Package ‘DNAtools’

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Title Tools for Analysing Forensic Genetic DNA Data
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       the multicore/CPU implementation.
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Description Computationally efficient tools for comparing all pairs of profiles
       in a DNA database. The expectation and covariance of the summary statistic
       is implemented for fast computing. Routines for estimating proportions of
       close related individuals are available. The use of wildcards (also called F-
       designation) is implemented. Dedicated functions ease plotting the results.
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DNAtools-package ................................................. 2
dbCollapse ....................................................... 3
dbCompare ........................................................ 4
dbExample .......................................................... 5
dbExpect ............................................................ 6
dbSimulate ........................................................... 7
dbVariance ........................................................... 9
DNAtools-package

Description

Computational efficient tools for comparing all pairs of profiles in a DNA database. The expectation and covariance of the summary statistic is implemented for fast computing. Routines for estimating proportions of close related individuals are available. The use of wildcards (also called F-designation) is implemented. Dedicated functions ease plotting the results.

Details

Package: DNAtools
Type: Package
Version: 0.1
Date: 2014-08-25
License: GPL (>= 2)

dbCompare: Compares make all \(\frac{n(n-1)}{2}\) pairwise comparisons between profiles of a database with \(n\) DNA profiles. dbExpect: Computes the expected number of matching and partial matching loci for a given number of profiles in a database. dbVariance: Calculates the associated covariance matrix.

Author(s)

Torben Tvedebrink <tvede@math.aau.dk> and James Curran <j.curran@auckland.ac.nz> wrote the package. Mikkel Meyer Andersen <mikl@math.aau.dk> assisted on the multicore/CPU implementation.

Maintainer: Torben Tvedebrink <tvede@math.aau.dk>
References


Read the vignette: vigette("DNAtools")

Examples

```r
## Not run:
data(dbExample)
dbCompare(dbExample, hit=5, trace=TRUE)
## End(Not run)
```

---

**dbCollapse**

*Collapse m/p output to vector*

**Description**

Collapse a m/p-matrix from dbCompare/dbExpect to a vector.

**Usage**

```r
dbCollapse(x)
```

**Arguments**

- `x` Either an object of class "dbcompare" (result from dbCompare) or "matrix".

**Details**

Collaps a m/p-matrix from dbCompare/dbExpect to a vector with entry i being the sum of all entries from m/p-matrix satisfying 2*m+p=i.

**Value**

A vector of length 2*max(m)+1 with entries begin the sum of entries i in m/p-matrix satisfying i=2*m+p.

**Author(s)**

Torben Tvedebrink
Examples

```r
## Not run:
data(dbExample)
res <- dbCompare(dbExample, hit=5, trace=TRUE)
dbCollapse(res) ## same as dbCompare(dbExample, hit=5, trace=TRUE, collapse=TRUE)
## End(Not run)
```

---

**dbCompare**

**Compare DNA profiles**

**Description**

Compare DNA profiles

**Usage**

```r
dbCompare(x, profiles=NULL, hit=7, trace=TRUE, vector=FALSE, collapse=FALSE,
wildcard=FALSE, wildcard.effective=FALSE, wildcard.impose=FALSE,
Rallele=FALSE, threads=2)
```

**Arguments**

- **x** Database with DNA profiles. The database format is expected to be a data frame with each column containing an allelic number such that for each DNA marker there are two columns in the data frame. See `data(dbExample)` for an example of the format.

- **profiles** One or more profiles to be compared with all profiles in the database. Input is a vector, matrix or data frame of same length/width as a row in the database `x`. If profiles is non-null only one CPU will be used. In case threads>1 a warning will be given but computations performed using single core.

- **hit** The number of matching loci for further investigation

- **trace** Shows a progress bar

- **vector** Logical. Whether the result should be returned as vector or a matrix. Note if 'collapse' is TRUE vector is ignored.

- **collapse** Logical (default FALSE). If TRUE the (m,p)-matrix will be collapsed into a (2*m+p)-vector containing the total number of matching alleles.

- **wildcard** Use the wildcard comparing.

- **wildcard.effective** Compare result of wildcard and no wildcard.

- **wildcard.impose** Force homozygous profiles (aa) to have wildcard (aF).

- **Rallele** Implementation of 'Rare allele'designation matching.

- **threads** The number of threads to use for performing comparisons in parallel for increased computation time. Use 0 for using the same number as the computer has CPU cores. NOTE: Only available on Linux and MacOS operating systems.
Details

Computes the distance between DNA profiles in terms of matching and partially-matching STR loci.

Value

Returns a matrix with the number of pairs matching/partially-matching at (i,j)-loci.

Author(s)

James Curran and Torben Tvedebrink. The multicore/CPU implementation was provided by Mikkel Meyer Andersen.

