# Package 'KinMixLite'

October 9, 2024

| Type Package   |
|--|
| Title Inference About Relationships from DNA Mixtures  |
| Version 2.1.1  |
| <b>Date</b> 2024-10-08   |
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| <b>Description</b> Methods for inference about relationships between contributors to a DNA mixture and other individuals of known genotype: a basic example would be testing whether a contributor to a mixture is the father of a child of known genotype. This provides most of the functionality of the 'KinMix' package, but with some loss of efficiency and restriction on problem size, as the latter uses 'RHugin' as the Bayes net engine, while this package uses 'gRain'. The package implements the methods introduced in Green, P. J. and Mortera, J. (2017) <doi:10.1016 j.fsigen.2017.02.001=""> and Green, P. J. and Mortera, J. (2021) <doi:10.1111 rssc.12498="">.</doi:10.1111></doi:10.1016> |
| License GPL (>= 2)   |
| Depends DNAmixturesLite, gRaven  |
| <b>Imports</b> statnet.common, gRbase, Rsolnp, numDeriv, Matrix, ribd, pedtools, methods   |
| <pre>URL https://petergreenweb.wordpress.com/kinmix/</pre>   |
| NeedsCompilation no  |
| Author Peter Green [aut, cre]  |
| Repository CRAN  |
| <b>Date/Publication</b> 2024-10-09 08:50:02 UTC  |
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KinMixLite-package

Inference About Relationships from DNA Mixtures

#### Description

Methods for inference about relationships between contributors to a DNA mixture and other individuals of known genotype: a basic example would be testing whether a contributor to a mixture is the father of a child of known genotype. This provides most of the functionality of the 'Kin-Mix' package, but with some loss of efficiency and restriction on problem size, as the latter uses 'RHugin' as the Bayes net engine, while this package uses 'gRain'. The package implements the methods introduced in Green, P. J. and Mortera, J. (2017) <doi:10.1016/j.fsigen.2017.02.001> and Green, P. J. and Mortera, J. (2021) <doi:10.1111/rssc.12498>.

#### **Details**

This package is a toolkit for inference about mixtures and familial relationships, either between contributors or between a contributor and other typed individuals. It extends the functionality proposed in Green and Mortera (2017) by allowing more general relationships, specified in general by an IBD pattern distribution - the generalisation to more than two individuals of the coefficients of identity of Jacquard (1974). Details are in the paper by Green and Mortera (2021). KinMixLite

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extends the capability of the **DNAmixturesLite** package, and intimately relies on that package; as with that package, instead of the **RHugin** package, it uses **gRaven** and **gRain** for Bayes Net calculations. This version implements the ALN, MBN and WLR as well as RPT methods; see Green and Mortera (2017).

#### **Formats**

See formats for formats of the various data objects used in this package.

#### Author(s)

Maintainer: Peter Green < P.J. Green @ bristol.ac.uk>

#### References

Green, P. J. and Mortera, J. (2017). Paternity testing and other inference about relationships from DNA mixtures. *Forensic Science International: Genetics*. <doi:10.1016/j.fsigen.2017.02.001>.

Green, P. J. and Mortera, J. (2021). Inference about complex relationships using peak height data from DNA mixtures. *Applied Statistics*. <doi:10.1111/rssc.12498>.

Jacquard, A. (1974) The genetic structure of populations. Springer-Verlag.

#### See Also

DNAmixturesLite

```
require(ribd)
data(test2data)
data(NGMDyes)
C<-50
## Fit 2-person mixture - baseline model
mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db)</pre>
pars < mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.7, U2=0.3)))
baseline<-logL(mixD)(pars)</pre>
## Fit 2-person mixture model in which contributor 1 is parent of a typed individual Cgt
mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db,triangulate=FALSE,compile=FALSE)</pre>
rpt.IBD(mixD,'parent',list(c=Cgt),targets=c('f','c'),contrib='f')
log10LR<-(logL(mixD)(pars)-baseline)/log(10)</pre>
cat('log10 LR',log10LR,'\n')
## Fit 2-person mixture, where contributors are siblings
mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db)</pre>
pars < -mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.6, U2=0.3, U3=0.1)))
baseline<-logL(mixD)(pars)</pre>
```

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```
\label{limixD} $$\min xD < -DNAmixture(list(epg), k=2, C=list(C), database=db, triangulate=FALSE, compile=FALSE)$$ $$\operatorname{rpt.IBD(mixD,'sibs', targets=c('b1','b2'), contribs=c('b1','b2'))}$$ $$\log 10LR < -(\operatorname{protected(logL(mixD)(pars))-baseline)/log(10)}$$ $$\operatorname{cat('log10 LR', log10LR,'\n')}$$
```

add.child.meiosis.nodes

Replace CPTs for mixture contributor a Father, given Child genotype, by MBN method

#### Description

loop over markers, and alleles within markers to create nodes for child allele count nodes, for paternity model with only Child genotyped then compile all domains. Implements method MBN.

## Usage

```
add.child.meiosis.nodes(mixture,aca,ind=1)
```

# **Arguments**

mixture A compiled DNAmixture object

aca Child's genotype profile as an allele count array

ind Index of contributor regarded as Parent (or Child): which 'unknown' contributor

are we modelling by amending his/her CPTs?

# Details

To calculate the likelihood of this model, conditional on the child's genotype, a call to this function should be followed by (a) setting the finding of the child's genotype by defining extra.findings, (b) evaluating the loglikelihood using logLX, and (c) correcting the result by subtracting the log probability of the child's genotype, all as in the example below. Without (c), the value returned is the likelihood for the peak heights *and* the child's genotype.

#### Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

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#### **Examples**

