer vector of even length.

**Value**

An integer vector of the same length as `x`.

**Examples**

```r
v = c(1,2,2,3)
stopifnot(identical(minimalPattern(v), c(1,2,1,3)))
```

---

**multiPersonIBD**

**Multi-person IBD coefficients**

**Description**

Computes the probabilities (coefficients) of all possible patterns of identity by descent (IBD) sharing at a single locus, among N>1 non-inbred members of a pedigree. The reported coefficients are "condensed" in the sense that allele ordering within each individual is ignored. For N = 2, the result should agree with the traditional "kappa" coefficients, as computed by `kappaIBD()`. This function is under development, and should be regarded as experimental. For now, the only cases handled are those with: N = 2 or 3, autosomal locus.

**Usage**

```r
multiPersonIBD(x, ids, complete = FALSE, verbose = FALSE)
```

**Arguments**

- `x` A ped object.
- `ids` A vector of ID labels.
- `complete` A logical. If FALSE, only IBD patterns with nonzero probability are included in the output.
- `verbose` A logical. If TRUE, some computational details are printed.
Details

Consider N members of a pedigree, i1, i2, ... iN. A pattern of IBD sharing between these individuals is a sequence of N ordered pairs of labels, (a1_1, a1_2), (a2_1, a2_2), ... (aN_1, aN_2), where ai_1 and ai_2 represent the paternal and maternal allele of individual i, respectively. Equality of labels means that the corresponding alleles are IBD, and vice versa.

We say that two IBD patterns are equivalent if one can be transformed into the other by some combination of

- renaming the labels (without changing the structure)
- swapping the paternal/maternal labels of some individuals

Each equivalence class has a "minimal" element, using integer labels, and being minimal with respect to standard sorting. For example, the minimal element equivalent to (a,c),(d,c),(b,b) is (1,2),(2,3),(4,4).

Value

A data frame in which each row corresponds to an equivalence class of multi-person IBD patterns. The first column gives the calculated probability, followed by one column for each ids individual, describing the minimal element of the equivalence class. (See Details.) If complete = FALSE (the default) rows with probability 0 are removed.

Examples

```r
### Trivial example: Trio ###
x = nuclearPed(1)
ids = 1:3
multiPersonIBD(x, ids, complete = TRUE)

### Example due to Peter Green ###
# Three (pariwise) cousins arranged in two different ways,
# with different 3-way IBD coefficients.

threeCousins1 = ped(
    id = c('gf', 'gm', 'gf1', 'gf2', 'gf3', 'gm1', 'gm2', 'gm3',
           'f1', 'f2', 'f3', 'm1', 'm2', 'm3', 'c1', 'c2', 'c3'),
    fid = c(0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0),
    mid = c(0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0),
    sex = c(1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1)),

threeCousins2 = ped(
    id = c('gf1', 'gf2', 'gf3', 'gm1', 'gm2', 'gm3', 'f1', 'f2', 'f3',
           'm1', 'm2', 'm3', 'c1', 'c2', 'c3'),
    fid = c(0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0),
    mid = c(0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0,0),
    sex = c(1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1))
```
ids = c('c1','c2','c3')
multiPersonIBD(threeCousins1, ids)
multiPersonIBD(threeCousins2, ids)

**ribd**

*ribd: Computation of pedigree-based relatedness coefficients*

**Description**

Recursive algorithms for computing various relatedness coefficients, including Jacquard’s condensed identity coefficients. The standard algorithms are extended to allow inbred founders. Both autosomal and X-linked coefficients are computed.

**twoLocusIBD**

*Two-locus IBD coefficients*

**Description**

Computes the 3*3 matrix of two-locus IBD coefficients of a pair of non-inbred pedigree members, for a given recombination rate.

