

Package ‘pedigree’

October 14, 2022

Type Package

Title Pedigree Functions

Version 1.4.2

Date 2022-08-13

Depends Matrix

Imports methods,HaploSim (>= 1.8.4),reshape

Description Pedigree related functions.

License GPL (>= 2)

Author Albart Coster [aut, cre]

Maintainer Albart Coster <albart@dairyconsult.nl>

NeedsCompilation yes

Repository CRAN

Date/Publication 2022-08-13 20:50:02 UTC

R topics documented:

| | |
|------------------|----|
| pedigree-package | 2 |
| add.Inds | 2 |
| blup | 3 |
| calcG | 4 |
| calcInbreeding | 4 |
| countGen | 5 |
| countOff | 6 |
| gblup | 6 |
| makeA | 8 |
| makeAinv | 8 |
| orderPed | 9 |
| trimPed | 10 |

Index

11

pedigree-package *Package to deal with pedigree data*

Description

Package with functions to analyse and transform pedigree data. A pedigree is a `data.frame` where the first column contains an ID, and the second and third columns contain ID of first and second parent.

Author(s)

Albart Coster: <albart.coster@wur.nl>

See Also

[trimPed](#) [orderPed](#) [countGen](#) [makeA](#) [makeAinv](#) [calcInbreeding](#) [add.Inds](#)

add.Inds *Function to add missing individuals to a pedigree*

Description

Function `add.Inds()` adds missing individuals to a pedigree and returns the complete pedigree as a `data.frame` with the same headers as the original pedigree. Remember to check for errors beforehand with function `errors.ped`. Unknown parents should be coded as NA.

Usage

`add.Inds(ped)`

Arguments

`ped` `data.frame` with three columns: id,id parent1,id parent2

Value

`data.frame` of three columns with identical header as input.

Author(s)

Albart Coster, Albart.Coster@wur.nl

See Also

[orderPed](#)

Examples

```
ID <- 3:5
DAM <- c(1,1,3)
SIRE <- c(2,2,4)
pedigree <- data.frame(ID,DAM,SIRE)
pedigree <- add.Inds(pedigree)
```

blup

Function to calculate breeding values using an animal model

Description

Fit an animal model to data, use a given variance ratio ($\alpha = \frac{\sigma_e^2}{\sigma_a^2}$). Calculate inverse of the additive genetic relationship matrix using function `makeInv()` of this package.

Usage

```
blup(formula, ped, alpha, trim = FALSE)
```

Arguments

- `formula` formula of the model, do not include the random effect due to animal (generally ID).
- `ped` `data.frame` with columns corresponding to ID, SIRE, DAM and the columns in the formula.
- `alpha` Variance ratio ($\frac{\sigma_e^2}{\sigma_a^2}$).
- `trim` If TRUE, trims the pedigree using the available phenotype data using function `trimPed`.

Value

Vector of solutions to the model, including random animal effects.

See Also

[SamplePedigree](#), [gblup](#), [makeAinv](#), [blup](#)

Examples

```
example(gblup)
sol <- blup(P~1,ped = ped,alpha = 1/h2 - 1)
```

| | |
|--------------------|--|
| <code>calcG</code> | <i>Function to calculate a relationship matrix from marker data (usually allele count data), G matrix.</i> |
|--------------------|--|

Description

Function to calculate a relationship matrix from marker data. Option to return the inverse of matrix. Inverse calculated using `Matrix` package.

Usage

```
calcG(M, data = NULL, solve = FALSE)
```

Arguments

| | |
|--------------------|---|
| <code>M</code> | Matrix of marker genotypes, usually the count of one of the two SNP alleles at each markers (0, 1, or 2). |
| <code>data</code> | Optional logical vector which can tell of which individuals we have phenotypes. |
| <code>solve</code> | Logic, if TRUE then function returns the inverse of the relationship matrix. |

Value

Matrix of class `dgeMatrix`.

See Also

[SamplePedigree](#), [gblup](#), [makeAinv.blup](#)

Examples

```
example(gblup)
G <- calcG(M)
Ginv <- calcG(M, solve = TRUE)
```

| | |
|-----------------------------|--|
| <code>calcInbreeding</code> | <i>Calculates inbreeding coefficients for individuals in a pedigree.</i> |
|-----------------------------|--|

Description

Calculates inbreeding coefficients of individuals in a pedigree.