```
data(test2data)
# set threshold C
C<-0.001
mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db)</pre>
pars < mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.7, U2=0.3)))
baseline<-logL(mixD)(pars)</pre>
mixMBN<-DNAmixture(list(epg),k=2,C=list(C),database=db,triangulate=FALSE,compile=FALSE)</pre>
cgtcaca<-gt2aca(mixMBN,Cgt)</pre>
add.child.meiosis.nodes(mixMBN,cgtcaca,1)
log10LR<-(logLX(mixMBN,</pre>
expr.make.findings(list(
list('Male',ind=1),
list('Caca',aca='cgtcaca')
))
)(pars)-attr(cgtcaca, 'logGt')-baseline)/log(10)
cat('log10 LR',log10LR,'\n')
```

add.motherchild.likd.node

Replace CPTs for mixture contributor a Father, given Child and Mother genotypes, by ALN method

## **Description**

loop over markers, and alleles within markers to create node Rlikd for relative likelihood for individual i, for paternity model with Mother and Child genotyped then compile all domains. Implements method ALN.

#### Usage

```
add.motherchild.likd.node(mixture,Cgt,Mgt,db,ind=1)
```

#### **Arguments**

| mixture | A DNAmixture object   |
|---------|---|
| Cgt     | Child's genotype profile as a data frame containing variables marker, allele1 and allele2                   |
| Mgt     | Mother's genotype profile as a data frame containing variables marker, allele1 and allele2                  |
| db      | Allele frequency database   |
| ind     | Index of contributor regarded as Father: which 'unknown' contributor are we modelling by amending his CPTs? |

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#### Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### **Examples**

```
data(test2data)
# set threshold C
C<-0.001
mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db)</pre>
pars < mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.7, U2=0.3)))
baseline<-logL(mixD)(pars)</pre>
mixD3 < -DNAmixture(list(epg), k=2, C=list(0.001), database=db, triangulate=FALSE, compile=FALSE)
cgtcaca<-gt2aca(mixD3,Cgt)
add.motherchild.likd.node(mixD3,Cgt,Mgt,db,1)
log10LR<-(logLX(mixD3,
expr.make.findings(list(
list('Male',ind=1),
list('Rlikd',aca='cgtcaca',cgt='Cgt',evid='Revid')
))
)(pars)-baseline)/log(10)
cat('log10 LR',log10LR,'\n')
```

add.relative.likd.node

Replace CPTs for mixture contributor a Father, given Child genotype, by ALN method

# Description

loop over markers, and alleles within markers to create node Rlikd for relative likelihood for individual i, for paternity model with only Child genotyped then compile all domains. Implements method ALN.

# Usage

```
add.relative.likd.node(mixture,aca,ind=1)
```

#### **Arguments**

mixture A compiled DNAmixture object

aca Child's genotype profile as an allele count array

ind Index of contributor regarded as Parent (or Child): which 'unknown' contributor

are we modelling by amending his/her CPTs?

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#### Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Examples**

```
data(test2data)
# set threshold C
C<-0.001

mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db)
pars<-mixpar(rho=list(2),eta=list(100),xi=list(0.1),phi=list(c(U1=0.7,U2=0.3)))
baseline<-logL(mixD)(pars)

mixALN<-DNAmixture(list(epg),k=2,C=list(C),database=db,triangulate=FALSE,compile=FALSE)
cgtcaca<-gt2aca(mixALN,Cgt)
add.relative.likd.node(mixALN,cgtcaca,1)
log10LR<-(logLX(mixALN,
expr.make.findings(list(
list('Male',ind=1),
list('Rlikd',aca='cgtcaca',cgt='Cgt',evid='Revid')
))
)(pars)-baseline)/log(10)
cat('log10 LR',log10LR,'\n')</pre>
```

as.gt

Extract genotype profile for a single contributor from rGTs output

#### **Description**

Extract genotype profile for a single contributor from rGTs output

# Usage

```
as.gt(res,ind)
```

#### **Arguments**

| res | Output from rGTs |
|-----|------------------|
|     |                  |

ind Integer, which individual's genotype profile should be extracted

#### Value

Data frame, genotype profile for selected individual, for format see formats.

8 checkpeaks

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

checkpeaks Check and modify database to have positive frequencies for all observed peaks/alleles

## **Description**

Check whether database has positive frequencies for all peaks/alleles observed in epg and genotype profiles, and optionally modify db by addition of small positive frequencies so that it does, followed by renormalisation of frequencies for each allele to sum to 1.

#### Usage

```
checkpeaks(x,db,fix=0.003)
```

## **Arguments**

x data frame, the epg or genotype profile; see formats.

db data frame, the db; see formats.

fix numeric: if positive, increment to db frequency for each identified discrepant

peak

#### Value

```
(possibly modified) db
```

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

```
data(test2data)

db<-checkpeaks(epg,db)

db<-checkpeaks(Cgt,db)

Xgt<-data.frame(marker=c('D10','D12'),allele1=c(8,13),allele2=c(13,10))
db<-checkpeaks(Xgt,db)
db</pre>
```

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| convertIBD Convert relationship information to IBD pattern distribution | convertIBD | Convert relationship information to IBD pattern distribution |
|---|------------|--|
|---|------------|--|

#### **Description**

Construct IBD pattern distribution from one of several alternative representations of multi-person condensed coefficients of identity

#### Usage