Examples

```r
## Not run:
data(dbExample)
dbCompare(dbExample,hit=5,trace=TRUE)
## End(Not run)
```

---

dbExample Simulated database with 1,000 individuals

Description

Database containing 1,000 simulated DNA profiles typed on ten autosomal markers.

Usage

dbExample

Format

A data frame with each row being a DNA profile and each column a part of a genetic marker. Note that homozygote profiles has the same allelic value in the two columns associated to the same marker.
**dbExpect**

*Expected value of cell counts in DNA database comparison*

**Description**

Computes the expected number of cell counts when comparing DNA profiles in a DNA database. For every pair of DNA profiles in a database the number of matching and partial matching loci is recorded. A match is declared if the two DNA profiles coincide for both alleles in a locus and a partial-match is recorded if only one allele is shared between the profiles. With a total of L loci the number of matching loci is 0,...,L and partial number of matches is 0,...,L-m, where m is the number of matching loci.

**Usage**

`dbExpect(probs, theta=0, k=(c(0,0,1), n=1, r=0, R=0, round=FALSE, na=TRUE, vector=FALSE, collapse=FALSE, wildcard=FALSE, no.wildcard=NULL, rare.allele=FALSE, no.rare.allele=NULL)`

**Arguments**

- `probs` List of vectors with allele probabilities for each locus
- `theta` The coancestery coefficient
- `k` The vector of identical-by-descent probabilities, k=(k2,k1,k0), where for full-siblings k=c(1,2,1)/4. The default is k=c(0,0,1) refering to unrelated individuals.
- `n` Number of DNA profiles in the database
- `r` The probability assigned to the rare alleles (see rare allele matching). If a vector must be of same length as `probs`.
- `R` The probability assigned to alleles shorter or longer than allelic ladder (see rare allele matching). If a vector must be of length 1 or 2, and if a list it must be same length as `probs`.
- `round` Whether or not the results should be rounded or not
- `na` Whether or not the off-elements should be returned as 0 or NA
- `vector` Whether or not the result should be returned as a matrix or vector. Note if 'collapse' is TRUE vector is ignored.
- `collapse` Logical (default FALSE). If TRUE the (m,p)-matrix will be collapased into a (2*m+p)-vector containing the total number of matching alleles.
- `wildcard` Should wildcards be used?
- `no.wildcard` Should 'w' wildcards be used?
- `rare.allele` Should rare allele matching be used?
- `no.rare.allele` Should 'r' rare allele loci be used?
**dbSimulate**

**Details**

Computes the expected cell counts using a recursion formula. See Tvedebrink et al (2011) for details.

**Value**

Returns a matrix (or vector, see above) of expected cell counts.

**Author(s)**

James Curran and Torben Tvedebrink

**References**


**Examples**

```r
## Not run:
## Simulate some allele frequencies:
freqs <- replicate(10, ( g = rgamma(n=10,scale=4,shape=3); g/sum(g)),
                 simplify=FALSE)
## Compute the expected number for a DB with 10000 profiles:
dbExpect(freqs,theta=0,n=10000)
## End(Not run)
```

---

### dbSimulate

**Simulate a DNA database**

**Description**

Simulates a DNA database given a set of allele probabilities and theta value. It is possible to have close relatives in the database simulated in pairs, such that within each pair the profiles are higher correlated due to close familial relationship, but between pairs of profiles the correlation is only modelled by theta.

**Usage**

```r
dbSimulate(probs,theta=0,n=1000,relatives=NULL)
```
Arguments

probs  List of allele probabilities, where each element in the list is a vector of allele probabilities.
theta  The coancestry coefficient
n      The number of profiles in the database
relatives  A vector of length 4. Determining the number of PAIRS of profiles in the database: (FULL-SIBLINGS, FIRST-COUSINS, PARENT-CHILD, AVUNCULAR). They should obey that 2*sum(relatives)<=n.

Details

Simulates a DNA database with a given number of DNA profiles (and possibly relatives) with a correlation between profiles governed by theta.

Value

A data frame where each row represents a DNA profile. The first column is a profile identifier (id) and the next 2*L columns contains the simulated genotype for each of the L loci. L is determined by the length of the list 'probs' with allele probabilities

Author(s)

James Curran and Torben Tvedebrink

Examples

```r
## Not run:
## Simulate some allele frequencies:

freq <- replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g)}, simplify=FALSE)
## Simulate a single database with 5000 DNA profiles:
simdb <- dbSimulate(freq,theta=0,n=5000)
## Simulate a number of databases, say N=50. For each database compute
## the summary statistic using dbCompare:
N <- 50
Msummary <- matrix(0,N,(length(freq)+1)*(length(freq)+2)/2)
for(i in 1:N)
  Msummary[i,] <- dbCompare(dbSimulate(freq,theta=0,n=1000),
                            vector=TRUE,trace=FALSE)
## Give the columns representative names:
dimnames(Msummary)[[2]] <- DNAtools:::dbCats(length(freq),vector=TRUE)
## Plot the simulations using a boxplot
boxplot(log10(Msummary))
## There might come some warnings due to taking log10 to zero-values (no counts)
## Add the expected number to the plot:
points(1:ncol(Msummary),log10(dbExpect(freq,theta=0,n=1000, vector=TRUE)),
      col=2,pch=16)
## End(Not run)
```
Description