Usage

```
calcInbreeding(ped)
```

`countGen`

5

Arguments

`ped` data.frame with three columns: id,id parent1,id parent2

Value

Logical.

Examples

```
id <- 1:6
dam <- c(0,0,1,1,4,4)
sire <- c(0,0,2,2,3,5)
ped <- data.frame(id,dam,sire)
(F <- calcInbreeding(ped))
```

`countGen`

Count generation number for each individual in a pedigree.

Description

Counts generation number for individuals in a pedigree.

Usage

`countGen(ped)`

Arguments

`ped` data.frame with three columns: id,id parent1,id parent2

Value

Numeric vector

Examples

```
id <- 1:5
dam <- c(0,0,1,1,4)
sire <- c(0,0,2,2,3)
ped <- data.frame(id,dam,sire)
(gens <- countGen(ped))
```

| | |
|----------|--|
| countOff | <i>Function that counts the number of offspring (and following generations for each individual in a pedigree).</i> |
|----------|--|

Description

Function to count the number of offspring for each individual in a pedigree. With loops, offspring of later generations will be counted several times.

Usage

```
countOff(ped)
```

Arguments

| | |
|-----|---|
| ped | data.frame with three columns: id,id parent1,id parent2 |
|-----|---|

Value

Numeric vector with number of offspring for each individual in the pedigree.

Author(s)

Albart Coster

Examples

```
example(countGen)
countOff(ped)
```

| | |
|-------|---|
| gblup | <i>Function to calculate breeding values using an animal model and a relationship matrix calculated from the markers (G matrix)</i> |
|-------|---|

Description

Fit an animal model to data, use a given variance ratio ($\alpha = \frac{\sigma_e^2}{\sigma_a^2}$). Calculate genetic relationship matrix using the function calcG of this package.

Usage

```
gblup(formula, data, M, lambda)
```

Arguments

| | |
|---------|---|
| formula | formula of the model, do not include the random effect due to animal (generally ID). |
| data | data.frame with columns corresponding to ID and the columns mentioned in the formula. |
| M | Matrix of marker genotypes, usually the count of one of the two SNP alleles at each markers (0, 1, or 2). |
| lambda | Variance ratio ($\frac{\sigma_e^2}{\sigma_a^2}$) |

Value

Vector of solutions to the model, including random animal effects.

See Also

[SamplePedigree](#), [gblup](#), [makeAinv](#), [blup](#)

Examples

```
## Example Code from SampleHaplotypes
hList <- HaploSim::SampleHaplotypes(nHaplotypes = 20,genDist =
1,nDec = 3,nLoc = 20) ## create objects
h <- HaploSim::SampleHaplotype(H0 = hList[[1]],H1 = hList[[2]],genDist =
1,nDec = 3)

## code from the Example SamplePedigree
ID <- 1:10
pID0 <- c(rep(0,5),1,1,3,3,5)
pID1 <- c(rep(0,4),2,2,2,4,4,6)
ped <- data.frame(ID,pID0,pID1)
phList <- HaploSim::SamplePedigree(orig = hList,ped = ped)

## own code
h2 <- 0.5
ped <- phList$ped
hList <- phList$hList
qtllist <- HaploSim::ListQTL(hList = hList,frqtl = 0.1,sigma2qtl = 1)
qtllist <- tapply(unlist(qtllist),list(rep(names(qtllist),times = unlist(lapply(qtllist,length)))), 
unlist(lapply(qtllist,function(x)seq(1,length(x))))),mean,na.rm = TRUE)
qtllist <- reshape::melt(qtllist)
names(qtllist) <- c("POS","TRAIT","a")
HH <- HaploSim::getAll(hList,translatePos = FALSE)
rownames(HH) <- sapply(hList,function(x)x@hID)
QQ <- HH[,match(qtllist$POS,colnames(HH))]
g <- QQ
ped$G <- with(ped,g[match(hID0,rownames(g))]+g[match(hID1,rownames(g))])
sigmae <- sqrt(var(ped$G)/h2 - var(ped$G))
ped$P <- ped$G + rnorm(nrow(ped),0,sigmae)
M <- with(ped,HH[match(hID0,rownames(HH)),] + HH[match(hID1,rownames(HH)),])
rownames(M) <- ped$ID
```

```
sol <- gblup(P~1,data = ped[,c('ID','P')],M = M,lambda = 1/h2 - 1)
```

makeA*Makes the A matrix for a part of a pedigree***Description**

Makes the A matrix for a part of a pedigree and stores it in a file called A.txt.