**Usage**

twoLocusIBD(
  x,
  ids,
  rho,
  coefs = NULL,
  detailed = FALSE,
  uniMethod = 1,
  verbose = FALSE
)

**Arguments**

- **x**: A pedigree in the form of a `pedtools::ped` object.
- **ids**: A character (or coercible to character) containing ID labels of two pedigree members.
- **rho**: A number in the interval [0, 0.5]; the recombination rate between the two loci.
- **coefs**: A character indicating which coefficient(s) to compute. A subset of c('k00', 'k01', 'k02', 'k10', 'k11', 'k12'). By default, all coefficients are computed.
- **detailed**: A logical, indicating whether the condensed (default) or detailed coefficients should be returned.
- **uniMethod**: Either 1 or 2 (for testing purposes)
- **verbose**: A logical.
Details

Let A, B be two pedigree members, and L1, L2 two loci with a given recombination rate $\rho$. The two-locus IBD coefficients $\kappa_{i,j}(\rho)$, for $0 \leq i, j \leq 2$ are defined as the probability that A and B have $i$ alleles IBD at L1 and $j$ alleles IBD at L2 simultaneously. Note that IBD alleles at the two loci are not required to be in cis (or in trans for that matter).

The method of computation depends on the (single-locus) IBD coefficient $\kappa_2$. If this is zero (e.g. if A is a direct ancestor of B, or vice versa) the two-locus IBD coefficients are easily computable from the two-locus kinship coefficients, as implemented in `twoLocusKinship()`. In the general case, the computation is more involved, requiring *generalised two-locus kinship* coefficients. This is implemented in the function `twoLocusGeneralisedKinship()`, which is not exported yet.

Value

By default, a symmetric 3*3 matrix containing the two-locus IBD coefficients $\kappa_{i,j}$.

If either `coefs` is explicitly given (i.e., not NULL), or `detailed = TRUE`, the computed coefficients are returned as a named vector.

See Also

twoLocusKinship()

Examples

# Some variables used in several examples below
rseq = seq(0, 0.5, length = 11) # recombination values

xlab = "Recombination rate"
main = expression(paste("Two-locus IBD: \( k_{1,1} \)\))

# Example 1: A classic example of three relationships with the same
# one-locus IBD coefficients, but different two-locus coefficients.
# As a consequence, these relationships cannot be separated with unlinked markers, but are (theoretically) separable with linked markers.
# Compute \( k_{1,1} \) for each rho
peds = list(
  GrandParent = list(ped = linearPed(2), ids = c(1, 5)),
  HalfSib = list(ped = halfSibPed(), ids = c(4, 5)),
  Uncle = list(ped = cousinPed(0, 1), ids = c(3, 6))
)

# Plot
matplot(rseq, kvals, type = "l", xlab = xlab, ylab = "", main = main)
legend("topright", names(peds), col = 1:3, lty = 1:3)
twoLocusIBD

# Example 2: Inspired by Fig. 3 in Thompson (1988),
# and its erratum: https://doi.org/10.1093/imammb/6.1.1.
#
# These relationships are also analysed in ?twoLocusKinship,
# where we show that they have identical two-locus kinship
# coefficients. Here we demonstrate that they have different
# two-locus IBD coefficients.

# List of pedigrees and ID pairs
peds = list(
   GreatGrand = list(ped = linearPed(3), ids = c(1, 7)),
   HalfUncle = list(ped = halfCousinPed(0, 1), ids = c(3, 7))
)

# Compute 'k11' for each rho
kvals = sapply(peds, function(x)
   sapply(rseq, function(r) twoLocusIBD(x$ped, x$ids, r, coefs = "k11")))

# Plot
matplot(rseq, kvals, type = "l", xlab = xlab, ylab = "", main = main)
legend("topright", names(peds), col = 1:2, lty = 1:2)

# Example 3: Two-locus IBD of two half sisters whose mother have
# inbreeding coefficient 1/4. We compare two different realisations
# of this:
# PO: the mother is the child of parent-offspring
# SIB: the mother is the child of full siblings
#
# We show below that these relationships have different two-locus
# coefficients. This exemplifies that a single-locus inbreeding
# coefficient cannot replace the genealogy in analyses of linked loci.

xPO = addChildren(nuclearPed(1, sex = 2), 1, 3, nch = 1, sex = 2)
xPO = addDaughter(addDaughter(xPO, 4), 4)
xSIB = addChildren(nuclearPed(2, sex = 1:2), 3, 4, nch = 1)
xSIB = addDaughter(addDaughter(xSIB, 5), 5)

plotPedList(list(xPO, xSIB), new = TRUE, title = c("PO", "SIB"))

