Package ‘pedtools’

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Type Package
Title Creating and Working with Pedigrees and Marker Data
Version 1.3.0
Description A comprehensive collection of tools for creating, manipulating and visualising pedigrees and genetic marker data. Pedigrees can be read from text files or created on the fly with built-in functions. A range of utilities enable modifications like adding or removing individuals, breaking loops, and merging pedigrees. Pedigree plots are produced by wrapping the plotting functionality of the 'kinship2' package. A Shiny app for creating pedigrees, based on 'pedtools', is available at <https://magnusdv.shinyapps.io/quickped>. 'pedtools' is the hub of the 'ped suite', a collection of packages for pedigree analysis. A detailed presentation of the 'ped suite' is given in the book 'Pedigree Analysis in R' (Vigeland, 2021, ISBN:9780128244302).

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**Description**

Extends the allele set of a marker attached to a pedigree, by adding a single allele.

**Usage**

```r
addAllele(x, marker, allele, freq = 0.001, adjust = c("previous", "all"))
```

**Arguments**

- **x**: A ped object or a list of such, or a frequency database (list of numeric vectors).
- **marker**: The name or index of a marker attached to x.
- **allele**: The name of the new allele.
- **freq**: The frequency of the new allele, by default 0.001.
- **adjust**: Either "previous" or "all", indicating how the frequencies should be adjusted so that they sum to 1. If "previous" (default), the frequencies of the original alleles are multiplied with 1 - freq. If "all", scaling is performed after adding the new allele, i.e., dividing all frequencies by 1 + freq.

**Value**

A copy of x with modified marker attributes.

**Examples**

```r
## Ped input
x = nuclearPed() |> 
  addMarker(geno = c(NA, NA, "b/c"), afreq = c(b = 0.5, c = 0.5))

y = addAllele(x, marker = 1, allele = "a")
afreq(y, 1)

z = addAllele(y, marker = 1, allele = "d", freq = 0.1, adjust = "all")
afreq(z, 1)

## Database input
db = list(M1 = c(a = 0.2, b = 0.3, c = 0.5),
```

---

**addAllele**

- [73]
- [74]
- [75]
- [76]

**Index**

- addAllele 78
M2 = c("7" = .9, "8.3" = .1))
addAllele(db, marker = "M2", allele = "8")

as.data.frame.ped  Convert ped to data.frame

Description
Convert a ped object to a data.frame. The first columns are id, fid, mid and sex, followed by
genotype columns for all (or a selection of) markers.

Usage
## S3 method for class 'ped'
as.data.frame(x, ..., markers, sep = "/", missing = "-")

Arguments
x  Object of class ped.
...  Further parameters
markers  Vector of marker names or indices. By default, all markers are included.
sep  A single string to be used as allele separator in marker genotypes.
missing  A single string to be used for missing alleles.

Details
Note that the output of as.data.frame.ped() is quite different from that of as.matrix.ped().
This reflects the fact that these functions have different purposes.
Conversion to a data frame is primarily intended for pretty printing. It uses correct labels for pedi-
gree members and marker alleles, and pastes alleles to form nice-looking genotypes.
The matrix method, on the other hand, is a handy tool for manipulating the pedigree structure.
It produces a numeric matrix, using the internal index labelling both for individuals and alleles,
making it very fast. In addition, all necessary meta information (loop breakers, allele frequencies
a.s.o) is kept as attributes, which makes it possible to recreate the original ped object.

Value
A data.frame with pedsize(x) rows and 4 + nMarkers(x) columns.

See Also
as.matrix.ped()
**Description**

Converts a ped object to a numeric matrix using internal labels, with additional info necessary to recreate the original ped attached as attributes.

**Usage**

```r
## S3 method for class 'ped'
as.matrix(x, include.attrs = TRUE, ...)

restorePed(x, attrs = NULL, validate = TRUE)
```

**Arguments**

- `include.attrs`: a logical indicating if marker annotations and other info should be attached as attributes. See Value.
- `...`: not used.
- `attrs`: a list containing labels and other ped info compatible with `x`, in the format produced by `as.matrix`. If `NULL`, the attributes of `x` itself are used.
- `validate`: a logical, forwarded to `ped()`. If `FALSE`, no checks for pedigree errors are performed.

**Details**

`restorePed` is the reverse of `as.matrix.ped`.

**Value**

For `as.matrix`: A numerical matrix with `pedsize(x)` rows. If `include.attrs = TRUE` the following attributes are added to the matrix, allowing `x` to be exactly reproduced by `restorePed`:

- `FAMID` the family identifier (a string)
- `LABELS` the ID labels (a character vector)
- `UNBROKEN_LOOPS` a logical indicating whether `x` has unbroken loops
- `LOOPBREAKERS` a numerical matrix, or `NULL`
- `markerattr` a list of length `nMarkers(x)`, containing the attributes of each marker

For `restorePed`: A ped object.

**Author(s)**

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See Also

`ped()`

Examples

```r
x = relabel(nuclearPed(1), letters[1:3])

# To examplify the ped -> matrix -> ped trick, we show how to
# reverse the internal ordering of the pedigree.
 m = as.matrix(x, include.attrs = TRUE)
m[] = m[3:1, ]

# Must reverse the labels also:
attrs = attributes(m)
attrs$LABELS = rev(attrs$LABELS)

# Restore ped:
y = restorePed(m, attrs = attrs)

# Of course a simpler way is use reorderPed():
z = reorderPed(x, 3:1)
stopifnot(identical(y, z))
```

---

as.ped

Conversions to `ped` objects

Description

Conversions to ped objects

Usage

```r
as.ped(x, ...)
```

```r
## S3 method for class 'data.frame'
as.ped(
  x,
  famid_col = NA,
  id_col = NA,
  fid_col = NA,
  mid_col = NA,
  sex_col = NA,
  marker_col = NA,
  locusAttributes = NULL,
  missing = 0,
  sep = NULL,
```
as.ped

    validate = TRUE,
    ...
)

Arguments

x             Any object.
...             Not used.
famid_col     Index of family ID column. If NA, the program looks for a column named "famid" (ignoring case).
id_col        Index of individual ID column. If NA, the program looks for a column named "id" (ignoring case).
fid_col       Index of father ID column. If NA, the program looks for a column named "fid" (ignoring case).
mid_col       Index of mother ID column. If NA, the program looks for a column named "mid" (ignoring case).
sex_col       Index of column with gender codes (0 = unknown; 1 = male; 2 = female). If NA, the program looks for a column named "sex" (ignoring case). If this is not found, genders of parents are deduced from the data, leaving the remaining as unknown.
marker_col    Index vector indicating columns with marker alleles. If NA, all columns to the right of all pedigree columns are used. If sep (see below) is non-NULL, each column is interpreted as a genotype column and split into separate alleles with strsplit(..., split = sep, fixed = TRUE).
locusAttributes     Passed on to setMarkers() (see explanation there).
missing       Passed on to setMarkers() (see explanation there).
sep           Passed on to setMarkers() (see explanation there).
validate      A logical indicating if the pedigree structure should be validated.

Value

A ped object or a list of such.

Examples

df = data.frame(famid = c("S1", "S2"),
      id = c("A", "B"),
      fid = 0,
      mid = 0,
      sex = 1)

# gives a list of two singletons
as.ped(df)

# Trio
df1 = data.frame(id = 1:3, fid = c(0,0,1), mid = c(0,0,2), sex = c(1,2,1))
as.ped(df1)

# Disconnected example: Trio (1-3) + singleton (4)
df2 = data.frame(id = 1:4, fid = c(2,0,0,0), mid = c(3,0,0,0),
                 M = c("1/2", "1/1", "2/2", "3/4"))
as.ped(df2)

# Two singletons
df3 = data.frame(id = 1:2, fid = 0, mid = 0, sex = 1)
as.ped(df3)

---

connectedComponents  Connected pedigree components

Description

Compute the connected parts of a pedigree. This is an important step when converting pedigree data from other formats (where disconnected pedigrees may be allowed) to pedtools (which requires pedigrees to be connected).

Usage

connectedComponents(id, fid = NULL, mid = NULL, fidx = NULL, midx = NULL)

Arguments

id  A vector of ID labels (character or numeric).
fid  The ID labels of the fathers (or "0" if missing).
mid  The ID labels of the mothers (or "0" if missing).
fidx, midx  (For internal use mostly.) Integer vectors with paternal (resp. maternal) indices. These may be given instead of id, fid, mid.

Value

A list, where each element is a subset of id constituting a connected pedigree.

Examples

# A trio (1-3) and a singleton (4)
x = data.frame(id = 1:4, fid = c(2,0,0,0), mid = c(3,0,0,0))
connectedComponents(x$id, x$fid, x$mid)
distributeMarkers

Distribute markers evenly along a set of chromosomes

Description

Create and attach identical (empty) marker objects, distributed along a set of chromosomes.

Usage

distributeMarkers(
  x,  
  n = NULL,  
  dist = NULL,  
  chromLen = NULL,  
  alleles = 1:2,  
  afreq = NULL,  
  prefix = "M"
)

Arguments

x               A ped object.
n              The total number of markers. Either this or dist must be NULL.
dist            A positive number; the distance (in megabases) between markers.
chromLen        A numeric vector indicating chromosome lengths (in Mb). By default, the
                 lengths of the human chromosomes 1-22 are used, as returned by
                 sapply(ibdsim2::loadMap("decode"),
                          ibdsim2::physRange).
ableles, afreq   Passed onto marker().
prefix          A string used as prefix for marker names. Default: "M".

Details

Note: When using the dist parameter, the function treats each chromosome separately, places one
marker at the start and then every dist megabases. (See Examples.)

Value

A copy of x with the indicated markers attached.

Examples

x = distributeMarkers(nuclearPed(), n = 10)
getMap(x)

y = distributeMarkers(nuclearPed(), dist = 100)
getMap(y)
### famid  
*Family identifier*

**Description**

Functions for getting or setting the family ID of a `ped` object.

**Usage**

```r
famid(x, ...)  
## S3 method for class 'ped'
famid(x, ...)  
famid(x, ...) <- value  
## S3 replacement method for class 'ped'
famid(x, ...) <- value
```

**Arguments**

- `x`: A `ped` object
- `...`: (Not used)
- `value`: The new family ID, which must be (coercible to) a character string.

**Examples**

```r
x = nuclearPed(1)  
famid(x) # empty string  
famid(x) = "trio"  
famid(x)
```

### founderInbreeding  
*Inbreeding coefficients of founders*

**Description**

Functions to get or set inbreeding coefficients for the pedigree founders.

**Usage**

```r
founderInbreeding(x, ids, named = FALSE, chromType = "autosomal")  
founderInbreeding(x, ids, chromType = "autosomal") <- value
```
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td>A ped object.</td>
</tr>
<tr>
<td>ids</td>
<td>Any subset of founders(x). If ids is missing in founderInbreeding(), it is set to founders(x).</td>
</tr>
<tr>
<td>named</td>
<td>A logical: If TRUE, the output vector is named with the ID labels.</td>
</tr>
<tr>
<td>chromType</td>
<td>Either &quot;autosomal&quot; (default) or &quot;x&quot;.</td>
</tr>
<tr>
<td>value</td>
<td>A numeric of the same length as ids, entries in the interval ([0, 1]). If the vector is named, then the names are interpreted as ID labels of the founders whose inbreeding coefficients should be set. In this case, the ids argument should not be used. (See examples.)</td>
</tr>
</tbody>
</table>

Value

For founderInbreeding, a numeric vector of the same length as ids, containing the founder inbreeding coefficients.

For founderInbreeding<-, the updated ped object is returned.

Examples

```r
x = nuclearPed(father = "fa", mother = "mo", child = 1)
founderInbreeding(x, "fa") = 1
founderInbreeding(x, named = TRUE)

# Setting all founders at once (replacement value is recycled)
founderInbreeding(x, ids = founders(x)) = 0.5
founderInbreeding(x, named = TRUE)

# Alternative syntax, using a named vector
founderInbreeding(x) = c(fa = 0.1, mo = 0.2)
founderInbreeding(x, named = TRUE)
```

freqDatabase

**Allele frequency database**

Description

Functions for reading, setting and extracting allele frequency databases, in either "list" format or "allelic ladder" format.

