Package 'ggroups'

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Title Pedigree and Genetic Groups

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Description Calculates additive genetic relationship matrix and its inverse, in matrix and tabular-sparse formats. It includes functions for checking and processing pedigree, as well as functions to calculate the matrix of genetic group contributions (Q), and adding those contributions to the genetic merit of animals (Quaas (1988) <doi:10.3168 jds.s0022-0302(88)79691-5="">). Calculation of Q is computationally extensive. There are computationally optimized functions to calculate Q.</doi:10.3168>
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Pedigree and genetic groups

Description

ggroups-package

This package contains pedigree processing and analyzing functions, including functions for checking and renumbering the pedigree, making the pedigree relationship matrix and its inverse, in matrix and tabular formats, as well as functions related to genetic groups.

Details

First, it is recommended to check the pedigree data. frame with the pedcheck function. Pedigree relationship matrix and its inverse are fundamentals in the conventional and modern animal breeding. The concept of genetic groups stems from the fact that not all the unknown parents are of the same genetic level. The genetic group contribution matrix (\mathbf{Q}) is required to weight and add genetic group effects ($\hat{\mathbf{g}}$) to the genetic merit of animals ($\hat{\mathbf{u}}$), which is equal to $\mathbf{Q}\hat{\mathbf{g}}+\hat{\mathbf{u}}$ (Quaas, 1988). Calculating \mathbf{Q} is computationally challenging, and for large pedigree, large RAM and long computational time is required. Therefore, the functions qmatL and its parallel version, qmatXL are introduced. Overlap between sire and dam genetic groups is supported.

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References

Quaas, R. L. 1988. Additive Genetic Model with Groups and Relationships. J. Dairy Sci., 71:1338-1345. <doi:10.3168.jds.S0022-0302(88)79691-5>

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buildA

Relationship matrix A

Description

Builds the pedigree-based additive genetic relationship matrix.

Usage

```
buildA(ped)
```

Arguments

ped

: data. frame with integer columns corresponding to ID, SIRE, DAM. Missing value is $\boldsymbol{0}$.

Value

Relationship matrix A

Examples

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) buildA(ped)
```

gghead

Append genetic groups to the pedigree

Description

This function appends parents that are not available in the first column of the pedigree, to the head of the pedigree, and sorts it. Given a pedigree with all missing parents replaced with the corresponding genetic groups, this functions appends genetic groups to the head of the pedigree.

Usage

```
gghead(ped)
```

Arguments

ped

: data.frame with integer columns corresponding to ID, SIRE, DAM. Missing value is 0.

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Details

Consider this simple pedigree:

3 0 0

4 3 0

6 4 5

5 0 0

First, unknown parents are replaced with the corresponding genetic groups.

Please note that unknown parent IDs should be smaller than animal IDs.

3 1 2

4 3 2

6 4 5

5 1 2

Then, gghead is applied to this pedigree (see the example).

Value

Processed pedigree data.frame

Examples

```
ped = data.frame(ID=c(3,4,6,5), SIRE=c(1,3,4,1), DAM=c(2,2,5,2)) gghead(ped)
```

inb

Inbreeding coefficient

Description

Calculate inbreeding coefficient for an individual.

Usage

```
inb(ped, id)
```

Arguments

ped : data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

id : Numeric ID of an individual

Value

Inb: Inbreeding coefficient of the individual

pedcheck 5

Examples

```
ped = data.frame(ID=1:7, SIRE=c(0,0,1,1,3,1,5), DAM=c(0,0,0,2,4,4,6)) inb(ped, 7)
```

pedcheck

Basic pedigree checks

Description

Performs basic pedigree checks.

Usage

```
pedcheck(ped)
```

Arguments

ped

: data. frame with integer columns corresponding to ID, SIRE, DAM. Missing value is 0.

Examples

peddown

Downward pedigree extraction

Description

Extract pedigree downward for one or a group of animals to find their descendants

Usage

```
peddown(ped, parents)
```

Arguments

ped

: data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

parents

: Vector of animal(s), from which the new pedigree is being extracted.

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Value

```
newped: Extracted pedigree data.frame
```

Examples

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) peddown(ped, c(1,4))
```

pedup

Upward pedigree extraction

Description

Extract pedigree upward for one or a group of animals to find their ascendants

Usage

```
pedup(ped, progeny)
```

Arguments

ped

: data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

progeny

: Vector of animal(s), from which the new pedigree is being extracted.

Value

```
newped: Extracted pedigree data.frame
```

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) pedup(ped, c(1,4))
```

pruneped 7

Description

Pruning pedigree in two different modes (strict, loose)

Usage

```
pruneped(ped, pheno, mode)
```

Arguments

ped : data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

pheno : Vector of phenotyped animals

mode : strict or loose

Details

In strict pruning, animals without progeny and phenotype are recursively deleted from the pedigree, and then animals without known parent and without progeny (if any) are deleted. Therefore, all uninfluential animals are deleted. The downside is that animals without phenotype or phenotyped progeny cannot receive any genetic merit based on the information from their phenotyped relatives. In loose pruning, the pedigree is upward extracted for phenotyped animals to thier founders, and then the pedigree is downward extracted from the founders.

Value