# List of pedigrees and ID pairs
peds = list(PO = list(ped = xPO, ids = c(6, 8)),
             SIB = list(ped = xSIB, ids = c(7, 9)))

# Compute 'k11' for each rho
kvals = sapply(peds, function(x)
   sapply(rseq, function(r) twoLocusIBD(x$ped, x$ids, r, coefs = "k11")))
twoLocusIBD

```r
# Plot
matplot(rseq, kvals, type = "l", xlab = xlab, ylab = "", main = main)
legend("topright", names(peds), col = 1:2, lty = 1:2)

# Check against exact formula
r = rseq
k11_PO = 1/8*(-4*r^5 + 12*r^4 - 16*r^3 + 16*r^2 - 9*r + 5)
stopifnot(all.equal(kvals[, "PO"], k11_PO, check.names = FALSE))

k11_S = 1/16*(8*r^6 - 32*r^5 + 58*r^4 - 58*r^3 + 43*r^2 - 20*r + 10)
stopifnot(all.equal(kvals[, "SIB"], k11_S, check.names = FALSE))

################################################
# Example 4:
# The complete two-locus IBD matrix of full sibs
################################################

x = nuclearPed(2)
k2_mat = twoLocusIBD(x, ids = 3:4, rho = 0.25)
k2_mat

# Compare with explicit formulas
IBDSibs = function(rho) {
  R = rho^2 + (1-rho)^2
  nms = c("ibd0", "ibd1", "ibd2")
  m = matrix(0, nrow = 3, ncol = 3, dimnames = list(nms, nms))
  m[1,1] = m[3,3] = 0.25 * R^2
  m[2,1] = m[1,2] = 0.5 * R * (1-R)
  m[3,1] = m[1,3] = 0.25 * (1-R)^2
  m[2,2] = 0.5 * (1 - 2 * R * (1-R))
  m[3,2] = m[2,3] = 0.5 * R * (1-R)
  m
}

stopifnot(all.equal(k2_mat, IBDSibs(0.25)))

#####################################################
# Example 5: Two-locus IBD of quad half first cousins
# We use this to exemplify two simple properties of
# the two-locus IBD matrix.
#####################################################

x = quadHalffirstCousins()
ids = c(9, 10)

# First compute the one-locus IBD coefficients (= c(17, 14, 1)/32)
k1 = kappaIBD(x, ids)

### Case 1: Complete linkage (\(\rho = 0\)).
# In this case the two-locus IBD matrix has `k1` on the diagonal,
```
twoLocusIdentity

Two-locus identity coefficients

Description

Computes the 9x9 matrix of two-locus condensed identity coefficients of a pair of pedigree members, for a given recombination rate.

Usage

twoLocusIdentity(x, ids, rho, coefs = NULL, detailed = FALSE, verbose = FALSE)

Arguments

x  A pedigree in the form of a pedtools::ped object.
ids  A character (or coercible to character) containing ID labels of two pedigree members.
rho  A number in the interval [0, 0.5]; the recombination rate between the two loci.
coefs  A character indicating which coefficient(s) to compute. A subset of c('d00', 'd01', ..., 'd99'). By default, all coefficients are computed.
detailed  A logical, indicating whether the condensed (default) or detailed coefficients should be returned.
verbose  A logical.
Details

Let A, B be two pedigree members, and L1, L2 two loci with a given recombination rate \( \rho \). The two-locus identity coefficients \( \Delta_{i,j}(\rho) \), for \( 1 \leq i, j \leq 9 \) are defined as the probability that the identity state of the alleles of A and B are \( \Sigma_i \) at L1 and \( \Sigma_j \) at L2 simultaneously. (The ordering of the 9 states follows Jacquard (1974).)

For details about the algorithm, see Vigeland (2019).

Value

By default, a symmetric 9*9 matrix containing the two-locus condensed identity coefficients \( \Delta_{i,j} \).

If either coefs is explicitly given (i.e., not NULL), or detailed = TRUE, the computed coefficients are returned as a named vector.