Usage

```r
getFreqDatabase(x, markers = NULL, format = c("list", "ladder"))

setFreqDatabase(x, database, format = c("list", "ladder"), ...)
```
freqDatabase

readFreqDatabase(filename, format = c("list", "ladder"), ...)
writeFreqDatabase(x, filename, markers = NULL, format = c("list", "ladder"))

Arguments

x A ped object, or a list of such.
markers A character vector (with marker names) or a numeric vector (with marker indices).
format Either "list" or "ladder".
database Either a list or matrix/data frame with allele frequencies, or a file path (to be passed on to readFreqDatabase()).
... Optional arguments passed on to read.table().
filename The path to a text file containing allele frequencies either in "list" or "allelic ladder" format.

Details

A frequency database in "list" format is a list of numeric vectors; each vector named with the allele labels, and the list itself named with the marker names.

Text files containing frequencies in "list" format should look as follows, where "marker1" and "marker2" are marker names, and "a1","a2"... are allele labels (which may be characters or numeric, but will always be converted to characters):

```
marker1
a1 0.2
a2 0.5
a3 0.3
```

```
marker2
a1 0.9
a2 0.1
```

A database in "allelic ladder" format is rectangular, i.e., a numeric matrix (or data frame), with allele labels as row names and markers as column names. NA entries correspond to unobserved alleles.

Value

- getFreqDatabase: either a list (if format = "list") or a data frame (if format = "ladder")
- readFreqDatabase: a list (also if format = "ladder") of named numeric vectors
- setFreqDatabase: a modified version of x

See Also

setLocusAttributes(), setMarkers(), setAlleles()
Examples

loc1 = list(name = "m1", afreq = c(a = .1, b = .9))
loc2 = list(name = "m2", afreq = c("1" = .2, "10.2" = .3, "3" = .5))
x = setMarkers(singleton(1), locus = list(loc1, loc2))
db = getFreqDatabase(x)
db
y = setFreqDatabase(x, database = db)
stopifnot(identical(x, y))

# The database can also be read directly from file
tmp = tempfile()
write("m1\na 0.1\nb 0.9\nm2\n1 0.2\n3 0.5\n10.2 0.3", tmp)

z = setFreqDatabase(x, database = tmp)
stopifnot(all.equal(x, z))

getAlleles  Allele matrix manipulation

Description

Functions for getting and setting the genotypes of multiple individuals/markers simultaneously

Usage

getAlleles(x, ids = NULL, markers = NULL)
setAlleles(x, ids = NULL, markers = NULL, alleles)

Arguments

x  A ped object or a list of such
ids  A vector of ID labels. If NULL (default) all individuals are included.
markers  A vector of indices or names of markers attaches to x. If NULL (default) all markers are included.
alleles  A character of the same format and dimensions as the output of getAlleles(x, ids, markers), or an object which can be converted by as.matrix() into such a matrix. See Details.

Details

If the alleles argument of setAlleles() is not a matrix, it is recycled (if necessary), and converted into a matrix of the correct dimensions. For example, setting alleles = 0 gives a simple way of removing the genotypes of some or all individuals (while keeping the markers attached).
Value

getAlleles() returns a character matrix with length(ids) rows and 2 * length(markers) columns. The ID labels of x are used as rownames, while the columns are named <m1>.1, <m1>.2, ... where <m1> is the name of the first marker, a.s.o.

setAlleles() returns a ped object identical to x, except for the modified alleles. In particular, all locus attributes are unchanged.

See Also

transferMarkers()

Examples

# Setup: Pedigree with two markers
x = nuclearPed(1)
x = addMarker(x, `2` = "1/2", alleles = 1:2, name = "m1")
x = addMarker(x, `3` = "2/2", alleles = 1:2, name = "m2")

# Extract allele matrix
mat1 = getAlleles(x)
mat2 = getAlleles(x, ids = 2:3, markers = "m2")
stopifnot(identical(mat1[2:3, 3:4], mat2))

# Remove all genotypes
y = setAlleles(x, alleles = 0)
y

# Setting a single genotype
z = setAlleles(y, ids = "1", marker = "m2", alleles = 1:2)

# Alternative: In-place modification with `genotype()`
genotype(y, id = "1", marker = "m2") = "1/2"
stopifnot(identical(y, z))

### Manipulation of pedlist objects
s = transferMarkers(x, singleton("s"))
peds = list(x, s)

getAlleles(peds)

setAlleles(peds, ids = "s", marker = "m1", alleles = 1:2)
getGenotypes

Description

Given a list of ped objects (called pedigree components), and a vector of ID labels, find the index of the component holding each individual.

Usage

getComponent(x, ids, checkUnique = FALSE, errorIfUnknown = FALSE)

Arguments

x  A list of ped objects
ids  A vector of ID labels (coercible to character)
checkUnique  If TRUE an error is raised if any element of ids occurs more than once in x. Default: FALSE.
errorIfUnknown  If TRUE, the function stops with an error if not all elements of ids are recognised as names of members in x. Default: FALSE.

Value

An integer vector of the same length as ids, with NA entries where the corresponding label was not found in any of the components.

See Also

internalID()

Examples

x = list(nuclearPed(1), singleton(id = "A"))
getComponent(x, c(3, "A"))

getGenotypes

Genotype matrix

Description

Extract the genotypes of multiple individuals/markers in form of a matrix.

Usage

getGenotypes(x, ids = NULL, markers = NULL, sep = "/", missing = "-"
Arguments

x  A ped object or a list of such
ids  A vector of ID labels. If NULL (default) all individuals are included.
markers  A vector of indices or names of markers attaches to x. If NULL (default) all markers are included.
sep  A single string to be used as allele separator in marker genotypes.
missing  A single string to be used for missing alleles.

Value

getGenotypes() returns a character matrix with length(ids) rows and length(markers) columns.

See Also

getAlleles()

Examples

x = nuclearPed(1)
m1 = marker(x, `2` = "1/2", alleles = 1:2, name = "m1")
m2 = marker(x, `3` = "2/2", alleles = 1:2, name = "m2")
x = setMarkers(x, list(m1, m2))
getGenotypes(x)

### A list of pedigrees

s = transferMarkers(x, singleton("s"))
peds = list(x, s)
getGenotypes(peds)

---

getMap  Tabulate marker positions

Description

Return a map of the markers attached to a pedigree.

Usage

getMap(x, markers = NULL, na.action = 0, merlin = FALSE, verbose = TRUE)

setMap(x, map, matchNames = NA, ...)

hasLinkedMarkers(x)
getMap

Arguments

- **x**: An object of class `ped` or a list of such.
- **markers**: A vector of names or indices referring to markers attached to `x`. By default, all markers are included.
- **na.action**: Either 0 (default), 1 or 2. (See Details.)
- **merlin**: A logical mostly for internal use: If TRUE the function returns a matrix instead of a data frame.
- **verbose**: A logical.
- **map**: Either a data frame or the path to a map file. See Details regarding format.
- **matchNames**: A logical; if TRUE, pre-existing marker names of `x` will be used to assign chromosome labels and positions from `map`.
- **...**: Further arguments passed to `read.table()`.

Details

The `na.action` argument controls how missing values are dealt with:

- `na.action = 0`: Return map unmodified
- `na.action = 1`: Replace missing values with dummy values.
- `na.action = 2`: Remove markers with missing data.

In `setMap()`, the `map` argument should be a data frame (or file) with the following columns in order:

1. chromosome
2. marker name
3. position (Mb)

Column names are ignored, as are any columns after the first three.

Value

- `getMap()` returns a data frame with columns CHROM, MARKER and MB.
- `setMap()` returns `x` with modified marker attributes.
- `hasLinkedMarkers()` returns TRUE if two markers are located (with set position) on the same chromosome, and FALSE otherwise.

Examples

```r
x = singleton(1)
m1 = marker(x, chrom = 1, posMb = 10, name = "m1")
m2 = marker(x, chrom = 1, posMb = 11)
m3 = marker(x, chrom = 1)
x = setMarkers(x, list(m1, m2, m3))

# Compare effect of 'na.action'
getMap(x, na.action = 0)
getMap(x, na.action = 1)
```
getSex(x, na.action = 2)

# Getting and setting map are inverses
y = setMap(x, getMap(x))
stopifnot(identical(x,y))

hasLinkedMarkers(x)

---

getSex

Get or set the sex of pedigree members

Description

Functions for retrieving or changing the sex of specified pedigree members.

Usage

getSex(x, ids = NULL, named = FALSE)

setSex(x, ids = NULL, sex)

swapSex(x, ids, verbose = TRUE)

Arguments

x A ped object or a list of such.
ids A character vector (or coercible to one) containing ID labels. If NULL, defaults to all members of x.
named A logical: return a named vector or not.
sex A numeric vector with entries 1 (= male), 2 (= female) or 0 (= unknown). If ids is NULL, sex must be named with ID labels. If sex is unnamed and shorter than ids it is recycled to length(ids).
verbose A logical: Verbose output or not.

Value

- getSex(x, ids) returns an integer vector of the same length as ids, with entries 0 (unknown), 1 (male) or 2 (female).
- setSex(x, ids, sex) returns a ped object similar to x, but where the sex of ids is set according to the entries of sex
- swapSex(x, ids) returns a ped object identical to x, but where the sex of ids (and their spouses) are swapped (1 <-> 2).

See Also

ped()
Examples

```r
x = nuclearPed(father = "fa", mother = "mo", children = "ch")

stopifnot(all.equal(
    getSex(x, named = TRUE),
    c(fa = 1, mo = 2, ch = 1)
))

# Make child female
setSex(x, ids = "ch", sex = 2)

# Same, using a named vector
setSex(x, sex = c(ch = 2))

# Swapping sex is sometimes easier,
# since spouses are dealt with automatically
swapSex(x, ids = "fa")

# setting/getting sex in a pedlist
y = list(singleton(1, sex = 2), singleton(2), singleton(3))
sx = getSex(y, named = TRUE)
y2 = setSex(y, sex = sx)

stopifnot(identical(y, y2))
```

---

**inbreedingLoops**

**Pedigree loops**

### Description

Functions for identifying, breaking and restoring loops in pedigrees.

### Usage

- `inbreedingLoops(x)`
- `breakLoops(x, loopBreakers = NULL, verbose = TRUE, errorIfFail = TRUE)`
- `tieLoops(x, verbose = TRUE)`
- `findLoopBreakers(x)`
- `findLoopBreakers2(x, errorIfFail = TRUE)`

### Arguments

- `x` a `ped()` object.
loopBreakers   either NULL (resulting in automatic selection of loop breakers) or a numeric containing IDs of individuals to be used as loop breakers.
verbose        a logical: Verbose output or not?
errorIfFail     a logical: If TRUE an error is raised if the loop breaking is unsuccessful. If FALSE, the pedigree is returned unchanged.

Details

Pedigree loops are usually handled (by pedtools and related packages) under the hood - using the functions described here - without need for explicit action from end users. When a ped object x is created, an internal routine detects if the pedigree contains loops, in which case x$UNBROKEN_LOOPS is set to TRUE.

In cases with complex inbreeding, it can be instructive to plot the pedigree after breaking the loops. Duplicated individuals are plotted with appropriate labels (see examples).

The function findLoopBreakers identifies a set of individuals breaking all inbreeding loops, but not marriage loops. These require more machinery for efficient detection, and pedtools does this is a separate function, findLoopBreakers2, utilizing methods from the igraph package. Since this is rarely needed for most users, igraph is not imported when loading pedtools, only when findLoopBreakers2 is called.

In practice, breakLoops first calls findLoopBreakers and breaks at the returned individuals. If the resulting ped object still has loops, findLoopBreakers2 is called to break any marriage loops.

Value

For breakLoops, a ped object in which the indicated loop breakers are duplicated. The returned object will also have a non-null loopBreakers entry, namely a matrix with the IDs of the original loop breakers in the first column and the duplicates in the second. If loop breaking fails, then depending on errorIfFail either an error is raised, or the input pedigree is returned, still containing unbroken loops.

For tieLoops, a ped object in which any duplicated individuals (as given in the x$LOOPBREAKERS entry) are merged. For any ped object x, the call tieLoops(breakLoops(x)) should return x.

For inbreedingLoops, a list containing all inbreeding loops (not marriage loops) found in the pedigree. Each loop is represented as a list with elements top, bottom, pathA (individuals forming a path from top to bottom) and pathB (creating a different path from top to bottom, with no individuals in common with pathA). Note that the number of loops reported here counts all closed paths in the pedigree and will in general be larger than the genus of the underlying graph.