```
as.IBD(x='sibs', targets=NULL, ped=FALSE)
convertIBD(x='sibs', targets=NULL, ped=FALSE)
```

#### **Arguments**

| x       | A string, a vector of length 3 or 9, a list with components pr and patt, or a list with two components, a pedigree and a vector of target id's; see Details |
|---------|---|
| targets | character vector of individual tags   |
| ped     | logical, should complete pedigree be added as an attribute to the output, if available?   |

#### **Details**

Possible formats for the input x are:

- 1. certain verbal mnemonics; currently one of the following (or an unambiguous partial match): c('sibs','parent-child','half-sibs', 'cousins','half-cousins','second-cousins', 'double-first-cousins', 'quadruple-half-first-cousins', '3cousins-cyclic','3cousins-star','trio')
- 2. a vector of 3 kappas
- 3. a vector of 9 Deltas
- 4. a list with matrix or vector valued component patt, with or without component pr
- 5. a list with 2 components, the first being a pedigree in the sense of the pedtools package, the second a vector of target id's
- 6. a 3-column character matrix of individual tags, each row denoting a child/mother/triple an alternative compact representation of a pedigree

#### Value

IBD pattern distribution as a list with components pr and patt

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### See Also

```
pedtools, formats
```

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#### **Examples**

```
data(test2data)

IBD<-convertIBD('parent-child')

IBD<-convertIBD(c(0.5,0.5,0.0))</pre>
```

delete.DQnodes

Delete D and Q dummy nodes from all Bayes nets in mixture

#### **Description**

Delete D and Q dummy nodes and associated edges from all Bayes nets in mixture, to save space; these nodes would only be needed for specific follow-up analyses

## Usage

```
delete.DQnodes(mixture,which="DQ")
```

#### **Arguments**

mixture A compiled DNAmixture object

which character string

## **Details**

The function removes the D and/or Q nodes from the DNA mixture object, depending on whether which includes "D", "Q" or both

## Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

```
data(test2data)
data(NGMDyes)

mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db,dyes=list(NGMDyes),
triangulate=FALSE,compile=FALSE)
delete.DQnodes(mixD)
replace.tables.for.UAF(mixD,40)
size(mixD)</pre>
```

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emperors

IBD pattern distribution in the Iulius-Claudius pedigree

## **Description**

IBD pattern distribution in the Iulius-Claudius pedigree

# Usage

```
data("emperors")
```

#### **Format**

IBD object

#### See Also

formats.

# **Examples**

```
data(emperors)
```

expr.make.findings

Coding additional findings as expression

# Description

Returns an expression that will be evaluated in logL.UKX whenever the likelihood of the model is calculated using the current method, and encodes the additional findings needed to implement the method; the details of the model and the extra information needed are held in the list z

## Usage

```
expr.make.findings(z)
```

#### **Arguments**

z A list specifying the additional findings; for the format, see Details

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#### **Details**

Each component of the list z is a list encoding a particular type of additional finding: the first component of this (sub-)list being a character string specifying the type of finding, and the remainder of its components being named parameters giving details of the finding. The types of finding and the valid parameters of each are as follows:

Male ind: index of relevant contributor: which 'unknown' contributor are we modelling by amending his CPTs?

Female ind: index of relevant contributor

Rlikd aca: allele count array, cgt: character string naming genotype profile data frame, evid: character string naming list with one component for each marker, whose value is the evidence

Aca ind: index of relevant contributor, aca: allele count array

Caca ind: index of relevant contributor, aca: allele count array

Denom no parameters

If z is NULL, then there are no additional findings.

#### Value

Expression encoding the additional findings.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

formats

**Formats** 

#### Description

Formats for data objects in KinMix and KinMixLite

#### **Formats**

An **allele frequency database** is a data frame containing variables marker, allele and frequency (character, numeric and numeric respectively).

A **mixture profile** is a data frame containing variables marker, allele and height (character, numeric and numeric respectively).

A **genotype profile** is a data frame containing variables marker, allele1 and allele2 (character, numeric and numeric respectively).

Examples of these 3 data formats are objects db, epg and Cgt, respectively, in test2data.

A **allele count array** is an alternative format for a genotype as a named list of vectors, one for each marker. Each vector gives the number of each allele in the genotype, with the alleles listed in the order in which they appear in the data component of the relevant mixture object.

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An **IBD pattern distribution** or **IBD object** is a list with components pr (a numerical vector) and patt (an integer matrix with nrow(patt)==length(pr) and an even number of columns). The elements of pr are the probabilities of the IBD patterns in the corresponding rows of patt. Adjacent pairs of columns encode the genotypes of different individuals; equal elements in any row determine equality of the alleles; different elements denote independent draws from the gene pool. If the component pr is missing, functions rpt.IBD and rpt.typed.relatives assume the probabilities are equal.

# Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

gt2aca

Converts genotype profiles to allele count arrays

## **Description**

Returns list of vectors of allele counts corresponding to genotype profile in gt

#### Usage

```
gt2aca(mixture,gt,eps=0)
```

## **Arguments**

| mixture | A compiled DNAmixture object  |
|---------|---|
| gt      | Genotype profile as a data frame containing variables marker, allele1 and allele2   |
| eps     | If non-zero, the function creates the output allele count arrays in a different format, that mitigates subsequent propagation errors in some situations. Instead of a vector of allele counts, each element of the list is a matrix with 3 columns, corresponding to allele counts 0, 1 and 2, with entries 1 or eps. |

#### Value

Returns list of vectors of allele counts. The log probability for the genotype is returned in its attribute 'logGt'.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

logL.UKX

#### **Examples**