Usage

```
makeA(ped,which)
```

Arguments

| | |
|--------------------|---|
| <code>ped</code> | data.frame with three columns: id,id parent1,id parent2 |
| <code>which</code> | Logical vector specifying between which individuals additive genetic relationship is required. Goes back through the whole pedigree but only for subset of individuals. |

Value

Logical.

Examples

```
id <- 1:6
dam <- c(0,0,1,1,4,4)
sire <- c(0,0,2,2,3,5)
ped <- data.frame(id,dam,sire)
makeA(ped,which = c(rep(FALSE,4),rep(TRUE,2)))
A <- read.table("A.txt")

if(file.exists("A.txt"))
file.remove("A.txt")
```

makeAinv*Makes inverted A matrix for a pedigree***Description**

Makes inverted A matrix for a pedigree and stores it in a file called Ainv.txt.

Usage

```
makeAinv(ped)
```

orderPed

9

Arguments

ped data.frame with three columns: id,id parent1,id parent2

Value

Logical.

Examples

```
id <- 1:6
dam <- c(0,0,1,1,4,4)
sire <- c(0,0,2,2,3,5)
ped <- data.frame(id,dam,sire)
makeAinv(ped)
Ai <- read.table('Ainv.txt')
nInd <- nrow(ped)
Ainv <- matrix(0,nrow = nInd,ncol = nInd)
Ainv[as.matrix(Ai[,1:2])] <- Ai[,3]
dd <- diag(Ainv)
Ainv <- Ainv + t(Ainv)
diag(Ainv) <- dd

if(file.exists("Ainv.txt"))
file.remove("Ainv.txt")
```

orderPed

Orders a pedigree

Description

Orders a pedigree so that offspring follow parents.

Usage

`orderPed(ped)`

Arguments

ped data.frame with three columns: id,id parent1,id parent2

Value

numerical vector

Examples

```
id <- 1:6
dam <- c(0,0,1,1,4,4)
sire <- c(0,0,2,2,3,5)
pedigree <- data.frame(id,dam,sire)
(ord <- orderPed(pedigree))
pedigree <- pedigree[6:1,]
(ord <- orderPed(pedigree))
pedigree <- pedigree[order(ord),]
pwroneg <- pedigree
pwroneg[1,2] <- pwroneg[6,1]
```

trimPed

Function to trim a pedigree based on available data

Description

Trims a pedigree given a vector of data. Branches without data are trimmed off the pedigree.

Usage

```
trimPed(ped, data, ngenback = NULL)
```

Arguments

| | |
|----------|---|
| ped | data.frame with three columns: id,id parent1,id parent2 |
| data | TRUE-FALSE vector. Specifies if data for an individual is available. |
| ngenback | Number of generations back. Specifies the number of generations to keep before the individuals with data. |

Value

Logical vector specifying if an individual should stay in the pedigree.

Examples

```
id <- 1:5
dam <- c(0,0,1,1,4)
sire <- c(0,0,2,2,3)
data <- c(FALSE,FALSE,TRUE,FALSE,FALSE)
ped <- data.frame(id,dam,sire)
yn <- trimPed(ped,data)
ped <- ped[yn,]
```

Index

* utilities

add.Inds, 2
blup, 3
calcG, 4
calcInbreeding, 4
countGen, 5
countOff, 6
gblup, 6
makeA, 8
makeAinv, 8
orderPed, 9
pedigree-package, 2
trimPed, 10

add.Inds, 2, 2
blup, 3, 3, 4, 7
calcG, 4
calcInbreeding, 2, 4
countGen, 2, 5
countOff, 6
gblup, 3, 4, 6, 7
makeA, 2, 8
makeAinv, 2–4, 7, 8
orderPed, 2, 9
pedigree (pedigree-package), 2
pedigree-package, 2
SamplePedigree, 3, 4, 7
trimPed, 2, 3, 10