For findLoopBreakers and findLoopBreakers2, a numeric vector of individual ID’s.

Author(s)

Magnus Dehli Vigeland

Examples

x = cousinPed(1, child = TRUE)
plot(breakLoops(x))
is.marker

Test if something is a marker

Description

Functions for testing if something is a marker object, or a list of such objects.

Usage

is.marker(x)

is.markerList(x)

Arguments

x Any object

Value

A logical

is.ped

Is an object a ped object?

Description

Functions for checking whether an object is a ped() object, a singleton() or a list of such.
Usage

is.ped(x)

is.singleton(x)

is.pedList(x)

Arguments

x Any R object.

Details

Note that the singleton class inherits from ped, so if x is a singleton, is.ped(x) returns TRUE.

Value

For is.ped(): TRUE if x is a ped or singleton object, otherwise FALSE.
For is.singleton(): TRUE if x is a singleton object, otherwise FALSE.
For is.pedList(): TRUE if x is a list of ped and/or singleton objects, otherwise FALSE.

Author(s)

Magnus Dehli Vigeland

See Also

ped()

Examples

x1 = nuclearPed(1)
x2 = singleton(1)
stopifnot(is.ped(x1), !is.singleton(x1),
             is.ped(x2), is.singleton(x2),
             is.pedList(list(x1, x2)))

locusAttributes Get or set locus attributes

Description

Retrieve or modify the attributes of attached markers
Usage

getLocusAttributes(
  x,  
  markers = NULL, 
  attribs = c("alleles", "afreq", "name", "chrom", "posMb", "mutmod")
)

setLocusAttributes( 
  x, 
  markers = NULL, 
  locusAttributes, 
  matchNames = NA, 
  erase = FALSE
)

Arguments

x
A ped object, or a list of such.

markers
A character vector (with marker names) or a numeric vector (with marker in-
dices). If NULL (default), the behaviour depends on matchNames, see Details.

attribs
A subset of the character vector c("alleles", "afreq", "name", "chrom", "posMb", "mutmod", "rate").

locusAttributes
A list of lists, with attributes for each marker.

matchNames
A logical, only relevant if markers = NULL. If TRUE, then the markers to be
modified are identified by the 'name' component of each locusAttributes
entry. If FALSE, all markers attached to x are selected in order.

erase
A logical. If TRUE, all previous attributes of the selected markers are erased.
If FALSE, attributes not affected by the submitted locusAttributes remain
untouched.

Details

The default setting markers = NULL select markers automatically, depending on the matchNames
argument. If matchNames = FALSE, all markers are chosen If matchNames = TRUE, markers will be
matched against the name entries in locusAttributes (and an error issued if these are missing).

Note that the default value NA of matchNames is changed to TRUE if all entries of locusAttributes
have a name component which matches the name a an attached marker.

Possible attributes given in locusAttributes are as follows (default values in parenthesis):

- alleles: a character vector with allele labels
- afreq: a numeric vector with allele frequencies(rep.int(1/L, L), where L = length(alleles))
- name: marker name (NA)
- chrom: chromosome number (NA)
- posMb: physical location in megabases (NA)
- mutmod: mutation model, or model name (NULL)
- rate: mutation model parameter (NULL)
Value

- `getLocusAttributes`: a list of lists
- `setLocusAttributes`: a modified version of x.

Examples

```r
x = singleton(1)
x = addMarkers(x, marker(x, name = "m1", alleles = 1:2))
x = addMarkers(x, marker(x, name = "m2", alleles = letters[1:2], chrom = "X"))

# Change frequencies at both loci
y = setLocusAttributes(x, markers = 1:2, loc = list(afreq = c(.1, .9)))
getMarkers(y, 1)

# Set the same mutation model at both loci
z = setLocusAttributes(x, markers = 1:2, loc = list(mutmod = "proportional", rate = .1))
mutmod(z, 1)

# By default, the markers to be modified are identified by name
locs = list(list(name = "m1", alleles = 1:10),
            list(name = "m2", alleles = letters[1:10]))
w = setLocusAttributes(x, loc = locs)
getMarkers(w, 1:2)

# If `erase = TRUE` attributes not explicitly given are erased
w2 = setLocusAttributes(x, loc = locs, erase = TRUE)
chrom(w2, 2) # not "X" anymore

# The getter and setter are inverses
newx = setLocusAttributes(x, loc = getLocusAttributes(x))
stopifnot(identical(x, newx))
```

marker

**Marker objects**

Description

Creating a marker object associated with a pedigree. The function `marker()` returns a marker object, while `addMarker()` first creates the marker and then attaches it to x.

Usage

```r
marker(
  x,
  ...
  geno = NULL,
  allelematrix = NULL,
```
Marker

```r
addMarker(
  x,
  ..., 
  geno = NULL,
  alleleMatrix = NULL,
  alleles = NULL,
  afreq = NULL,
  chrom = NA,
  posMb = NA,
  name = NA,
  NAstrings = c(0, "", NA, "-"),
  mutmod = NULL,
  rate = NULL,
  validate = TRUE
)
```

Arguments

- `x` a `ped` object
- `...` one or more expressions of the form `id = genotype`, where `id` is the ID label of a member of `x`, and `genotype` is a numeric or character vector of length 1 or 2 (see Examples).
- `geno` a character vector of length `pedsize(x)`, with genotypes written in the format "a/b".
- `alleleMatrix` a matrix with 2 columns and `pedsize(x)` rows. If this is non-NULL, then ... must be empty.
- `alleles` a character (or coercible to character) containing allele names. If not given, and `afreq` is named, `names(afreq)` is used. The default action is to take the sorted vector of distinct alleles occurring in `alleleMatrix`, `geno` or ... .
- `afreq` a numeric of the same length as `alleles`, indicating the population frequency of each allele. A warning is issued if the frequencies don’t sum to 1 after rounding to 3 decimals. If the vector is named, and `alleles` is not NULL, an error is raised if `setequal(names(afreq), alleles)` is not TRUE. If `afreq` is not specified, all alleles are given equal frequencies.
- `chrom` a single integer: the chromosome number. Default: NA.
posMb        a nonnegative real number: the physical position of the marker, in megabases. Default: NA.
name         a character string: the name of the marker. Default: NA.
NAstrings    A character vector containing strings to be treated as missing alleles. Default: c("", "0", NA, "+").
mutmod, rate mutation model parameters. These are passed directly to `pedmut::mutationModel()`: see there for details. Note: `mutmod` corresponds to the model parameter. Default: NULL (no mutation model).
validate     if TRUE, the validity of the created marker object is checked.

Value

An object of class marker. This is an integer matrix with 2 columns and one row per individual, and the following attributes:

- alleles (a character vector with allele labels)
- afreq (allele frequencies; default `rep.int(1/length(alleles), length(alleles))`)
- chrom (chromosome number; default = NA)
- posMb (physical location in megabases; default = NA)
- name (marker identifier; default = NA)
- mutmod (a list of two (male and female) mutation matrices; default = NULL)

See Also

`marker_attach`

Examples

```r
x = nuclearPed(father = "fa", mother = "mo", children = "child")

# An empty SNP with alleles "A" and "B"
marker(x, alleles = c("A", "B"))

# Creating and attaching to `x`
addMarker(x, alleles = c("A", "B"))

# Alleles/frequencies can be given jointly or separately
stopifnot(identical(
  marker(x, afreq = c(A = 0.01, B = 0.99)),
  marker(x, alleles = c("A", "B"), afreq = c(0.01, 0.99)),
))

# Genotypes can be assigned individually ...
marker(x, fa = "1/1", mo = "1/2")

# ... or using the `geno` vector (all members in order)
marker(x, geno = c("1/1", "1/2", NA))
```
# Attaching a marker to the pedigree
m = marker(x)  # By default a SNP with alleles 1,2
x = setMarkers(x, m)

# A marker with a "proportional" mutation model,
# with different rates for males and females
mutrates = list(female = 0.1, male = 0.2)
marker(x, alleles = 1:2, mutmod = "prop", rate = mutrates)

---

**marker_attach**

**Attach markers to pedigrees**

### Description

In many applications it is useful to *attach* markers to their associated ped object. In particular for bigger projects with many markers, this makes it easier to manipulate the dataset as a unit. The function `setMarkers()` replaces all existing markers with the supplied ones, while `addMarkers()` appends the supplied markers to any existing ones. Note that there is also the function `addMarker()`, which creates and attaches a single marker in one go.

### Usage

```r
setMarkers(
  x,
  m = NULL,
  alleleMatrix = NULL,
  locusAttributes = NULL,
  missing = 0,
  sep = NULL,
  checkCons = TRUE
)
```

```r
addMarkers(
  x,
  m = NULL,
  alleleMatrix = NULL,
  locusAttributes = NULL,
  missing = 0,
  sep = NULL,
  checkCons = TRUE
)
```

### Arguments

- **x**  
  A ped object

- **m**  
  Either a single marker object or a list of marker objects
alleleMatrix  A matrix with pedsize(x) rows, containing the observed alleles for one or several markers. The matrix must have either 1 or 2 columns per marker. If the former, then a sep string must be a given, and will be used to be split all entries.

locusAttributes  A list of lists, with attributes for each marker. See Details for possible attributes.

missing  A single character (or coercible to one) indicating the symbol for missing alleles.

sep  If this is a single string, each entry of alleleMatrix is interpreted as a genotype, and will be split by calling strsplit(..., split = sep, fixed = TRUE). If alleleMatrix contains entries with "/", this will be taken as separator by default. (To override this behaviour, put sep = FALSE.)

checkCons  A logical. If TRUE (default), each marker is checked for consistency with x.

Details

The most general format of locusAttributes a list of lists, one for each marker, where possible entries in the inner lists are as follows (default values in parenthesis):

- alleles: a character vector with allele labels
- afreq: a numeric vector with allele frequencies (rep.int(1/L, L), where L = length(alleles))
- chrom: chromosome number (NA)
- posMb: physical location in megabases (NA)
- name: marker name (NA)
- mutmod: mutation model, or model name (NULL)
- rate: mutation model parameter (NULL)

If locusAttributes is just a single list of attributes (not a list of lists), then it is repeated to match the number of markers. In particular, the shortcut locusAttributes = "snp-12" sets all markers to be SNPs with alleles 1 and 2.

Two alternative format of locusAttributes are allowed: If a data.frame or matrix is given, an attempt is made to interpret it as a frequency database in allelic ladder format. Such an interpretation is also attempted if locusAttributes is a list of named frequency vectors (where the names are the allele labels).

Value

A ped object.

See Also

addMarker()

Examples

x = singleton(1)
m1 = marker(x, "1" = "1/2")
m2 = marker(x, "1" = "a/b")
# Attach to x
x1 = setMarkers(x, list(m1, m2))

# Reversing the order of the markers
setMarkers(x, list(m2, m1))

# Alternative syntax, adding one marker at a time
x2 = x |> addMarker(`1` = "1/2") |> addMarker(`1` = "a/b")

stopifnot(identical(x1, x2))

---

**marker_getattr**  
*Get marker attributes*

**Description**

S3 methods retrieving marker attributes. They work on single marker objects and markers attached to ped objects (or lists of such).

**Usage**

```r
genotype(x, ...)
## S3 method for class 'marker'
genotype(x, id, ...)

## S3 method for class 'ped'
genotype(x, markers = NULL, id, ...)

mutmod(x, ...)
## S3 method for class 'marker'
mutmod(x, ...)

## S3 method for class 'ped'
mutmod(x, marker, ...)

## S3 method for class 'list'
mutmod(x, marker, ...)

alleles(x, ...)
## S3 method for class 'marker'
alleles(x, ...)

## S3 method for class 'ped'
alleles(x, marker, ...)
```
## S3 method for class 'list'
alleles(x, marker, ...)

afreq(x, ...)

## S3 method for class 'marker'
afreq(x, ...)

## S3 method for class 'ped'
afreq(x, marker, ...)

## S3 method for class 'list'
afreq(x, marker, ...)

name(x, ...)

## S3 method for class 'marker'
name(x, ...)

## S3 method for class 'ped'
name(x, markers = NULL, ...)

## S3 method for class 'list'
name(x, markers = NULL, ...)

chrom(x, ...)