```
data(test2data)
data(NGMDyes)

# set threshold C
C<-0.001

mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db,dyes=list(NGMDyes))
cgtcaca<-gt2aca(mixD,Cgt)

print(Cgt)
print(cgtcaca)</pre>
```

intoMix

Edit output from rGTs to omit individuals with NA amounts of DNA

# **Description**

Edit output from rGTs to omit individuals with NA amounts of DNA

#### Usage

```
intoMix(res)
```

#### **Arguments**

res

Output from rGTs

#### Value

The edited data structure

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

logL.UKX

Evaluates mixture log likelihood for unknown contributors with extra findings

# Description

Replacement for logL.UK in DNAmixtures that calls extra.findings immediately before propagating all findings and returning the normalising constant for the network.

logL.UKX

## Usage

```
logL.UKX(mixture, expr.extra.findings, initialize = FALSE)
```

# Arguments

#### Value

The log likelihood.

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### See Also

See also logL.UK.

```
data(test2data)

# set threshold C
C<-0.001

pars<-mixpar(rho=list(2),eta=list(100),xi=list(0.1),phi=list(c(U1=0.9,U2=0.1)))

mixMBN<-DNAmixture(list(epg),k=2,C=list(C),database=db,triangulate=FALSE,compile=FALSE)
cgtcaca<-gt2aca(mixMBN,Cgt)
add.child.meiosis.nodes(mixMBN,cgtcaca,1)
logL.UKX(mixMBN,
expr.make.findings(list(
list('Male',ind=1),
list('Caca',aca='cgtcaca')
)))(pars)</pre>
```

logLX

logLX

Evaluates mixture log likelihood when extra findings present

## **Description**

Replacement for logL in DNAmixtures that calls calls LogL.UKX instead of logL.UK.

## Usage

```
logLX(mixture, expr.extra.findings, presence.only = FALSE, initialize = FALSE)
```

#### **Arguments**

#### Value

The log likelihood.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### See Also

See also logL.

```
data(test2data)
# set threshold C
C<-0.001
mixD<-DNAmixture(list(epg), k=2,C=list(0.001),database=db)
pars<-mixpar(rho=list(2),eta=list(100),xi=list(0.1),phi=list(c(U1=0.7,U2=0.3)))
baseline<-logL(mixD)(pars)
mixMBN<-DNAmixture(list(epg),k=2,C=list(C),database=db,triangulate=FALSE,compile=FALSE)
cgtcaca<-gt2aca(mixMBN,Cgt)
add.child.meiosis.nodes(mixMBN,cgtcaca,1)
log10LR<-(logLX(mixMBN,</pre>
```

loop.rpt.IBD

```
expr.make.findings(list(
list('Male',ind=1),
list('Caca',aca='cgtcaca')
))
)(pars)-attr(cgtcaca,'logGt')-baseline)/log(10)
cat('log10 LR',log10LR,'\n')
```

loop.rpt.IBD

Analysis of DNA mixtures with familial relationships by looping

#### **Description**

Analysis of DNA mixtures with familial relationships, by looping over traces, markers, and IBD patterns, to reduce total BN table size, at some price in execution time

#### Usage

```
loop.rpt.IBD(listdata, pars, IBD, typed.gts = NULL, inds = 1,
    jtyped = ncol(IBD$patt)/2 - length(typed.gts) + seq_along(typed.gts),
    jcontr = seq_along(inds), targets = NULL, contribs,
    quiet=FALSE, verbose=FALSE, presence.only=FALSE, ...)
```

#### **Arguments**

listdata as in call to DNAmixture

pars parameter structure, in mixpar format

IBD multi-person coefficients of identity, in any of the formats accepted by convertIBD

typed.gts, inds, jtyped, jcontr, targets, contribs, quiet

as in call to rpt.IBD

verbose should per-marker and overall log10LR's be reported?

presence.only Set to TRUE to ignore peak heights and evaluate the likelihood function con-

sidering peak presence and absence (heights above and below threshold) only.

Defaults to FALSE.

... other arguments to DNAmixture, particularly including k, C, database

#### Value

The value of the overall log10 LR, and the contributions of individual markers in the form of a vector-valued attribute 'log10LR', are returned invisibly; individual marker/pattern values are also printed out.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

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#### **Examples**

```
data(test2data)
data(NGMDyes)
C<-0.001
## Fit 3-person mixture - baseline model
mixD<-DNAmixture(list(epg),k=3,C=rep(list(C),length(list(epg))),database=db)</pre>
pars3 < mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.6, U2=0.3, U3=0.1)))
baseline3<-logL(mixD)(pars3)</pre>
size(mixD)
## Fit 3-person mixture - in which U1 and U2 have a parent-child relationship
mixD<-DNAmixture(list(epg),k=3,C=rep(list(C),length(list(epg))),database=db,
triangulate=FALSE,compile=FALSE)
delete.DQnodes(mixD)
rpt.IBD(mixD,IBD=c(0,1,0),typed.gts=list(),inds=1:2,jtyped=NULL)
size(mixD)
log10LR<-(logL(mixD)(pars3)-baseline3)/log(10)</pre>
cat('log10 LR',log10LR,'\n')
## the same analysis by loop.rpt.IBD
listdata<-list(epg)</pre>
print(loop.rpt.IBD(listdata,pars3,IBD=c(0,1,0),
k=3,C=rep(list(C),length(listdata)),database=db,
typed.gts=list(),inds=1:2,jtyped=NULL))
```

make.profile

Convert genotype profile to reference profile format

#### **Description**

Convert genotype profile to reference profile format

#### Usage