Computes the covariance matrix for the cell counts when comparing DNA profiles in a DNA database. For every pair of DNA profiles in a database the number of matching and partial matching loci is recorded. A match is declared if the two DNA profiles coincide for both alleles in a locus and a partial-match is recorded if only one allele is shared between the profiles. With a total of L loci the number of matching loci is 0,...,L and partial number of matches is 0,...,L-m, where m is the number of matching loci. The expression is given by:

\[ \text{latex} \]

Usage

\[ \text{dbVariance(probs,theta=0,n=1,collapse=FALSE)} \]

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>probs</td>
<td>List of vectors with allele probabilities for each locus</td>
</tr>
<tr>
<td>theta</td>
<td>The coancestery coefficient. If a vector of different theta values are supplied a list of covariance matrices is returned. Note it is faster to give a vector of theta values as argument than calculating each matrix at the time.</td>
</tr>
<tr>
<td>n</td>
<td>Number of DNA profiles in the database. If n=1 is supplied a list of the components for computing the variance is returned. That is, the variance and two covariances on the right hand side of the equation above.</td>
</tr>
<tr>
<td>collapse</td>
<td>Logical, default FALSE. If TRUE the covariance matrix is collapsed such that it relates to ((2*m+p))-vectors of total number of matching alleles rather than ((m,p))-matrix.</td>
</tr>
</tbody>
</table>

Details

Computes the covariance matrix of the cell counts using a recursion formula. See Tvedebrink et al (2011) for details.

Value

Returns a covariance matrix for the cell counts.

Author(s)

James Curran and Torben Tvedebrink
References


Examples

```r
## Not run:
## Not run:
freqs <- replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g) }, simplify=FALSE)
## List of elements needed to compute the covariance matrix.
comps <- dbVariance(freqs, theta=0, n=1)
## Covariance for a DB with 1000 DNA profiles
cov1000 <- dbVariance(freqs, theta=0, n=1000)
## The result is the same as:
comps1000 <- choose(1000,2)*comps$V1 + 6*choose(1000,3)*comps$V2 + 6*choose(1000,4)*comps$V3
```

estimatePD

Estimate the drop-out probability based on number of alleles

Description

An inferior may to estimate the drop-out probability compared to using the peak heights from the electropherogram. However, to compare the performance with Gill et al. (2007) this implements a theoretical approach based on their line of arguments.

Usage

```r
estimatePD(n0, m, pnoa=NULL, probs=NULL, theta=0, noa.tabs=NULL, locuswise=FALSE)
```

Arguments

- `n0` Vector of observed allele counts - same length as the number of loci.
- `m` The number of contributors
- `pnoa` The vector of P(N(m)=n) for n=1,...,2Lm, where L is the number of loci and m is the number of contributors.
- `probs` List of vectors with allele probabilities for each locus
- `theta` The coancesttery coefficient
- `noa.tabs` If noa.tabs has been computed by the noaTabs-function for m, then this will speed up the computations.
- `locuswise` Logical. Indicating whether computations should be done locuswise.
freqEst

Details

Computes the P(D) that maximises equation (10) in Tvedebrink (2013).

Value

Returns the MLE of P(D) based on equation (10) in Tvedebrink (2013)

Author(s)

Torben Tvedebrink

References


Examples

```r
## Simulate some allele frequencies:
freqs <- structure(replicate(10, ( g = rgamma(n=10, scale=4, shape=3); g/sum(g)),
                           simplify=FALSE),.Names=paste("locus",1:10,sep="."))
## Assume 15 alleles are observed in a 2-person DNA mixture with 10 loci:
estimatePD(n0=15,m=2,probs=freqs)
```

freqEst

Simple allele frequency estimation

Description

Estimates allele frequencies from a database with DNA profiles

Usage

freqEst(x)

Arguments

x A database of the form ["id","locus1 allele1","locus1 allele2",...,"locusN allele 1","locusN allele2"].

Details

Computes the allele frequencies for a given database.
**Value**

Returns a list of probability vectors - one vector for each locus.

**Author(s)**

James Curran and Torben Tvedebrink

**Examples**

```r
data(dbExample)
freqEst(dbExample)
```

---

**Description**

Computes the \( \alpha_{m} \)-vectors and their weights, \( c(\alpha_{m}) \), in order to compute the exact distribution of the number of alleles in a \( m \)-person DNA mixture.