## S3 method for class 'marker'
chrom(x, ...)

## S3 method for class 'ped'
chrom(x, markers = NULL, ...)

## S3 method for class 'list'
chrom(x, markers = NULL, ...)

posMb(x, ...)

## S3 method for class 'marker'
posMb(x, ...)

## S3 method for class 'ped'
posMb(x, markers = NULL, ...)

### Arguments

\textbf{x} \\
Either a \texttt{marker} object, a \texttt{ped} object or a list of \texttt{ped} objects.
Further arguments, not used.

**id**

The ID label of a single pedigree member.

**marker, markers**

The index or name of a marker (or a vector indicating several markers) attached to \( x \).

**Value**

The associated marker attributes.

**See Also**

Setting marker attributes: `marker_setattr` and `marker_inplace`.

**Examples**

```r
x = nuclearPed(1)
x = addMarker(x) # add empty marker

# Inspect default attributes
alleles(x, marker = 1)
afreq(x, marker = 1)
name(x, marker = 1) # NA
chrom(x, marker = 1) # NA
```

**Description**

These S3 methods perform in-place modifications of marker attributes. They work on single marker objects and markers attached to ped objects (or lists of such). Although these functions will continue to exist, we recommend the newer alternatives `setGenotype()`, `setAfreq()` ... in most cases.

**Usage**

```r
genotype(x, ...) <- value

## S3 replacement method for class 'marker'
genotype(x, id, ...) <- value

## S3 replacement method for class 'ped'
genotype(x, marker, id, ...) <- value

mutmod(x, ...) <- value

## S3 replacement method for class 'marker'
```
mutmod(x, ...) <- value

## S3 replacement method for class 'ped'
mutmod(x, marker = NULL, ...) <- value

## S3 replacement method for class 'list'
mutmod(x, marker = NULL, ...) <- value

afreq(x, ...) <- value

## S3 replacement method for class 'marker'
afreq(x, ...) <- value

## S3 replacement method for class 'ped'
afreq(x, marker, ...) <- value

## S3 replacement method for class 'list'
afreq(x, marker, ...) <- value

name(x, ...) <- value

## S3 replacement method for class 'marker'
name(x, ...) <- value

## S3 replacement method for class 'ped'
name(x, markers = NULL, ...) <- value

## S3 replacement method for class 'list'
name(x, markers = NULL, ...) <- value

chrom(x, ...) <- value

## S3 replacement method for class 'marker'
chrom(x, ...) <- value

## S3 replacement method for class 'ped'
chrom(x, markers = NULL, ...) <- value

## S3 replacement method for class 'list'
chrom(x, markers = NULL, ...) <- value

posMb(x, ...) <- value

## S3 replacement method for class 'marker'
posMb(x, ...) <- value

## S3 replacement method for class 'ped'
posMb(x, markers = NULL, ...) <- value
marker_prop

Arguments

- **x**: Either a marker object, a ped object or a list of ped objects.
- **...**: Further arguments, not used.
- **value**: Replacement value(s).
- **id**: The ID label of a single pedigree member.
- **marker, markers**: The index or name of a marker (or a vector indicating several markers) attached to ped. Used if x is a ped object.

Value

These functions perform in-place modification of x.

See Also


Examples

```r
x = nuclearPed(1)
x = addMarker(x, alleles = 1:2)

# Set genotypes
genotype(x, marker = 1, id = 1) = "1/2"

# Set marker name
name(x, 1) = "M"

# Change allele freqs
afreq(x, "M") = c("1" = 0.1, "2" = 0.9)

# Set position
chrom(x, "M") = 1
posMb(x, "M") = 123.45

# Check result
m = marker(x, "1" = "1/2", name = "M", afreq = c("1" = 0.1, "2" = 0.9),
            chrom = 1, posMb = 123.45)
stopifnot(identical(x$MARKERS[[1]], m))
```

---

**Description**

These functions are used to retrieve various properties of marker objects. Each function accepts as input either a single marker object, a ped object, or a list of ped objects.
Usage

emptyMarker(x, ...)

## Default S3 method:
emptyMarker(x, ...)

## S3 method for class 'marker'
emptyMarker(x, ...)

## S3 method for class 'ped'
emptyMarker(x, markers = NULL, ...)

## S3 method for class 'list'
emptyMarker(x, markers = NULL, ...)

nTyped(x, ...)

## Default S3 method:
nTyped(x, ...)

## S3 method for class 'marker'
nTyped(x, ...)

## S3 method for class 'ped'
nTyped(x, markers = NULL, ...)

## S3 method for class 'list'
nTyped(x, markers = NULL, ...)

nAlleles(x, ...)

## Default S3 method:
nAlleles(x, ...)

## S3 method for class 'marker'
nAlleles(x, ...)

## S3 method for class 'ped'
nAlleles(x, markers = NULL, ...)

## S3 method for class 'list'
nAlleles(x, markers = NULL, ...)

isXmarker(x, ...)

## Default S3 method:
isXmarker(x, ...)
## S3 method for class 'marker'
isXmarker(x, ...)

## S3 method for class 'ped'
isXmarker(x, markers = NULL, ...)

## S3 method for class 'list'
isXmarker(x, markers = NULL, ...)

allowsMutations(x, ...)

## Default S3 method:
al lowsMutations(x, ...)

## S3 method for class 'marker'
allowsMutations(x, ...)

## S3 method for class 'ped'
al lowsMutations(x, markers = NULL, ...)

## S3 method for class 'list'
al lowsMutations(x, markers = NULL, ...)

**Arguments**

x
A single marker object or a ped object (or a list of such)

... Not used.

markers A vector of names or indices of markers attached to x. By default all attached markers are selected.

**Details**

e mptyMarker() returns TRUE for markers with no genotypes. If the input is a list of pedigrees, all must be empty for the result to be TRUE.

nTyped() returns the number of typed individuals for each marker. Note that if the input is a list of pedigrees, the function returns the sum over all components.

nAlleles() returns the number of alleles of each marker.

isXmarker() returns TRUE for markers whose chrom attribute is either "X" or 23.
al lowsMutations returns TRUE for markers whose mutmod attribute is non-NULL and differs from the identity matrix.

**Value**

If x is a single marker object, the output is a vector of length 1.

Otherwise, a vector of length nMarkers(x) (default) or length(markers), reporting the property of each marker.
Examples

cmp1 = nuclearPed(1)
cmp2 = singleton(10)
loc = list(alleles = 1:2)
x = setMarkers(list(cmp1, cmp2), locus = rep(list(loc), 3))

-------- nAlleles() ------------
# All markers have 2 alleles
stopifnot(identical(nAlleles(x), c(2L,2L,2L)))

-------- emptyMarkers() ------------
# Add genotype for indiv 1 at marker 1
genotype(x[[1]], 1, 1) = "1/2"

# Check that markers 2 and 3 are empty
stopifnot(identical(emptyMarker(x), c(FALSE,TRUE,TRUE)),
          identical(emptyMarker(x[[1]]), c(FALSE,TRUE,TRUE)),
          identical(emptyMarker(x[[2]]), c(TRUE,TRUE,TRUE)),
          identical(emptyMarker(x, markers = c(3,1)), c(TRUE,FALSE)))

-------- nTyped() ------------
stopifnot(identical(nTyped(x), c(1L,0L,0L)))

# Add genotypes for third marker
genotype(x[[1]], marker = 3, id = 1:3) = "1/1"
genotype(x[[2]], marker = 3, id = 10) = "2/2"

# nTyped() returns total over all components
stopifnot(identical(nTyped(x), c(1L,0L,4L)))

-------- allowsMutations() ------------
# Marker 2 allows mutations
mutmod(x, 2) = list("prop", rate = 0.1)
stopifnot(identical(allowsMutations(x), c(FALSE,TRUE,FALSE)),
          identical(allowsMutations(x, markers = 2:3), c(TRUE,TRUE,FALSE)))

-------- isXmarker() ------------
# Make marker 3 X-linked
chrom(x[[1]], 3) = "X"
chrom(x[[2]], 3) = "X"

stopifnot(identical(isXmarker(x), c(FALSE,FALSE,TRUE)))

marker_select
Select or remove attached markers

Description

Functions for manipulating markers attached to ped objects.
Usage

```
selectMarkers(x, markers = NULL, chroms = NULL, fromPos = NULL, toPos = NULL)
getMarkers(x, markers = NULL, chroms = NULL, fromPos = NULL, toPos = NULL)
removeMarkers(x, markers = NULL, chroms = NULL, fromPos = NULL, toPos = NULL)
whichMarkers(x, markers = NULL, chroms = NULL, fromPos = NULL, toPos = NULL)
```

Arguments

- **x**: A ped object, or a list of such
- **markers**: Either a character vector (with marker names), a numeric vector (with marker indices), a logical (of length `nMarkers(x)`), or NULL.
- **chroms**: A vector of chromosome names, or NULL
- **fromPos**: A single number or NULL
- **toPos**: A single number or NULL

Details

If `markers` consists of negative integers, it will be converted to its complement within `1:nMarkers(x)`.

Value

The return values of these functions are:

- **selectMarkers()**: an object identical to `x`, but where only the indicated markers are kept
- **removeMarkers()**: an object identical to `x`, but where the indicated markers are removed
- **getMarkers()**: a list of marker objects. Note: If `x` is a list of pedigrees, the marker objects attached to the first component will be returned.
- **whichMarkers()**: an integer vector with indices of the indicated markers. If `x` is a list of pedigrees an error is raised unless `whichMarkers()` gives the same result for all components.

See Also

- `setMarkers()`
marker_setattr  

*Set marker attributes*

**Description**

These functions set or modify various attributes of markers attached to a pedigree. They are sometimes more convenient (and pipe-friendly) than the in-place modifiers described in `marker_inplace`.

**Usage**

```r
setGenotype(x, marker = NULL, id, geno)
setAfreq(x, marker, afreq, strict = TRUE)
setMarkername(x, marker = NULL, name)
setChrom(x, marker = NULL, chrom)
setPosition(x, marker = NULL, posMb)
```

**Arguments**

- **x**: A ped object or a list of ped objects.
- **marker**: A vector of indices or names of one or several markers attached to x.
- **id**: The ID label of a single pedigree member.
- **geno**: a character vector of length `pedsize(x)`, with genotypes written in the format "a/b".
- **afreq**: a numeric of the same length as alleles, indicating the population frequency of each allele. A warning is issued if the frequencies don’t sum to 1 after rounding to 3 decimals. If the vector is named, and alleles is not NULL, an error is raised if `setequal(names(afreq), alleles)` is not TRUE. If `afreq` is not specified, all alleles are given equal frequencies.
- **strict**: A logical. If TRUE (default) the new frequencies cannot remove or add any alleles.
- **name**: A character of the same length as marker, containing marker names.
- **chrom**: A character of the same length as marker, containing chromosome labels.
- **posMb**: A numeric of the same length as marker, containing the physical marker positions in megabases (or NA).

**Value**

A copy of x with modified attributes.
Examples

```r
x = nuclearPed(1) |> addMarker(alleles = 1:2) |> setMarkernname(marker = 1, name = "M") |> setGenotype(marker = "M", id = 1, geno = "1/2") |> setAfreq(marker = "M", afreq = c("1" = 0.1, "2" = 0.9)) |> setChrom(marker = "M", chrom = 1) |> setPosition(marker = "M", posMb = 123.45)
```

# Of course, all of this could have been done on creation:
y = addMarker(nuclearPed(), "1" = "1/2", afreq = c("1" = 0.1, "2" = 0.9), name = "M", chrom = 1, posMb = 123.45)
stopifnot(identical(x, y))

---

mendelianCheck

Check for Mendelian errors

Description

Check marker data for Mendelian inconsistencies

Usage

`mendelianCheck(x, remove = FALSE, verbose = !remove)`

Arguments

- `x`: a `ped()` object
- `remove`: a logical. If FALSE, the function returns the indices of markers found to incorrect. If TRUE, a new `ped` object is returned, where the incorrect markers have been deleted.
- `verbose`: a logical. If TRUE, details of the markers failing the tests are shown.

Value

A numeric containing the indices of the markers that did not pass all tests, or (if `remove` = TRUE) a new `ped` object where the failing markers are removed.