References


See Also

twoLocusIBD()

Examples

```r
### Full sibs ###
x = nuclearPed(2)
kapp = twoLocusIBD(x, ids = 3:4, rho = 0.25)
jacq = twoLocusIdentity(x, ids = 3:4, rho = 0.25)
stopifnot(all.equal(jacq[9:7,9:7], kapp, check.attributes = FALSE))

#' ### Parent-child ###
x = nuclearPed(1)
jacq = twoLocusIdentity(x, ids = c(1,3), rho = 0.25)
stopifnot(jacq[8,8] == 1)

### Full sib mating ###
x = fullSibMating(1)
j = condensedIdentity(x, ids = 5:6)
j2 = twoLocusIdentity(x, ids = 5:6, rho = 0.25)
stopifnot(identical(unname(rowSums(j2)), j))
```
twoLocusKinship

**Description**
Computes the two-locus kinship coefficient of a pair of pedigree members, at a given recombination rate.

**Usage**
```
twoLocusKinship(
  x,
  ids,
  rho,
  recombinants = NULL,
  verbose = FALSE,
  debug = FALSE
)
```

**Arguments**
- `x`: A pedigree in the form of a `pedtools::ped` object.
- `ids`: A character (or coercible to character) containing ID labels of two or more pedigree members.
- `rho`: A numeric vector of recombination rates; all entries must be in the interval [0, 0.5].
- `recombinants`: A logical of length 2, applicable only when `ids` has length 2. When given, it indicates whether each of the two gametes is a recombinant or non-recombinant. This parameter is mainly used by `twoLocusIBD()`.
- `verbose`: A logical.
- `debug`: A logical. If TRUE, detailed messages are printed during the recursion process.

**Details**
Let A, B be two pedigree members, and L1, L2 two loci with a given recombination rate rho. The two-locus kinship coefficient $\phi_{AB}(\rho)$ is defined as the probability that random gametes segregating from A and B has IBD alleles at both L1 and L2 simultaneously.

The implementation is based on the recursive algorithm described by Thompson (1988).

**References**
**Examples**

```
# Example 1: Full sibs
x = nuclearPed(2)

k_0 = twoLocusKinship(x, ids = 3:4, rho = 0)
k_0.5 = twoLocusKinship(x, ids = 3:4, rho = 0.5)

stopifnot(k_0 == 1/4, k_0.5 == 1/16)
```

```
# Example 2: Reproducing Fig. 3 in Thompson (1988)
# Note that in the article, curve (a) is wrong.
# See Erratum: https://doi.org/10.1093/imammb/6.1.1

# Pedigrees (a) - (d)
peds = list(
a = list(ped = linearPed(3), ids = c(1,7)),
b = list(ped = halfCousinPed(0, 1), ids = c(3,7)),
c = list(ped = cousinPed(1), ids = c(5,8)),
d = list(ped = doubleCousins(1, 1, half1 = TRUE, half2 = TRUE), ids = c(5,9))
)

# Recombination values
rseq = seq(0, 0.5, length = 20)

# Compute two-locus kinship coefficients
kvals = sapply(peds, function(x) twoLocusKinship(x$ped, x$ids, rseq))

# Plot
matplot(rseq, kvals, type = "l", lwd = 2)
legend("topright", names(peds), col = 1:4, lty = 1:4)
```

**twoLocusPlot**

**Two-locus coefficient plot**

**Description**

Plot two-locus kinship or IBD coefficients as function of the recombination rate.

**Usage**

```r
twoLocusPlot(
peds,
coeff = "k11",
```
twoLocusPlot

xlab = "Recombination rate",
ylab = NA,
col = seq_along(peds),
lty = 1,

)

Arguments

peds        A list of lists. See details.
coeff       A string identifying which coefficient to compute. See Details for legal values.
xlab, ylab, col, lty
            Plotting parameters
            ...
            Further parameters passed on to matplot()

Details

Each entry of peds must be a list with the following (named) entries:

- ped: A ped object
- ids: A pair of labels identifying two members of ped

The coeff parameter must be either a character naming the coefficient to compute, or a function. If a character, it must be one of the following names: "kinship", "phi", "phi11", "k00", "k01", "k02", "k10", "k11", "k12", "k20", "k21" or "k22".