```
make.profile(gt,name='K')
```

#### **Arguments**

gt genotype profile

name character string used to name profile in output data frame

#### Value

data frame containing reference profile

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#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Examples**

```
data(test2data)
S1prof<-make.profile(S1gt,'S1')
C<-0.001
mixD<-DNAmixture(list(epg),k=3,K='S1',reference.profile=S1prof,C=list(C),database=db)</pre>
```

mixMLX

Maximises mixture likelihood when extra findings present

#### **Description**

Replacement for mixML in DNAmixtures that calls logLX instead of logL.

## Usage

```
mixMLX(mixture, expr.extra.findings, pars, constraints = NULL, phi.eq = FALSE, val = NULL, trace = FALSE, order.unknowns = TRUE, initialize = FALSE, ...)
```

#### **Arguments**

```
mixture
                  Compiled DNAmixture object.
expr.extra.findings
                 expression containing the extra findings
pars
                 Parameters, in mixpar format.
constraints
                 as in mixML
phi.eq
                  as in mixML
val
                  as in mixML
trace
                  as in mixML
order.unknowns as in mixML
initialize
                 should all entered evidence be removed from the networks in mixture
                  as in mixML
```

#### Value

```
A list containing
```

```
mle The (suggested) MLE.lik The log of the likelihood (log e).as well as the output from the optimisation.
```

20 pedigreeIBD

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### See Also

See also mixML.

#### **Examples**

```
data(test2data)
# set threshold C
C<-0.001
mixD<-DNAmixture(list(epg),k=2,C=list(C),database=db)
# find MLE's and maximised likelihood
# adding evidence individual 1 is Male
expr.extra.findings<-expr.make.findings(list(list('Male',ind=1)))
startpar<-mixpar(rho=list(60),eta=list(24),xi=list(0.16),phi=list(c(U1=0.75,U2=0.25)))
mlDM<-mixMLX(mixD,expr.extra.findings,startpar,trace=FALSE)
pars<-mlDM$mle
cat('\nBaseline model maximised likelihood:',mlDM$lik,'\n')
cat('and MLEs:\n')
print(mlDM$mle)</pre>
```

pedigreeIBD

Construct IBD pattern distribution from pedigree

#### **Description**

Construct IBD pattern distribution from a pedigree and a target list of individuals

#### Usage

```
pedigreeIBD(x, targets, cond = TRUE, ped=FALSE, quiet = TRUE, verbose = FALSE)
```

#### **Arguments**

| х       | A pedigree in pedtools format   |
|---------|---|
| targets | Character vector, some or all of the individual identifiers in the pedigree x           |
| cond    | should IBD pattern be condensed?  |
| ped     | logical, should complete pedigree be added as an attribute to the output, if available? |
| quiet   | should resulting IBD pattern distribution be printed?                                   |
| verbose | should trace information be printed?  |

plot.IBD 21

#### **Details**

This function computes the multi-person condensed coefficients of identity for an arbitrary set of individuals, in the sparse notation of the IBD pattern distribution of Green and Vigeland (2019).

#### Value

IBD pattern distribution as a list with components pr and patt

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### References

*Multi-person condensed coefficients of identity*, by Peter J. Green and Magnus Dehli Vigeland, University of Bristol technical report, 2019.

#### See Also

```
pedtools, formats
```

#### **Examples**

```
require(ribd)
id<-c('gf','gm','b1','b2','m','c')
fid<-c(0,0,'gf','gf',0,'b1')
mid<-c(0,0,'gm','gm',0,'m')
sex<-c(1,2,1,1,2,0)
x<-ped(id,fid,mid,sex)

IBD<-pedigreeIBD(x,c('m','c','b1','b2'))</pre>
```

plot.IBD

Plot IBD patterns and pattern distributions

#### **Description**

Plot IBD patterns and pattern distributions

#### **Usage**

```
## S3 method for class 'IBD'
plot(x,labels=NULL,probs=NULL,order=NULL,colrs=c('black','red','blue'),
digits=3,nr=ceiling(sqrt(np)),...)
```

plot.IBD

# Arguments

| X      | A matrix whose rows are IBD patterns, or a list whose components are patt, such a matrix, together with pr, a vector of the corresponding probabilities   |
|--------|---|
| labels | Vector of numerical or character labels for the patterns, if NA, labels are constructed from the patterns by catenation, if NULL, the labels are not displayed.   |
| probs  | Vector of probabilities of the patterns, if not provided as a component of pattern; if NULL, the probabilities are not displayed.   |
| order  | A character string, partially matched using pmatch to one of 'pattern', 'probs', or 'labels', requesting ordering diagram accordingly (in the case of probs in decreasing order, <b>or</b> a numeric, complex, character or logical vector of length the number of patterns, requesting ordering by this variable, <b>or</b> NULL (the default), requesting no re-ordering. |
| colrs  | A vector of colours: ties in the ordering variable are indicated by coloured groups, with colours chosen cyclically from this vector.   |
| digits | Integer, overwriting default number of significant digits for probs   |
| nr     | Integer, overwriting default number of rows for plotted array, default a rounding up of the square root of the number of patterns.  |
|        | additional arguments to plot  |

#### Value

No value is returned, the function is called for its side effect, a plot on the current display device.

# Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