```
newped: Pruned pedigree data.frame
```

```
ped = data.frame(ID=1:7, SIRE=c(0,0,1,3,1,4,0), DAM=c(0,0,2,2,2,5,0)) pheno = c(1,4) pruneped(ped, pheno, mode="strict") pruneped(ped, pheno, mode="loose")
```

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Qgpu

 $Vector \mathbf{Qg} + \mathbf{u}$

Description

Adds genetic group contributions to the genetic merit of animals.

Usage

```
Qgpu(Q, sol)
```

Arguments

Q : The output matrix from qmat (for more details: ?qmat)

sol : data.frame with 2 numeric columns corresponding to ID, EBV ([$\hat{\mathbf{g}},~\hat{\mathbf{u}}]$),

where $\hat{\mathbf{g}}$ and $\hat{\mathbf{u}}$ are the genetic group and genetic merit solutions, respectively. The order of solutions must be the order of columns and the order of rows in

matrix Q.

Value

```
Vector of \mathbf{Q}\hat{\mathbf{g}} + \hat{\mathbf{u}}
```

Examples

qmat

Matrix Q

Description

Creates the genetic group contribution matrix.

Usage

```
qmat(ped2)
```

Arguments

ped2

: The output data. frame from gghead (for more details: ?gghead)

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Value

Matrix **Q**

Examples

```
ped = data.frame(ID=c(3,4,6,5), SIRE=c(1,3,4,1), DAM=c(2,2,5,2))
ped2 = gghead(ped)
qmat(ped2)
```

qmatL

Matrix **Q** for large pedigrees

Description

Creates the genetic group contribution matrix for large pedigrees.

Usage

```
qmatL(ped2)
```

Arguments

ped2

: The output data.frame from gghead (for more details: ?gghead)

Details

Calculation of the genetic group contribution matrix for large pedigrees requires a lot of memory and time. This might not be possible on ordinary computers, using the function qmat. The function qmatL takes less RAM and time, making the calculation possible for ordinary computers.

Value

Matrix Q

```
ped = data.frame(ID=c(3,4,6,5), SIRE=c(1,3,4,1), DAM=c(2,2,5,2)) ped2 = gghead(ped) qmatL(ped2)
```

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qmatXL

Matrix **Q** for large pedigrees (parallel processing)

Description

Creates the genetic group contribution matrix for large pedigrees, with parallel processing.

Usage

```
qmatXL(ped2, ncl)
```

Arguments

ped2 : The output data. frame from gghead (for more details: ?gghead)

ncl : User defined number of nodes; if the number of user defined nodes is greater

than the number of genetic groups, the number genetic groups is considered as

the number of nodes.

Details

This function is the parallel version of qmatL. It requires foreach and doParallel packages.

Value

Matrix Q

Examples

```
ped = data.frame(ID=c(3,4,6,5), SIRE=c(1,3,4,1), DAM=c(2,2,5,2)) ped2 = gghead(ped) qmatXL(ped2, 2)
```

renum

Pedigree renumbering

Description

Renumbering pedigree to numerical IDs, so that progeny's ID is smaller than parents' IDs.

Usage

```
renum(ped)
```

rg 11

Arguments

ped : data.frame with columns corresponding to ID, SIRE, DAM. Missing value

is 0.

Value

newped: Pedigree data.frame with renumberred IDs.

Cross-reference data. frame with 2 columns for original and renumberred IDs.

Examples

```
ped = data.frame(ID=letters[1:6], SIRE=c(0,0,letters[c(1,3,1,4)]), DAM=c(0,0,letters[c(2,2,2,5)])) \\ renum(ped)
```

rg

Genetic relationship coefficient

Description

Calculate genetic relationship coefficient between two individuals.

Usage

```
rg(ped, id1, id2)
```

Arguments

ped : data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

id1 : Numeric ID of an individualid2 : Numeric ID of an individual

Value

rG: Genetic relationship coefficient between the two individuals

```
ped = data.frame(ID=1:7, SIRE=c(0,0,1,1,3,1,5), DAM=c(0,0,0,2,4,4,6)) rg(ped, 5, 6)
```

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tab2mat

Tabular to matrix

Description

Converts tabular data to matrix data.

Usage

```
tab2mat(tab)
```

Arguments

tab

: data.frame with 2 integer (IDs) and 1 numeric (values) columns.

Value

matrix

Examples

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) tab2mat(tabA(ped))
```

tabA

Relationship matrix A in a tabular format

Description

Creates the pedigree-based additive genetic relationship data. frame.

Usage

tabA(ped)

Arguments

ped

: data. frame with integer columns corresponding to ID, SIRE, DAM. Missing value is $\boldsymbol{0}.$

Value

Genetic relationship data. frame

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Examples

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) tabA(ped)
```

tabAinv

Inverse of the relationship matrix A in a tabular format

Description

Creates the inverse of the pedigree-based additive genetic relationship matrix in a data.frame.

Usage

```
tabAinv(ped, inbr)
```

Arguments

ped : data.frame with integer columns corresponding to ID, SIRE, DAM. Missing

value is 0.

inbr : Inbreeding coefficients in the order of animals in the relationship matrix.

Value

Inverse of the genetic relationship data. frame

```
ped = data.frame(ID=1:6, SIRE=c(0,0,1,3,1,4), DAM=c(0,0,2,2,2,5)) inbr = c(0, 0, 0, 0.25, 0, 0.25) # or (inbr = diag(buildA(ped)) - 1) # or inbr = tabA(ped); (inbr = inbr[inbr[,1]==inbr[,2],]$a - 1) # or # For individual inbreeding values, use function inb. tabAinv(ped, inbr)
```

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