If coeff is a function, it must take three arguments named ped, ids and rho, and produce a single number for each set of input data. See Examples.

The first three are synonymous and indicate the two-locus kinship coefficient. The remaining choices are two-locus IBD coefficients. (See twoLocusIBD().)

Examples

###############################
# Classic example of three relationships with equal one-locus coeffs
peds = list(
  GrandParent = list(ped = linearPed(2), ids = c(1, 5)),
  HalfSib = list(ped = halfSibPed(), ids = c(4, 5)),
  Uncle = list(ped = cousinPed(0, 1), ids = c(3, 6)))

twoLocusPlot(peds, coeff = "kinship")
twoLocusPlot(peds, coeff = "k11")

###############################

peds = list(
  PO = list(ped = nuclearPed(1), ids = c(1,3)),
  S = list(ped = nuclearPed(2), ids = c(3,4)))

twoLocusPlot(peds, coeff = "kinship")
twoLocusPlot(peds, coeff = "k11")

#############################################################
peds = list(
  'H-sibs' = list(ped = addChildren(halfSibPed(sex2 = 2), 4, 5, nch = 2),
    ids = 6:7),
  'G-sibs' = list(ped = addChildren(addDaughter(nuclearPed(1), 3), 1, 5, nch = 2),
    ids = 6:7),
  'U-sibs' = list(ped = addChildren(addDaughter(nuclearPed(2), 4), 3, 6, nch = 2),
    ids = 7:8)
)
# plotPedList(peds)
twoLocusPlot(peds, coeff = "kinship")

#############################################################
### Example where 'coeff' is a function.
### Reproducing Fig 2 of Bishop & Williamson (1990)

# The coefficient d11(rho) is the conditional probability of IBD = 1
# in the first locus, given IBD = 1 in the second.
peds = list(
  GrandParent = list(ped = linearPed(2), ids = c(1, 5)),
  HalfSib = list(ped = halfSibPed(), ids = c(4, 5)),
  Uncle = list(ped = cousinPed(0, 1), ids = c(3, 6)),
  FirstCous = list(ped = cousinPed(1), ids = c(5, 8)),
  FirstCous1R = list(ped = cousinPed(1, 1), ids = c(5, 10)),
  SecondCous = list(ped = cousinPed(2), ids = c(7, 12)))
d11 = function(ped, ids, rho) {
  twoLocusIBD(ped, ids, rho, coefs = "k11")/kappaIBD(ped, ids)[2]
}
twoLocusPlot(peds, coeff = d11)
Index

*Topic datasets
  jicaque, 11
condensedIdentity, 2
condensedIdentity(), 4, 9, 12
condensedIdentityX, 3
condensedIdentityX(), 3, 9
external_coefs, 5
generalised_karigl, 8
generalisedKinship, 6
generalisedKinship22
  (generalised_karigl), 8
generalisedKinship3
  (generalised_karigl), 8
generalisedKinship4
  (generalised_karigl), 8
idcoefs (external_coefs), 5
idcoefs2 (external_coefs), 5
identity::identity.coefs(), 2, 5, 6
inbreeding, 9
inbreeding(), 14
inbreedingX (inbreeding), 9
jicaque, 11
kappa(), 3, 14
kappaIBD, 11
kappaIBD(), 15
kappaIbdX (kappaIBD), 11
kinPattern, 13
kinship, 14
kinship(), 9, 10, 12
kinship2::kinship(), 5, 6
kinship2_inbreeding (external_coefs), 5
kinship2_inbreedingX (external_coefs), 5
kinship2_kinship (external_coefs), 5
kinship2_kinshipX (external_coefs), 5
kinshipX (kinship), 14
kinshipX(), 4, 9
matplot(), 25
minimalPattern, 15
multiPersonIBD, 15
pedtools::founderInbreeding(), 2–4, 9,
  14
pedtools::ped, 2, 4, 5, 8, 10, 12, 14, 17, 21,
  23
ribd, 17
system(), 5
twoLocusIBD, 17
twoLocusIBD(), 22, 23, 25
twoLocusIdentity, 21
twoLocusKinship, 23
twoLocusKinship(), 18
twoLocusPlot, 24

27