```
require(ribd)
data(emperors)

plot.IBD(convertIBD('3cousins-star'),order='probs',col=c('blue','red','black'))

plot(attr(emperors,'ped'))

o<-order(emperors$pr)[1:12]
plot.IBD(emperors$patt[o,],probs=emperors$pr[o],labels=NA,order='probs')</pre>
```

protected 23

|--|

## **Description**

Attempts to catch numerical erros in evaluating the expression x, delivering a default result instead of NaN or other failures

#### Usage

```
protected(x,default=-Inf)
```

# Arguments

x expression to be evaluated, typically the log-likelihood of a modified mixture

model

default value to be delivered if numerical errors are encountered

#### Value

Returns -Inf in case of error, otherwise the value of x

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Description**

Attempts to catch numerical errors in maximum likelihood computation, by replacing logL values by a default value instead of NaN or other failures

# Usage

```
protected.mixML(mixture, pars, constraints = NULL, phi.eq = FALSE, val = NULL,
    trace = FALSE, order.unknowns = TRUE, default=-999999, ...)
```

24 require.compiled

#### **Arguments**

mixture A DNAmixture object.

pars A mixpar parameter used as a starting value for the optimisation.

constraints Equality constraint function as function of an array of parameters.

phi.eq Should the mixture proportions be the same for all traces? Defaults to FALSE.

val Vector of values to be satisfied for the equality constraints.

trace Print the evaluations of the likelihood-function during optimisation?

order.unknowns Should unknown contributors be ordered according to decreasing contributions?

Defaults to TRUE.

Further arguments to be passed on to solnp.

default value of logL to be used if numerical errors are encountered

#### Value

A list containing

mle The (suggested) MLE.

1ik The log of the likelihood (log e).

as well as the output from the optimisation.

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

require.compiled

Force compilation of all BNs in a DNA mixture model

## **Description**

Scan all Bayes nets in mixture, and compile any that are not already compiled

#### Usage

require.compiled(mixture)

# Arguments

mixture A DNAmixture object

#### Value

No value is returned, the function is called for its side effect

rGTs 25

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Examples**

```
data(test2data)
data(NGMDyes)
mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db,dyes=list(NGMDyes),</pre>
triangulate=FALSE,compile=FALSE)
replace.tables.for.UAF(mixD,40,compile=FALSE)
require.compiled(mixD)
```

| rGTs | Simulate random genotype profiles and DNA samples for related indi- |
|------|---|
|      | viduals   |

## **Description**

Simulate random genotype profiles and DNA samples for arbitrarily related individuals

#### Usage

```
rGTs(nreps, IBD, db, DNA, sex=rep(0, ncontr), nU=0)
```

#### **Arguments**

| nreps | Integer, number of replicates  |
|-------|--|
| IBD   | Specification of relationships, as in convertIBD   |
| db    | Data frame, database of alleles and their frequencies, for each marker; for format, see formats. |
| DNA   | Integer vector, numbers of DNA cells for the respective individuals, can be NA                   |
| sex   | Integer vector, sex of the respective contributors: 1=male, 2=female, 0=unspecified              |
| nU    | Integer, include also this number of unrelated individuals                                       |

#### **Details**

Genotype profiles are generated randomly, using the allele frequency database db, under the relationships specified by the IBD argument. In accordance with the underlying biology, allele values for the AMEL marker (if this is one of the markers included) are not influenced by relationships with other individuals; however they are influenced by the sex of the individuals, where this is known. Information on sex can be specified by the optional argument sex: a male is given the profile X-Y, a female X-X, and an individual with unspecified sex X-X or X-Y with equal probabilities.

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## Value

Data frame with variables Sim, Sample.name, Marker, Allele, and DNA, suitable for input to simExtraction, etc. See package pcrsim.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Examples**

```
data(test2data)
data(NGMDyes)
```

rni

Random number initialiser supporting spontaneous replication

# Description

Random number initialiser supporting spontaneous replication

#### Usage

rni(seed=0)

## **Arguments**

seed

Integer, seed

#### **Details**

This is a convenience front end to set.seed. A non-zero value of seed is passed directly to set.seed. Given a zero value (the default), the function calls Sys.time to generate an unpredictable starting value – but the value ultimately passed to set.seed is both output using cat and returned invisibly, so can be used for unanticipated replica runs of a simulation.

#### Value

Non-zero seed value that can be used to reproduce run subsequently

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

rpt.AMEL 27

#### **Examples**

rni(0)
runif(6)
rni(0)
runif(6)
rni(3456)
runif(6)
rni(3456)
runif(6)
keep<-rni(0)
print(keep)
runif(6)
rni(keep)
runif(6)</pre>

rpt.AMEL

Replace CPTs for AMEL marker in a DNA mixture

#### **Description**

Used after a call to DNAmixture with compile=FALSE, triangulate=FALSE, this function replaces the CPTs for the genotype allele count arrays for the AMEL marker in a DNA mixture to specify sex of contributors

# Usage