Author(s)

Magnus Dehli Vigeland
Examples

```r
x = nuclearPed()

# Add a SNP with Mendelian error
m = marker(x, '1' = "1/1", '2' = "1/1", '3' = "1/2")
x = setMarkers(x, m)
mendelianCheck(x)
```

mergePed  

**Merge two pedigrees**

Description

This function merges two ped objects, joining them at the indicated individuals. Only ped objects without marker data are supported.

Usage

```r
mergePed(x, y, by = NULL, relabel = FALSE, ...)
```

Arguments

- `x`, `y`  
  ped() objects
- `by`  
  The individuals to merge by. The most general form uses a named vector with entries of the form `id.x = id.y` (see Examples). If the vector is unnamed, it is assumed that the merging individuals have the same labels in both pedigrees. Finally, if `by = NULL` (default), it is set to `intersect(labels(x), labels(y))`.
- `relabel`  
  A logical, by default FALSE. If TRUE, `relabel(..., "asPlot")` is run on the merged pedigree before returning.
- `...`  
  further arguments passed along to ped(), e.g. famid, validate and reorder.

Details

Some internal checks are done to ensure that merging individuals have the same sex and the same parents.

If `relabel = FALSE`, some relabelling might still be performed in order to ensure unique labels for everyone. Specifically, this is the case if some ID labels occur in both `x` and `y` other than those given in the by argument. In such cases, the relevant members of `y` get a suffix `.y`.

Value

A ped object.
Examples

############
# Example 1
# A family trio where each parent have first cousin parents.
############

# Trio
x = nuclearPed(1)

# Add paternal family
pat = cousinPed(1, child = TRUE)
x = mergePed(x, pat, by = c("1" = "9"))

# Maternal family
mat = cousinPed(1, child = TRUE) |> swapSex("9")
x = mergePed(x, mat, by = c("2" = "9"))

# Relabel (Alternative: add `relabel = TRUE` in the previous call)
x = relabel(x, "asPlot")
plot(x)

##################################
# Example 2: Double first cousins
##################################

# First cousins, whose fathers are brothers
y = cousinPed(degree = 1)

# Create two sisters
motherPed = nuclearPed(2, sex = 2)

# Plot to see who is who: `plotPedList(list(y, motherPed))`

# Merge
z = mergePed(y, motherPed, by = c("4" = 3, "6" = 4), relabel = TRUE)
plot(z)
Description

This is the internal constructor of marker objects. It does not do any input validation and should only be used in programming scenarios, and only if you know what you are doing. Most users are recommended to use the regular constructor `marker()`.

Usage

```r
newMarker(
    alleleMatrixInt,
    alleles,
    afreq,
    name = NA_character_,
    chrom = NA_character_,
    posMb = NA_real_,
    mutmod = NULL,
    pedmembers,
    sex
)
```

Arguments

- `alleleMatrixInt`: An integer matrix.
- `alleles`: A character vector.
- `afreq`: A numeric vector.
- `name`: A character of length 1.
- `chrom`: A character of length 1.
- `posMb`: A numeric of length 1.
- `mutmod`: A mutation model.
- `pedmembers`: A character vector.
- `sex`: An integer vector.

Details

See `marker()` for more details about the marker attributes.

Value

A marker object.

Examples

```r
newMarker(matrix(c(1L, 0L, 1L, 1L, 0L, 2L), ncol = 2),
          alleles = c("A", "B"), afreq = c(0.1, 0.9), name = "M",
          pedmembers = c("1", "2", "3"), sex = c(1L, 2L, 1L))
```
newPed

**Internal ped constructor**

**Description**

This is the internal constructor of ped objects. It does not do any validation of input other than simple type checking. In particular it should only be used in programming scenarios where it is known that the input is a valid, connected pedigree. End users are recommended to use the regular constructor `ped()`.

**Usage**

```
newPed(ID, FIDX, MIDX, SEX, FAMID)
```

**Arguments**

- **ID**: A character vector.
- **FIDX**: An integer vector.
- **MIDX**: An integer vector.
- **SEX**: An integer vector.
- **FAMID**: A string.

**Details**

See `ped()` for details about the input parameters.

**Value**

A ped object.

**Examples**

```
newPed("a", 0L, 0L, 1L, ")
```
nMarkers

The number of markers attached to a pedigree

Description
The number of markers attached to a pedigree

Usage
nMarkers(x)

hasMarkers(x)

Arguments
x
A ped object or a list of such (see Value).

Value
The function nMarkers returns the number of marker objects attached to x. If x is a list of pedigrees, an error is raised unless all of them have the same number of markers.
The function hasMarkers returns TRUE if nMarkers(x) > 0.

ped

Pedigree construction

Description
This is the basic constructor of ped objects. Utility functions for creating many common pedigree structures are described in ped_basic.

Usage
ped(  
id,  
fid,  
mid,  
sex,  
famid = "",  
reorder = TRUE,  
validate = TRUE,  
isConnected = FALSE,  
verbose = FALSE  
)  

singleton(id = 1, sex = 1, famid = "")
**Arguments**

- `id`: a vector (numeric or character) of individual ID labels.
- `fid`: a vector of the same length as `id`, containing the labels of the fathers. In other words, `fid[i]` is the father of `id[i]`, or 0 if `id[i]` is a founder.
- `mid`: a vector of the same length as `id`, containing the labels of the mothers. In other words, `mid[i]` is the mother of `id[i]`, or 0 if `id[i]` is a founder.
- `sex`: a numeric of the same length as `id`, describing the genders of the individuals (in the same order as `id`). Each entry must be either 1 (=male), 2 (=female) or 0 (=unknown).
- `reorder`: a logical. If TRUE, the pedigree is reordered so that all parents precede their children.
- `validate`: a logical. If TRUE, `validatePed()` is run before returning the pedigree.
- `isConnected`: a logical, by default FALSE. If it is known that the input is connected, setting this to TRUE speeds up the processing.
- `verbose`: a logical.

**Details**

A singleton is a special `ped` object whose pedigree contains 1 individual. The class attribute of a singleton is `c('singleton', 'ped')`.

Selfing, i.e. the presence of pedigree members whose father and mother are the same individual, is allowed in `ped` objects. Any such "self-fertilizing" parent must have undecided sex (`sex = 0`).

If the pedigree is disconnected, it is split into its connected components and returned as a list of `ped` objects.

**Value**

A `ped` object, which is essentially a list with the following entries:

- `ID`: A character vector of ID labels. Unless the pedigree is reordered during creation, this equals `as.character(id)`.
- `FIDX`: An integer vector with paternal indices: For each $j = 1, 2, \ldots$, the entry `FIDX[j]` is 0 if `ID[j]` has no father within the pedigree; otherwise, `ID[FIDX[j]]` is the father of `ID[j]`.
- `MIDX`: An integer vector with maternal indices: For each $j = 1, 2, \ldots$, the entry `MIDX[j]` is 0 if `ID[j]` has no mother within the pedigree; otherwise, `ID[MIDX[j]]` is the mother of `ID[j]`.
- `SEX`: An integer vector with gender codes. Unless the pedigree is reordered, this equals `as.integer(sex)`.
- `FAMID`: The family ID.
- `UNBROKEN_LOOPS`: A logical: TRUE if the pedigree is inbred.
- `LOOP_BREAKERS`: A matrix with loop breaker ID's in the first column and their duplicates in the second column. All entries refer to the internal IDs. This is usually set by `breakLoops()`.
- `FOUNDER_INBREEDING`: A list of two potential entries, "autosomal" and "x"; both numeric vectors with the same length as `founders(x)`. `FOUNDER_INBREEDING` is always NULL when a new `ped` is created. See `founderInbreeding()`.
- `MARKERS`: A list of marker objects, or NULL.
Author(s)
Magnus Dehli Vigeland

See Also
ped_basic, ped_modify, ped_subgroups, relabel()

Examples

# Trio
x = ped(id = 1:3, fid = c(0,0,1), mid = c(0,0,2), sex = c(1,2,1))

# Female singleton
y = singleton(‘NN’, sex = 2)

# Selfing
z = ped(id = 1:2, fid = 0:1, mid = 0:1, sex = 0:1)
stopifnot(hasSelfing(z))

# Disconnected pedigree: Trio + singleton
w = ped(id = 1:4, fid = c(2,0,0,0), mid = c(3,0,0,0), sex = c(1,1,2,1))
stopifnot(is.pedList(w), length(w) == 2)

pedtools: Tools for working with pedigrees in R

Description
A comprehensive collection of tools for creating, manipulating and visualising pedigrees and genetic marker data. Pedigrees can be read from text files or created on the fly with built-in functions. A range of utilities enable modifications like adding or removing individuals, breaking loops, and merging pedigrees. Pedigree plots are produced by wrapping the plotting functionality of the kinship2 package. A Shiny app for creating pedigrees, based on pedtools, is available at https://magnusdv.shinyapps.io/quickped. pedtools is the hub of the ped suite, a collection of packages for pedigree analysis. A detailed presentation of the ped suite is given in the book Pedigree Analysis in R (Vigeland, 2021, ISBN:9780128244302).

ped_basic

Create simple pedigrees

Description
Utility functions for creating some common pedigree structures.
Usage

nuclearPed(nch = 1, sex = 1, father = "1", mother = "2", children = NULL)

halfSibPed(
    nch1 = 1,
    nch2 = 1,
    sex1 = 1,
    sex2 = 1,
    type = c("paternal", "maternal")
)

linearPed(n, sex = 1)

cousinPed(
    degree,
    removal = 0,
    side = c("right", "left"),
    half = FALSE,
    child = FALSE
)

avuncularPed(
    top = c("uncle", "aunt"),
    bottom = c("nephew", "niece"),
    side = c("right", "left"),
    removal = 1,
    half = FALSE
)

halfCousinPed(degree, removal = 0, side = c("right", "left"), child = FALSE)

ancestralPed(g)

selfingPed(s, sex = 1)

Arguments

nch  The number of children, by default 1. If children is not NULL, nch is set to length(children)

sex  A vector with integer gender codes (0=unknown, 1=male, 2=female). In nuclearPed(), it contains the genders of the children and is recycled (if necessary) to length nch. In linearPed() it also contains the genders of the children (1 in each generation) and should have length at most n (recycled if shorter than this). In selfingPed() it should be a single number, indicating the gender of the last individual (the others must necessarily have gender code 0.)

father  The label of the father. Default: "1".

mother  The label of the mother. Default: "2".
children A character with labels of the children. Default: "3", "4", ...

nch1, nch2 The number of children in each sibship.

sex1, sex2 Vectors of gender codes for the children in each sibship. Recycled (if necessary) to lengths nch1 and nch2 respectively.

type Either "paternal" or "maternal".

n The number of generations, not including the initial founders.

degree A non-negative integer: 0=siblings, 1=first cousins; 2=second cousins, a.s.o.

removal A non-negative integer. See Details and Examples.

side Either "right" or "left"; the side on which removals should be added.

half A logical indicating if the relationship should be "half-like". Default: FALSE.

child A logical: Should an inbred child be added to the two bottom individuals?

top, bottom Words indicating the gender combination in avuncular relationships. The first must be either "uncle" or "aunt", while the second is "nephew" or "niece". Both can be abbreviated.

g A nonnegative integer indicating the number of ancestral generations to include. The resulting pedigree has \(2^{(g+1)}-1\) members. The case \(g = 0\) results in a singleton.

s A nonnegative integer indicating the number of consecutive selfings. The case \(s = 0\) results in a singleton.

Details

halfSibPed(nch1, nch2) produces a pedigree containing two sibships (of sizes nch1 and nch2) with the same father, but different mothers. If maternal half sibs are wanted instead, add type = "maternal".

cousinPed(degree = n, removal = k) creates a pedigree with two \(n\)'th cousins, \(k\) times removed. By default, removals are added on the right side, but this can be changed by adding side = left.

halfCousinPed(...) is a synonym for cousinPed(..., half = TRUE).

avuncularPed() creates uncle/aunt - nephew/niece pedigrees. The empty call avuncularPed() is equivalent to avuncularPed("uncle", "nephew"). Note that the arguments can be abbreviated, so that e.g. avuncularPed("a", "ni") produces an aunt-niece relationship. Grand (and great-grand etc) uncles/aunts can be produced by specifying removal greater than 1.

ancestralPed(g) returns the family tree of a single individual, including all ancestors \(g\) generations back.

selfingPed(s) returns a line of \(s\) consecutive selfings.

Value

A ped object.