```
rpt.AMEL(mixture,sex,compile=TRUE)
```

## **Arguments**

mixture A DNAmixture object

sex Integer vector, sex of each contributor

compile Logical, should BN be compiled after modification?

#### **Details**

The sex of each contributor is coded as in pedtools, namely 0=unspecified, 1=male, 2=female.

# Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

28 rpt.IBD

#### **Examples**

rpt.IBD Replacing CPTs for selected mixture contributors with familial relationships

#### **Description**

Used after a call to DNAmixture with compile=FALSE, triangulate=FALSE, this function replaces the CPTs for the genotype allele count arrays for specified mixture contributors by those representing the specified relationship with each other and typed relatives

#### Usage

```
rpt.IBD(mixture, IBD="parent-child", typed.gts = NULL, inds = 1,
jtyped = ncol(IBD$patt)/2 - length(typed.gts) + seq_along(typed.gts),
jcontr = seq_along(inds),
targets=attr(IBD, 'targets'), contribs=NULL, quiet=FALSE, all.freq = NULL, compile = TRUE)

rpt.typed.relatives(mixture, IBD="parent-child", typed.gts = NULL, inds = 1,
jtyped = ncol(IBD$patt)/2 - length(typed.gts) + seq_along(typed.gts),
jcontr = seq_along(inds),
targets=attr(IBD, 'targets'), contribs=NULL, quiet=FALSE, all.freq = NULL, compile = TRUE)

rpt.typed.child(mixture, aca, ind=1)

replace.Ui.tables(mixture, aca, ind=1)

rpt.typed.parents(mixture, Mgt, Fgt, ind=1, compile=TRUE)

rpt.typed.relative(mixture, Rgt, IBD=c(0.25,0.5,0.25), ind=1, compile=TRUE)
```

#### Arguments

| mixture   | DNAmixtures object created by previous call to DNAmixture with triangulate=FALSE,compile=FALSE  |
|-----------|---|
| IBD       | relationships between the specified individuals, as multi-person condensed coefficients of identity, in one of several representation; see Details.   |
| typed.gts | list of 0 or more genotypes of relatives; the components of this list must be named (with the id's of the relevant individuals) if targets and contribs are used to code the correspondences (see Details). |

rpt.IBD

inds vector of 1 or more integers: which 'unknown' contributors are we modelling by amending their CPTs? The elements should be listed in the same order as the corresponding pairs of columns of the IBD patterns in IBD jtyped indices of pairs of columns of IBD\$patt that correspond to the typed relatives (if any); default the last length(typed.gts) pairs of columns indices of pairs of columns of IBD\$patt that correspond to the relevant mixture jcontr contributors; default the first length(inds) pairs of columns targets Character vector of the tags of the individuals referred to in IBD contribs Character vector of the tags of the individuals included in the mixture, in order quiet should calculated values of inds, jtyped and jcontr be reported? all.freq alternative allele frequency database(s), see Details. compile logical flag: should mixture object be compiled on exit? ind as inds, used above when only one allowed aca, Mgt, Fgt, Rgt

individual genotypes, as allele count arrays

#### **Details**

In using rpt.IBD or rpt.typed.relatives (which is identical), the correspondence between mixture contributors, specified relationships, and typed genotype profiles should be specified **either** (preferably) using targets, contribs and through the names of the components in typed.gts, **or** (to be deprecated) with inds, jcontr and jtyped: the two representations should not be mixed up. If either targets or contribs specified, the former representation is assumed.

Special cases are treated slightly more efficiently: rpt.typed.child: single contributor, single typed relative, parent or child; rpt.typed.parents: single contributor, both parents typed; rpt.typed.relative: single contributor, single typed relative.

Note that IBD\$patt always has an even number of columns, two for each individual in the joint relationship specified; jtyped and jcontr are vectors of indices of these individuals, i.e. to pairs of adjacent columns of IBD\$patt.

Multiple functions in this group can validly be called sequentially (with all but the last having compile=FALSE) providing they reference different sets of contributors among the targets, **and** that these sets are conditionally independent given the typed genotypes specified.

There are multiple valid representations for relationships in the argument IBD – as an IBD pattern distribution, via a pedigree, or. in the case of just two individuals. via either a vector of 3 kappas or 9 Deltas (Jacquard's condensed coefficients of identity). For full details, see convertIBD. If IBD is missing, the default value represents parent-child.

In the interests of upward compatibility, in rpt.typed.child and replace.Ui.tables (which are identical), the argument Cgt can be given as either a genotype profile data frame, or an allele count array.

By default, the allele frequency database used for the founding genes is that used when the mixture object is created, in an earlier call to DNAmixture. A non-null value for the all.freq argument allows the user to specify alternative database (s) for the founding genes. If its value is an allele frequency database (in the format specified in formats) then that database is used for all founding genes; if the value of the argument is a list of such databases, then component k of the list is used for

30 rpt.IBD

allele frequencies for the founding gene labelled k in the IBD argument. Note that this option allows modelling of mixtures where different contributors are drawn from different populations, whether or not there are relationships among individuals.

#### Value

Vector of marker-specific probabilities of the typed genotypes.

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