See Also

ped(), singleton(), ped_complex, ped_subgroups
Examples

# A nuclear family with 2 boys and 3 girls
nuclearPed(5, sex = c(1, 1, 2, 2, 2))

# A straight line of females
linearPed(3, sex = 2)

# Paternal half brothers
halfSibPed()

# Maternal half sisters
halfSibPed(sex1 = 2, sex2 = 2, type = "maternal")

# Larger half sibships: boy and girl on one side; 3 girls on the other
halfSibPed(nch1 = 2, sex = 1:2, nch2 = 3, sex2 = 2)

# Grand aunt:
cousinPed(degree = 0, removal = 2)

# Second cousins once removed.
cousinPed(degree = 2, removal = 1)

# Same, but with the 'removal' on the left side.
cousinPed(2, 1, side = "left")

# A child of half first cousins.
halfCousinPed(degree = 1, child = TRUE)

# The 'family tree' of a person
ancestralPed(g = 2)

Description

Functions for creating a selection of pedigrees that are awkward to construct from scratch or with the simple structures described in ped_basic.

Usage

doubleCousins(
degree1,
degree2,
removal1 = 0,
removal2 = 0,
half1 = FALSE,
half2 = FALSE,
child = FALSE
)

doubleFirstCousins()
quadHalfFirstCousins()
fullSibMating(n)
halfSibStack(n)
halfSibTriangle(g)

Arguments

degree1, degree2, removal1, removal2
Nonnegative integers.
half1, half2 Logical, indicating if the fathers (resp. mothers) should be full or half cousins.
child A logical: Should a child be added to the double cousins?
n A positive integer indicating the number of crossings.
g A positive integer; the number of generations.

Details

The function doubleCousins returns a pedigree linking two individuals who are simultaneous paternal and maternal cousins. More precisely, they are:

• paternal (full or half) cousins of type (degree1, removal1)
• maternal (full or half) cousins of type (degree2, removal2).

For convenience, a wrapper doubleFirstCousins is provided for the most common case, double first cousins.
quadHalfFirstCousins produces a pedigree with quadruple half first cousins.
fullSibMating crosses full sibs consecutively n times.
halfSibStack produces a breeding scheme where the two individuals in the final generation are simultaneous half k'th cousins, for each k = 0, ..., n-1.
halfSibTriangle produces a triangular pedigree in which every pair of parents are half siblings.

Value

A ped object.

See Also

ped_basic
Examples

# Consecutive brother-sister matings.
x = fullSibMating(2)
# plot(x)

# Simultaneous half siblings and half first cousins
x = halfSibStack(2)
# plot(x)

# Double first cousins
x = doubleFirstCousins()
# plot(x)

# Quadruple half first cousins
x = quadHalffirstCousins()
# plot(x) # Weird plotting behaviour for this pedigree.

# Triangular half-sib pattern
x = halfSibTriangle(4)
# plot(x)

Description

These functions give access to - and enable modifications of - the order in which the members of a pedigree are stored. (This is the order in which the members are listed when a ped object is printed to the screen.)

Usage

reorderPed(x, neworder = NULL)

parentsBeforeChildren(x)

hasParentsBeforeChildren(x)

foundersFirst(x)

internalID(x, ids, errorIfUnknown = TRUE)

Arguments

x A ped object. Most of these functions also accepts ped lists.
neworder A permutation of labels(x) or of vector 1:pedsize(x). By default, the sorting order of the ID labels is used.

ids A character vector (or coercible to one) of original ID labels.

errorIfUnknown A logical. If TRUE (default), the function stops with an error if not all elements of ids are recognised as names of members in x.

Details

The internal ordering is usually of little importance for end users, with one important exception: Certain pedigree-traversing algorithms require parents to precede their children. A special function, parentsBeforeChildren() is provided for this purpose. This is a wrapper of the more general reorderPed() which allows any permutation of the members.

It should be noted that ped() by default calls parentsBeforeChildren() whenever a pedigree is created, unless explicitly avoided with reorder = FALSE.

hasParentsBeforeChildren() can be used as a quick test to decide if it is necessary to call parentsBeforeChildren().

The foundersFirst() function reorders the pedigree so that all the founders come first.

The utility internalID() converts ID labels to indices in the internal ordering. If x is a list of pedigrees, the output is a data frame containing both the component number and internal ID (within the component).

See Also

ped()

Examples

x = ped(id = 3:1, fid = c(1,0,0), mid = c(2,0,0), sex = c(1,2,1), reorder = FALSE)
x

# The 'ids' argument is converted to character, hence these are equivalent:
internalID(x, ids = 3)
internalID(x, ids = "3")

hasParentsBeforeChildren(x)

# Fix the ordering
y = parentsBeforeChildren(x)
internalID(y, ids = 3)

# A different ordering
reorderPed(x, c(2,1,3))
Description

Functions for adding or removing individuals in a 'ped' object.

Usage

```r
addChildren(
  x,
  father = NULL,
  mother = NULL,
  nch = NULL,
  sex = 1,
  ids = NULL,
  verbose = TRUE
)
```

```r
addSon(x, parents, id = NULL, verbose = TRUE, parent = NULL)
```

```r
addDaughter(x, parents, id = NULL, verbose = TRUE, parent = NULL)
```

```r
addParents(x, id, father = NULL, mother = NULL, verbose = TRUE)
```

```r
removeIndividuals(x, ids, verbose = TRUE)
```

```r
branch(x, id)
```

```r
## S3 method for class 'ped'
subset(x, subset, ...)
```

Arguments

- `x`: A ped object.
- `father, mother`: Single ID labels. At least one of these must belong to an existing pedigree member. The other label may either: 1) belong to an existing member, 2) not belong to any existing member, or 3) be missing (i.e. not included in the function call). In cases 2 and 3 a new founder is added to the pedigree. In case 2 its label is the one given, while in case 3 a suitable label is created by the program (see Details).
- `nch`: A positive integer indicating the number of children to be created. Default: 1.
- `sex`: Gender codes of the created children (recycled if needed).
- `ids`: A character vector (or coercible to such) with ID labels. In `addChildren` the (optional) `ids` argument is used to specify labels for the created children. If given, its length must equal `nch`. If not given, labels are assigned automatically as explained in Details.
verbose       A logical: Verbose output or not.
parents       A vector of 1 or 2 ID labels, of which at least one must be an existing member of x.
id            The ID label of some existing pedigree member.
parent         Deprecated; renamed to parents.
subset       A character vector (or coercible to such) with ID labels forming a connected sub-pedigree.
...            Not used.

Details

In addChildren() and addParents(), labels of added individuals are created automatically if they are not specified by the user. In the automatic case, the labelling depends on whether the existing labels are integer-like or not (i.e. if `labels(x)` equals `as.character(as.integer(labels(x))))`.
If so, the new labels are integers subsequent to the largest of the existing labels. If not, the new labels are "NN_1", "NN_2", ... If any such label already exists, the numbers are adjusted accordingly.

addSon() and addDaughter() are wrappers for the most common use of addChildren(), namely adding a single child to a pedigree. Note that the parents can be given in any order. If only one parent is supplied, the other is created as a new individual.

In removeIndividuals() all descendants of ids are also removed. Any individuals (spouses) left unconnected to the remaining pedigree are also removed.

The branch() function extracts the sub-pedigree formed by id and all his/her spouses and descendants.

Finally, subset() can be used to extract any connected sub-pedigree. (Note that in the current implementation, the function does not actually check that the indicated subset forms a connected pedigree; failing to comply with this may lead to obscure errors.)

Value

The modified ped object.

See Also

ped(), relabel(), swapSex()

Examples

```r
x = nuclearPed(1)
# To see the effect of each command below, use plot(x) in between.
x = addSon(x, 3)
x = addParents(x, id = 4, father = 6, mother = 7)
x = removeIndividuals(x, 4)
```
**Description**

A collection of utility functions for identifying pedigree members with certain properties.

**Usage**

```r
founders(x, internal = FALSE)
nonfounders(x, internal = FALSE)
leaves(x, internal = FALSE)
males(x, internal = FALSE)
females(x, internal = FALSE)
typedMembers(x, internal = FALSE)
untypedMembers(x, internal = FALSE)
father(x, id, internal = FALSE)
mother(x, id, internal = FALSE)
children(x, id, internal = FALSE)
offspring(x, id, internal = FALSE)
spouses(x, id, internal = FALSE)
unrelated(x, id, internal = FALSE)
parents(x, id, internal = FALSE)
grandparents(x, id, degree = 2, internal = FALSE)
siblings(x, id, half = NA, internal = FALSE)
nephews_nieces(x, id, removal = 1, half = NA, internal = FALSE)
ancestors(x, id, inclusive = FALSE, internal = FALSE)
commonAncestors(x, ids, inclusive = FALSE, internal = FALSE)
```
descendants(x, id, inclusive = FALSE, internal = FALSE)

commonDescendants(x, ids, inclusive = FALSE, internal = FALSE)

descentPaths(x, ids = founders(x), internal = FALSE)

Arguments

x A ped() object or a list of such.
internal A logical indicating whether id (or ids) refers to the internal order.
id, ids A character (or coercible to such) with one or several ID labels.
degree, removal Non-negative integers.
half a logical or NA. If TRUE (resp. FALSE), only half (resp. full) siblings/cousins/nephews/nieces are returned. If NA, both categories are included.
inclusive A logical indicating whether an individual should be counted among his or her own ancestors/descendants

Value

The functions founders, nonfounders, males, females, leaves each return a vector containing the IDs of all pedigree members with the wanted property. (Recall that a founder is a member without parents in the pedigree, and that a leaf is a member without children in the pedigree.)

The functions father, mother, cousins, grandparents, nephews_nieces, children, parents, siblings, spouses, unrelated, each returns a vector containing the IDs of all pedigree members having the specified relationship with id.

The commands ancestors(x, id) and descendants(x, id) return vectors containing the IDs of all ancestors (resp. descendants) of the individual id within the pedigree x. If inclusive = TRUE, id is included in the output, otherwise not.

For commonAncestors(x, ids) and commonDescendants(x, ids), the output is a vector containing the IDs of common ancestors (descendants) to all of ids.

Finally, descentPaths(x, ids) returns a list of lists, containing all pedigree paths descending from each individual in ids (by default all founders).

Author(s)

Magnus Dehli Vigeland

Examples

x = ped(id = 2:9,
    fid = c(0,0,2,0,4,4,0,2),
    mid = c(0,0,3,0,5,5,0,8),
    sex = c(1,2,1,2,1,2,2,2))

spouses(x, id = 2) # 3, 8
children(x, 2) # 4, 9
descendants(x, 2) # 4, 6, 7, 9
siblings(x, 4) # 9 (full or half)
unrelated(x, 4) # 5, 8
father(x, 4) # 2
mother(x, 4) # 3

siblings(x, 4, half = FALSE) # none
siblings(x, 4, half = TRUE) # 9

leaves(x) # 6, 7, 9
founders(x) # 2, 3, 5, 8

---

**ped_utils**  
**Pedigree utilities**

**Description**
Various utility functions for ped objects.

**Usage**

- pedsize(x)
- generations(x, maxOnly = TRUE, maxComp = TRUE)
- hasUnbrokenLoops(x)
- hasInbredFounders(x, chromType = "autosomal")
- hasSelfing(x)
- hasCommonAncestor(x)
- subnucs(x)
- peelingOrder(x)

**Arguments**

- **x**: A ped object, or (in some functions - see Details) a list of such.
- **maxOnly**: A logical, by default TRUE. (See Value.)
- **maxComp**: A logical, by default TRUE. (See Value.)
- **chromType**: Either "autosomal" (default) or "x".
Value

• `pedsize(x)` returns the number of pedigree members in each component of `x`.

• `generations(x)` by default returns the number of generations in `x`, defined as the number of individuals in the longest line of parent-child links. (Note that this definition is valid also if `x` has loops.) If `maxOnly = FALSE`, the output is a named integer vector, showing the generation number of each pedigree member. If `x` has multiple components, the output depends on the parameter `maxComp`. If this is `FALSE`, the output is a vector containing the result for each component. If `TRUE` (default), only the highest number is returned.

• `hasUnbrokenLoops(x)` returns TRUE if `x` has loops, otherwise FALSE. (No computation is done here; the function simply returns the value of `x$UNBROKEN_LOOPS`).

• `hasInbredFounders(x)` returns TRUE is founder inbreeding is specified for `x` and at least one founder has positive inbreeding coefficient. See `founderInbreeding()` for details.