```
data(test2data)
data(NGMDyes)
## Fit 2-person mixture - baseline model
mixD<-DNAmixture(list(epg), k=2, C=list(0.001), database=db)
pars < mixpar(rho=list(2), eta=list(100), xi=list(0.1), phi=list(c(U1=0.7, U2=0.3)))
baseline<-logL(mixD)(pars)</pre>
## Fit 2-person mixture model in which contributor 1 is parent of a typed individual Cgt
mixD<-DNAmixture(list(epg), k=2,C=list(0.001),database=db,triangulate=FALSE,compile=FALSE)
rpt.IBD(mixD,,list(Cgt))
log10LR<-(logL(mixD)(pars)-baseline)/log(10)
cat('log10 LR',log10LR,'\n')
## Fit 2-person mixture model in which contributor 1 is father of a typed individual Cgt
## with mother Mgt
mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db,triangulate=FALSE,compile=FALSE)</pre>
rpt.IBD(mixD,,list(Mgt,Cgt))
log10LR<-(logL(mixD)(pars)-baseline)/log(10)</pre>
cat('log10 LR',log10LR,'\n')
## Fit 2-person mixture model in which contributors are two parents of a child with
## genotype Cgt, and a parent of one of them has genotype Rgt. Note the encoding of allele
## labels to reduce the complexity of the IBD pattern distribution IBD.
IBD<-list(patt=rbind(c(1,3,2,4,1,2,1,5),c(1,3,2,4,1,2,3,5)))
mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db,triangulate=FALSE,compile=FALSE)</pre>
rpt.IBD(mixD,IBD,list(Cgt,Rgt),1:2)
log10LR<-(logL(mixD)(pars)-baseline)/log(10)
cat('log10 LR',log10LR,'\n')
## the same, with individuals and relationships denoted by character tags
mixD<-DNAmixture(list(epg), k=2,C=list(0.001),database=db,triangulate=FALSE,compile=FALSE)
```

rpt.UAF

```
 rpt.IBD(mixD,IBD,list(c=Cgt,gf=Rgt),targets=c('f','m','c','gf'),contribs=c('f','m')) \\ log10LR<-(logL(mixD)(pars)-baseline)/log(10) \\ cat('log10 LR',log10LR,'\n') \\
```

rpt.UAF

Replace CPTs in a DNA mixture to model uncertain allele frequencies

#### **Description**

Replace CPTs in a DNA mixture to model uncertainty in allele frequencies

# Usage

```
replace.tables.for.UAF(mixture, M, compile = TRUE)
rpt.UAF(mixture, M, compile = TRUE)
```

# **Arguments**

mixture DNAmixtures object created by previous call to DNAmixture with

triangulate=FALSE,compile=FALSE

M Size of allele frequency database

compile logical flag: should mixture object be compiled on exit?

#### Value

No value is returned, the function is called for its side effect

#### Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

```
data(test2data)
data(NGMDyes)

## Fit 2-person mixture - baseline model

mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db)
pars<-mixpar(rho=list(2),eta=list(100),xi=list(0.1),phi=list(c(U1=0.7,U2=0.3)))
baseline<-logL(mixD)(pars)

## Fit 2-person mixture model under assumption that database size was only 40

mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db,triangulate=FALSE,compile=FALSE)
replace.tables.for.UAF(mixD,40)
log10LR<-(logL(mixD)(pars)-baseline)/log(10)
cat('log10 LR',log10LR,'\n')</pre>
```

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size

Calculate and display total size of BN tables for a DNA mixture

# **Description**

Calculate and display total size of BN tables for a DNA mixture

# Usage

```
size(mixture)
```

## **Arguments**

mixture

A compiled DNAmixture object

#### Value

Returns total size, typically to be printed by bespoke method

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

## **Examples**

```
data(test2data)
data(NGMDyes)

## Fit 2-person mixture - baseline model

mixD<-DNAmixture(list(epg),k=2,C=list(0.001),database=db)
size(mixD)</pre>
```

test2data

Small data set for demonstrating some capabilities of KinMix and Kin-MixLite

# Description

Small test data set (2 markers with 4 or 5 alleles each, plus AMEL), for demonstrating some capabilities of KinMix and KinMixLite

# Usage

```
data("test2data")
```

wlr 33

## **Format**

Data objects for demonstrating KinMix: epg (DNAmixtures peak height data), db (DNAmixtures allele frequency database), and Cgt, Fgt, Mgt, Rgt, S1gt, S2gt potential relative genotype data frames.

## **Examples**

```
data(test2data)
```

wlr

Computes paternity LR using WLR method

# Description

Computes overall LR from Ugt-specific LR's using estimated Ugt genotype profile in sep corresponding to contributor i in the mixture as Father; uses Child genotype information in Cgt data.frame and optionally Mother's genotype in Mgt. Implements method WLR.

# Usage

```
wlr(sep, Cgt, db, ind=1, Mgt=NULL)
```

# Arguments

| sep | Separation, a list of configurations of genotypes for some or all unknown contributors, output by map.genotypes. |
|-----|--|
| Cgt | Child's genotype profile as a data frame containing variables marker, allele1 and allele2                        |
| db  | Allele frequency database  |
| ind | Index of contributor regarded as Father  |
| Mgt | (optionally) Mother's genotype profile as a data frame containing variables marker, allele1 and allele2          |

## Value

Returns LR for paternity

## Author(s)

Peter Green (P.J.Green@bristol.ac.uk)

#### See Also

See also map.genotypes.

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```
data(test2data)
data(NGMDyes)

# set threshold C
C<-0.001

pars<-mixpar(rho=list(2),eta=list(100),xi=list(0.1),phi=list(c(U1=0.9,U2=0.1)))

mixWLR<-DNAmixture(list(epg),k=2,C=list(C),database=db,dyes=list(NGMDyes))
setPeakInfo(mixWLR,pars)
sepWLR<-map.genotypes(mixWLR,type="all",pmin=0.001,U=1)
LR<-wlr(sepWLR,Cgt,db)
cat('\nWLR LR:',LR,'; log10(LR):',log10(LR),'\n')</pre>
```

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