• `hasSelfing(x)` returns TRUE if the pedigree contains selfing events. This is recognised by father and mother begin equal for some child. (Note that for this to be allowed, the gender code of the parent must be 0.)

• `hasCommonAncestor(x)` computes a logical matrix `A` whose entry `A[i,j]` is TRUE if pedigree members `i` and `j` have a common ancestor in `x`, and FALSE otherwise. By convention, `A[i,i]` is TRUE for all `i`.

• `subnucs(x)` returns a list of all nuclear sub-pedigrees of `x`, wrapped as nucleus objects. Each nucleus is a list with entries `father`, `mother` and `children`.

• `peelingOrder(x)` calls `subnucs(x)` and extends each entry with a `link` individual, indicating a member linking the nucleus to the remaining pedigree. One application of this function is the fact that if `fails` to find a complete peeling order if and only if the pedigree has loops. (In fact it is called each time a new `ped` object is created by `ped()` in order to detect loops.) The main purpose of the function, however, is to prepare for probability calculations in other packages, as e.g. in pedprobr::likelihood.

Examples

```r
x = fullSibMating(1)
stopifnot(pedsize(x) == 6)
stopifnot(hasUnbrokenLoops(x))
stopifnot(generations(x) == 3)

# All members have common ancestors except the grandparents
CA = hasCommonAncestor(x)
stopifnot(!CA[1,2], !CA[2,1], sum(CA) == length(CA) - 2)

# Effect of breaking the loop
y = breakLoops(x)
stopifnot(!hasUnbrokenLoops(y))
stopifnot(pedsize(y) == 7)

# A pedigree with selfing (note the necessary `sex = 0`)
z1 = singleton(1, sex = 0)
z2 = addChildren(z1, father = 1, mother = 1, nch = 1)
stopifnot(!hasSelfing(z1), hasSelfing(z2))
```
# Nucleus sub-pedigrees
stopifnot(length(subnucs(z1)) == 0)
peelingOrder(cousinPed(1))

plot.ped

Plot pedigrees with genotypes

Description

This is the main function for pedigree plotting, with many options for controlling the appearance of pedigree symbols and accompanying labels. It wraps the plotting functionality in the kinship2 package.

Usage

```r
## S3 method for class 'ped'
plot(
   x,
   marker = NULL,
   sep = "/",
   missing = "-",
   showEmpty = FALSE,
   labs = labels(x),
   title = NULL,
   col = 1,
   aff = NULL,
   carrier = NULL,
   hatched = NULL,
   shaded = NULL,
   deceased = NULL,
   starred = NULL,
   twins = NULL,
   textInside = NULL,
   textAbove = NULL,
   hints = NULL,
   fouInb = "autosomal",
   margins = c(0.6, 1, 4.1, 1),
   keep.par = FALSE,
   ...
)

## S3 method for class 'singleton'
plot(
   x,
   marker = NULL,
   sep = "/",
   ...
)
```
missing = "-",
showEmpty = FALSE,
labs = labels(x),
title = NULL,
col = 1,
aff = NULL,
carrier = NULL,
hatched = NULL,
shaded = NULL,
deceased = NULL,
starred = NULL,
textInside = NULL,
textAbove = NULL,
fouInb = "autosomal",
margins = c(8, 0, 0, 0),
yadj = 0,
...
)

as_kinship2_pedigree(
  x,
  deceased = NULL,
  aff = NULL,
twins = NULL,
hints = NULL
)

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

Arguments

x          A \texttt{ped()} object.

marker     Either a vector of names or indices referring to markers attached to \texttt{x}, a \texttt{marker} object, or a list of such. The genotypes for the chosen markers are written below each individual in the pedigree, in the format determined by \texttt{sep} and \texttt{missing}. See also \texttt{showEmpty}. If \texttt{NULL} (the default), no genotypes are plotted.

sep        A character of length 1 separating alleles for diploid markers.

missing    The symbol (integer or character) for missing alleles.

showEmpty  A logical, indicating if empty genotypes should be included.

labs       A vector or function controlling the individual labels included in the plot. Alternative forms:

  - If \texttt{labs} is a vector with nonempty intersection with \texttt{labels(x)}, these individuals will be labelled. If the vector is named, then the (non-empty) names are used instead of the ID label. (See Examples.)
  - If \texttt{labs} is \texttt{NULL}, or has nonempty intersection with \texttt{labels(x)}, then no labels are drawn.
• If `labs` is the word "num", then all individuals are numerically labelled following the internal ordering.
• If `labs` is a function, it will be replaced with `labs(x)` and handled as above. (See Examples.)

`title`  The plot title. If NULL (default) or "", no title is added to the plot.

`col` A vector of colours for the pedigree members, recycled if necessary. Alternatively, `col` can be a list assigning colours to specific members. For example if `col = list(red = "a", blue = c("b", "c"))` then individual "a" will be red, "b" and "c" blue, and everyone else black. By default everyone is black.

`aff` A vector of labels identifying members whose plot symbols should be filled. (This is typically used in medical pedigrees to indicate affected members.)

`carrier` A vector of labels identifying members whose plot symbols should be marked with a dot. (This is typically used in medical pedigrees to indicate unaffected carriers of the disease allele.)

`hatched` A vector of labels identifying members whose plot symbols should be hatched.

`shaded` (Deprecated) synonym of hatched

`deceased` A vector of labels indicating deceased pedigree members.

`starred` A vector of labels indicating pedigree members that should be marked with a star in the pedigree plot.

`twins` A data frame with columns `id1`, `id2` and `code`, passed on to the relation parameter of `kinship2::plot.pedigree()`. 

`textInside`, `textAbove` Character vectors of text to be printed inside or above pedigree symbols.

`hints` A list with alignment hints passed on to `kinship2::align.pedigree()`. Rarely necessary, but see Examples.

`fouInb` Either "autosomal" (default), "x" or NULL. If "autosomal" or "x", inbreeding coefficients are added to the plot above the inbred founders. If NULL, or if no founders are inbred, nothing is added.

`margins` A numeric of length 4 indicating the plot margins. For singletons only the first element (the 'bottom' margin) is used.

`keep.par` A logical (default = FALSE). If TRUE, the graphical parameters are not reset after plotting, which may be useful for adding additional annotation.

`...` Arguments passed on to `kinship2::plot.pedigree()`. In particular `symbolsize` and `cex` can be useful.

`yadj` A tiny adjustment sometimes needed to fix the appearance of singletons.

**Details**

This plotting function is in essence an elaborate wrapper for `kinship2::plot.pedigree()`. 

**Author(s)**

Magnus Dehli Vigeland
See Also

kinship2::plot.pedigree()

Examples

```r
x = nuclearPed(father = "fa", mother = "mo", child = "boy")
m = marker(x, fa = "1/1", boy = "1/2", name = "SNP")

plot(x, marker = m)

# Markers attached to `x` may be called by name
x = setMarkers(x, m)
plot(x, marker = "SNP")

# Other options
plot(x, marker = "SNP", hatched = typedMembers(x),
     starred = "fa", deceased = "mo")

# Filled symbols
plot(x, aff = males(x))

# Label only some members
plot(x, labs = c("fa", "boy"))

# Label only some members; rename the father
plot(x, labs = c(FATHER = "fa", "boy"))

# Label males only
plot(x, labs = males)

# Colours
plot(x, col = list(red = "fa", green = "boy"), hatched = "boy")

# Founder inbreeding is shown by default
founderInbreeding(x, "mo") = 0.1
plot(x)

# ... but can be suppressed
plot(x, fouInb = NULL)

# Twins
x = nuclearPed(children = c("tw1", "tw2", "tw3"))
plot(x, twins = data.frame(id1 = "tw1", id2 = "tw2", code = 1)) # MZ
plot(x, twins = data.frame(id1 = "tw1", id2 = "tw2", code = 1)) # DZ

# Triplets
plot(x, twins = data.frame(id1 = c("tw1", "tw2"),
                           id2 = c("tw2", "tw3"),
                           code = 2))
```

# In some cases, the plotting machinery of `kinship2` needs a hint
# (see ?kinship2::align.pedigree)

# Example with 3/4-siblings
y = nuclearPed(2)
y = addChildren(y, 3, mother = 5, nch = 1)
y = addChildren(y, 4, mother = 5, nch = 1)

plot(y) # bad

hints = list(order = 1:7, spouse = rbind(c(3,5,0), c(5,4,0)))
plot(y, hints = hints) # good

---

**plotPedList**  
*Plot a collection of pedigrees.*

**Description**

This function creates a row of pedigree plots, each created by `plot_ped()`. Any parameter accepted by `plot_ped()` can be applied, either to all plots simultaneously, or to individual plots. Some effort is made to guess a reasonable window size and margins, but in general the user must be prepared to do manual resizing of the plot window. See various examples in the Examples section below.

**Usage**

```r
plotPedList(
  plots,
  widths = NULL,
  groups = NULL,
  titles = NULL,
  frames = TRUE,
  fmar = NULL,
  frametitles = NULL,
  source = NULL,
  dev.height = NULL,
  dev.width = NULL,
  newdev = !is.null(dev.height) || !is.null(dev.width),
  verbose = FALSE,
  ...
)
```

**Arguments**

- **plots**  
  A list of lists. Each element of `plots` is a list, where the first element is a pedigree, and the remaining elements are passed on to `plot_ped`. These elements must be correctly named. See examples below.
widths A numeric vector of relative widths of the subplots. Recycled to length(plots) if necessary, before passed on to layout(). Note that the vector does not need to sum to 1.
groups A list of vectors, each consisting of consecutive integers, indicating subplots to be grouped. By default the grouping follows the list structure of plots.
titles A character vector of titles for each group. Overrides titles given in individuals subplots.
frames A logical indicating if groups should be framed.
frametitles Deprecated; use titles instead.
source NULL (default), or the name or index of an element of plots. If given, marker data is temporarily transferred from this to all the other pedigrees. This may save some typing when plotting the same genotypes on several pedigrees.
dev.height, dev.width The dimensions of the new plot window. If these are NA suitable values are guessed from the pedigree sizes.
newdev A logical, indicating if a new plot window should be opened.
verbose A logical.
... Further arguments passed on to each call to plot.ped().

Details
Note that for tweaking dev.height and dev.width the function dev.size() is useful to determine the size of the active device.

Author(s)
Magnus Dehli Vigeland

See Also
plot.ped()

Examples

# Basic examples
# Simples use: Just give a list of ped objects.
peds = list(nuclearPed(3), cousinPed(2), singleton(12), halfSibPed())
plotPedList(peds, newdev = TRUE)

# Modify the relative widths (which are not guessed)
w = c(2, 3, 1, 2)
plotPedList(peds, widths = w)
# In most cases the guessed dimensions are ok but not perfect.
# Resize plot window manually and re-plot with `newdev = FALSE` (default)
# plotPedList(peds, widths = w)

## Remove frames
plotPedList(peds, widths = w, frames = FALSE)

## Non-default grouping
plotPedList(peds, widths = w, groups = list(1, 2:3), titles = 1:2)

## Parameters added in the main call are used in each sub-plot
plotPedList(peds, widths = w, margins = c(2, 4, 2, 4), labs = leaves,
           hatched = leaves, symbolsize = 1.3, col = list(red = 1))

dev.off()

### Example of automatic grouping

H1 = nuclearPed()
H2 = list(singleton(1), singleton(3))  # grouped!

plotPedList(list(H1, H2), dev.height = 2, dev.width = 4,
           titles = c(expression(H[1]), expression(H[2])))

dev.off()

### Complex example with individual parameters for each plot

# For more control of individual plots, each plot and all
# its parameters can be specified in its own list.
x1 = nuclearPed(nch = 3)
m1 = marker(x1, `3` = "1/2")
marg1 = c(7, 4, 7, 4)
plot1 = list(x1, marker = m1, margins = marg1, title = "Plot 1",
             deceased = 1:2, cex = 1.3)

x2 = cousinPed(2)
m2 = marker(x2, `11` = "A/A", `12` = "A/A")
marg2 = c(3, 4, 2, 4)
plot2 = list(x2, marker = m2, margins = marg2, title = "Plot 2",
             symbolsize = 1.2, labs = NULL)

x3 = singleton("Mr. X")
marg3 = c(10, 0, 0, 0)
plot3 = list(x3, margins = marg3, title = "Plot 3",
             symbolsize = 1, cex = 2)

x4 = halfSibPed()
hatched = 4:5
col = list(red = founders(x4), blue = leaves(x4))
marg4 = marg1
plot4 = list(x4, margins = marg4, title = "Plot 4", cex = 1.3,
             hatched = hatched, col = col)

plotPedList(list(plot1, plot2, plot3, plot4), widths = c(2,3,1,2),
            groups = list(1, 2:3, 4), newdev = TRUE)

dev.off()

################################
# Example with large pedigrees #
################################

# Important to set device dimensions here

plotPedList(list(halfCousinPed(4), cousinPed(7)),
            titles = c("Large", "Very large"),
            dev.height = 8, dev.width = 5)

dev.off()

print.nucleus

S3 methods

Description

S3 methods

Usage

## S3 method for class 'nucleus'
print(x, ...)

Arguments

x                   An object
...

Not used
print.ped

Printing pedigrees

Description

Print a ped object using original labels.

Usage

```r
## S3 method for class 'ped'
print(x, ..., markers, verbose = TRUE)
```

Arguments

- `x`: object of class ped.
- `...`: (optional) arguments passed on to `print.data.frame()`.
- `markers`: (optional) vector of marker indices. If missing, and `x` has less than 10 markers, they are all displayed. If `x` has 10 or more markers, the first 5 are displayed.
- `verbose`: If TRUE, a message is printed if only the first 5 markers are printed. (See above).

Details

This first calls `as.data.frame.ped()` and then prints the resulting data.frame. The data.frame is returned invisibly.

randomPed

Random pedigree

Description

Generate a random pedigree by applying random mating starting from a finite population. The resulting pedigree will have \( f + g \) members, where \( f \) is the number of founders and \( g \) is the number of matings.

Usage

```r
randomPed(g, founders = rpois(1, 3) + 1, selfing = FALSE, seed = NULL)
```

Arguments

- `g`: A positive integer: The number of matings.
- `founders`: A positive integer: The size of the initial population.
- `selfing`: A logical indicating if selfing is allowed.
- `seed`: A numerical seed for random number generation. (Optional.)
Details

The sampling scheme for choosing parents in each mating depends on the selfing parameter. If selfing = FALSE, a father is randomly sampled from the existing males, and a mother from the existing females. If selfing = TRUE then one parent P1 is sampled first (among all members), and then a second parent from the set consisting of P1 and all members of the opposite sex. The gender of the child is randomly chosen with equal probabilities.

Value

A ped object.

Examples

```r
randomPed(3, 3)
randomPed(3, 3, selfing = TRUE)
```

Description

Reads a text file in pedigree format, or something fairly close to it.

Usage

```r
readPed(
  pedfile,
  colSep = "",
  header = NA,
  famid_col = NA,
  id_col = NA,
  fid_col = NA,
  mid_col = NA,
  sex_col = NA,
  marker_col = NA,
  locusAttributes = NULL,
  missing = 0,
  sep = NULL,
  validate = TRUE,
  ...
)
```

readPed  

Read a pedigree from file

---

readPed

Read a pedigree from file

---

Description

Reads a text file in pedigree format, or something fairly close to it.

Usage

```r
readPed(
  pedfile,
  colSep = "",
  header = NA,
  famid_col = NA,
  id_col = NA,
  fid_col = NA,
  mid_col = NA,
  sex_col = NA,
  marker_col = NA,
  locusAttributes = NULL,
  missing = 0,
  sep = NULL,
  validate = TRUE,
  ...
)
```
**Arguments**

- **pedfile**: A file name
- **colSep**: A column separator character, passed on as the `sep` argument of `read.table()`. The default is to separate on white space, that is, one or more spaces, tabs, newlines or carriage returns. (Note: the parameter `sep` is used to indicate allele separation in genotypes.)
- **header**: A logical. If NA, the program will interpret the first line as a header line it contains both "id" and "sex" as part of some entries (ignoring case).
- **famid_col**: Index of family ID column. If NA, the program looks for a column named "famid" (ignoring case).
- **id_col**: Index of individual ID column. If NA, the program looks for a column named "id" (ignoring case).
- **fid_col**: Index of father ID column. If NA, the program looks for a column named "fid" (ignoring case).
- **mid_col**: Index of mother ID column. If NA, the program looks for a column named "mid" (ignoring case).
- **sex_col**: Index of column with gender codes (0 = unknown; 1 = male; 2 = female). If NA, the program looks for a column named "sex" (ignoring case). If this is not found, genders of parents are deduced from the data, leaving the remaining as unknown.
- **marker_col**: Index vector indicating columns with marker alleles. If NA, all columns to the right of all pedigree columns are used. If `sep` (see below) is non-NULL, each column is interpreted as a genotype column and split into separate alleles with `strsplit(..., split = sep, fixed = TRUE)`.
- **locusAttributes**: Passed on to `setMarkers()` (see explanation there).
- **missing**: Passed on to `setMarkers()` (see explanation there).
- **sep**: Passed on to `setMarkers()` (see explanation there).
- **validate**: A logical indicating if the pedigree structure should be validated.
- **...**: Further parameters passed on to `read.table()`, e.g. `comment.char` and `quote`.

**Details**

If there are no headers, and no column information is provided by the user, the program assumes the following column order:

- family ID (optional; guessed from the data)
- individual ID
- father’s ID
- mother’s ID
- sex
- marker data (remaining columns)
relabel

Get or modify pedigree labels

Description

Functions for getting or changing the ID labels of pedigree members.
Usage

relabel(x, new, old = labels(x), reorder = FALSE)

## S3 method for class 'ped'
labels(object, ...)

## S3 method for class 'list'
labels(object, ...)

Arguments

x 
A ped object or a list of such.

new, old 
Character vectors (or coercible to character) of the same length. ID labels in old are replaced by those in new.

reorder 
A logical. If TRUE, reorderPed() is called on x after relabelling. Default: FALSE.

object 
A ped object

... 
Not used

Value

- labels() returns a character vector containing the ID labels of all pedigree members. If the input is a list of ped objects, the output is a list of character vectors.
- relabel() returns ped object similar to the input except for the labels.

Author(s)

Magnus Dehli Vigeland

See Also

ped()

Examples

x = nuclearPed()
x
labels(x)
relabel(x, new = "girl", old = 3)
setSNPs

Attach SNP loci to a pedigree

Description
Create and attach a list of empty SNP markers with specified position and allele frequencies.

Usage

setSNPs(x, snpData)

Arguments

x
A ped object.
snpData
A data frame with 6 columns. See Details.

Details
The data frame snpData should contain the following columns, in order:

- CHROM: Chromosome (character)
- MARKER: Marker name (character)
- MB: Physical position in megabases (numeric)
- A1: First allele (single-letter character)
- A2: Second allele (single-letter character)
- FREQ1: Allele frequency of A1 (number in [0, 1])

The actual column names do not matter.
Each column must be of the stated type, or coercible to it. (For example, CHROM, A1 and A2 may be given as numbers, but will be internally converted to characters.)

Value
A copy of x with the indicated SNP markers attached.

Examples

snps = data.frame(
  CHROM = 1:2,
  MARKER = c("M1", "M2"),
  MB = c(1.23, 2.34),
  A1 = c("A", "G"),
  A2 = c("C", "C"),
  FREQ1 = c(0.7, 0.12))

x = setSNPs(nuclearPed(), snpData = snps)
sortGenotypes

# Inspect the results:
getMap(x)
getFreqDatabase(x)

sortGenotypes  Sort the alleles in each genotype

Description

Ensure that all genotypes are sorted internally. For example, if a marker attached to x has alleles 1 and 2, then running this function will replace all genotypes "2/1" by "1/2".

Usage

sortGenotypes(x)

Arguments

x  A ped object or a list of such

Value

An object identical to x except that the all genotypes are sorted.

Examples

x = singleton(1)

# Various markers with misordered genotypes
m1 = marker(x, '1' = "2/1")
m2 = marker(x, '1' = "b/a")
m3 = marker(x, '1' = "100.3/99.1")
x = setMarkers(x, list(m1, m2, m3))
x

# Sort all genotypes
y = sortGenotypes(x)
y

# Also works when input is a list of peds
sortGenotypes(list(x, x))
Transfer marker data between pedigrees. Any markers attached to the target are overwritten.

Usage

```r
transferMarkers(
  from,
  to,
  ids = NULL,
  idsFrom = ids,
  idsTo = ids,
  erase = TRUE,
  matchNames = TRUE,
  checkSex = FALSE
)
```

Arguments

- `from`: A ped or singleton object, or a list of such objects.
- `to`: A ped or singleton object, or a list of such objects.
- `ids`: A vector of ID labels. This should be used only if the individuals have the same name in both pedigrees; otherwise use `idsFrom` and `idsTo` instead.
- `idsFrom`, `idsTo`: Vectors of equal length, denoting source individuals (in the `from` pedigree) and target individuals (in the `to` pedigree), respectively.
- `erase`: A logical. If TRUE (default), all markers attached to `to` are erased prior to transfer, and new marker objects are created with the same attributes as in `from`. If FALSE no new marker objects are attached to `to`. Only the genotypes of the `ids` individuals are modified, while genotypes for other pedigree members - and marker attributes - remain untouched.
- `matchNames`: A logical, only relevant if `erase` = FALSE. If `matchNames` = TRUE (default) marker names are used to ensure genotypes are transferred into the right markers. The output contains only markers present in `from`, in the same order. (An error is raised if the markers are not named.)
- `checkSex`: A logical. If TRUE, it is checked that `fromIds` and `toIds` have the same sex. Default: FALSE.

Details

By default, genotypes are transferred between all individuals present in both pedigrees.


**validatePed** 75

**Value**

A ped object (or a list of such) similar to to, but where all individuals also present in from have marker genotypes copied over. Any previous marker data is erased.

**Examples**

```r
x = nuclearPed(fa = "A", mo = "B", child = "C")
x = addMarker(x, A = "1/2", B = "1/1", C = "1/2", name = "M1")

y = list(singleton("A"), nuclearPed(fa = "D", mo = "B", child = "C"))

# By default all common individuals are transferred
transferMarkers(x, y)

# Transfer data for the boy only
transferMarkers(x, y, ids = "C")

# Transfer without first erasing the target markers
z = nuclearPed(fa = "A", mo = "B", child = "C")
z = addMarker(z, A = "1/1", alleles = 1:2, name = "M1")

transferMarkers(x, z, ids = "C", erase = FALSE)
transferMarkers(x, z, ids = "C", erase = TRUE) # note the difference
```

**validatePed**  Pedigree errors

**Description**

Validate the internal structure of a ped object.

**Usage**

validatePed(x)

**Arguments**

- x : object of class ped.

**Value**

If no errors are detected, the function returns NULL invisibly. Otherwise, messages describing the errors are printed to the screen and an error is raised.
writePed Write a pedigree to file

Description

Write a pedigree to file

Usage

writePed(
  x,
  prefix,
  what = "ped",
  famid = is.pedList(x),
  header = TRUE,
  merlin = FALSE,
  verbose = TRUE
)

Arguments

x A ped object
prefix A character string giving the prefix of the files. For instance, if prefix = "myped" and what = c("ped", "map"), the output files are "myped.ped" and "myped.map" in the current directory. Paths to other folder may be included, e.g. prefix = "path-to-my-dir/myped".
what A subset of the character vector c("ped", "map", "dat", "freq"), indicating which files should be created. By default only the "ped" file is created. This option is ignored if merlin = TRUE.
famid A logical indicating if family ID should be included as the first column in the ped file. The family ID is taken from famid(x). If x is a pedlist, the family IDs are taken from names(x), or if this is NULL, the component-wise famid() values. Missing values are replaced by natural numbers. This option is ignored if merlin = TRUE.
header A logical indicating if column names should be included in the ped file. This option is ignored if merlin = TRUE.
merlin A logical. If TRUE, "ped", "map", "dat" and "freq" files are written in a format readable by the MERLIN software. In particular MERLIN requires non-numerical allele labels in the frequency file.
verbose A logical.

Value

A character vector with the file names.
Examples

```r
x = nuclearPed(1)
x = addMarker(x, "3" = "a/b", name = "m1")

# Write to file
fn = writePed(x, prefix = tempfile("test"))

# Read
y = readPed(fn)

stopifnot(identical(x, y))
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