**Usage**

```r
noaTabs(alpha=NULL, m=2, weight=1)
```

**Arguments**

- **alpha**: If `noaTabs` has been evaluated for \( n < m \), then providing this object to `noaTabs` reduces the number of recursions needed to evaluate for \( m \).
- **m**: The number of contributors
- **weight**: See the paper for details. Usually, this should be set to 1

**Details**

Computes the \( \alpha_{m} \)-vectors and their weights, \( c(\alpha_{m}) \), by recursion over the number of contributors \( m \). That is, \( \alpha_{m} \) is obtained based on \( \alpha_{m-1} \) — see paper for details.

**Value**

Returns a named vector of counts, where the names denotes the \( \alpha_{m} \)-vectors and the countes is the associated weights, \( c(\alpha_{m}) \).

**Author(s)**

Torben Tvedebrink
optim.relatedness

Estimate theta and the fraction of comparisons between close relatives

Description
Estimates the fraction of comparisons between pairs of close relatives while fitting the theta parameter minimising the object function. The function makes use of the R-package 'Rsolnp' which is an implementation of an solver for non-linear minimisation problems with parameter constraints.

Usage
```
optim.relatedness(obs, theta0=0, theta1=0.03, theta.tol=10^(-7), theta.step=NULL, max.bisect=15, probs, var.list=NULL, init.alpha=10^c(-4,-6,-8,-10), init.keep=FALSE, objfunction=c("T2","T1","C3","C2","C1"), collapse=FALSE, trace=FALSE, solnp.ctrl=list(tol=10^(-9), rho=10, delta=init.alpha*0.01, trace=FALSE))
```

Arguments
- `obs`: The matrix or vector of observed matches/partial-matches as returned by the dbCompare()-function
- `theta0`: The left value of the interval in which a bisection-like search is performed for theta
- `theta1`: Right value of interval (see theta0)
- `theta.tol`: A stopping criterion for the search. If the search narrows within theta.tol the function terminates

References
theta.step  Default is NULL. If not a grid search will be performed on seq(from = theta0, to = theta1, by = theta.step)
max.bisect  The maximum number of bisectional iterations perform prior to termination
probs  List of vectors with allele probabilities for each locus
var.list  A named list of components for computing variances, see dbVariance. The names of the elements are the associated theta-values, and each component is a list of (V1,V2,V3) - see dbVariance with n=1
init.alpha  Initial values for alpha, where the order is (First-cousins, Avuncular, Parent-child, Full-siblings). The value for Unrelated is computed as 1-sum(init.alpha)
init.keep  Whether the initial values should be used in successive steps for the current optimum should be used.
objFunction  Which of the five different object functions should be used to compare observed and expected
collapse  Not yet implemented
trace  Should iteration steps and other process indicators be printed
solnp.ctrl  See solnp for details

Details

Computes the proportion of comparisons between close relatives in a database matching exercise for each theta value under investigation.

Value

Returns a list of three components: value, solution and var.list. The first element, value, is a dataframe with the value of the objection function for each of the theta values investigated. Solution is the estimated alpha-vector where the objection function was minimised. Finally, var.list is a names list of components for computing variances. May be reused in later computations for increased speed in some iterations.

Author(s)

James Curran and Torben Tvedebrink

References


Examples

## Not run:
## Simulate some allele frequencies:
freqs <- replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g)}, simplify=FALSE)
## Load the sample database:
pContribution

```r
data(dbExample)
obs <- dbCompare(dbExample, trace=FALSE)$m
C3 <- optim.relatedness(obs, theta=0.0, theta1=0.03, probs=freqs,
  objFunction="C3", max.bisect=30, trace=TRUE)

## End(Not run)
```

---

**pContribution**

*Compute the posterior probabilities for $P(m|n_0)$ for a given prior $P(m)$ and observed vector $n_0$ of locus counts*

**Description**

where $m$ ranges from 1 to $m_{\text{max}}$ and $n_0$ is the observed locus counts.

**Usage**

```r
pContribution(n0, probs=NULL, m.prior=rep(1/m.max, m.max), m.max=8, pnoa=NULL, theta=0)
```

**Arguments**

- `n0` Vector of observed allele counts - same length as the number of loci.
- `probs` List of vectors with allele probabilities for each locus.
- `m.prior` A vector with prior probabilities (summing to 1), where the length of `m.prior` determines the plausible range of $m$.
- `m.max` Derived from the length of `m.prior`, and if `m.prior=NULL` a uniform prior is specified by `m.max`: `m.prior = rep(1/m.max, m.max)`.
- `pnoa` A named list of locus specific probabilities. Output from `pNoA` with `locuswise=TRUE`.
- `theta` The coancestry coefficient.

**Details**

Computes a vector $P(m|n_0)$ evaluated over the plausible range 1,...,$m_{\text{max}}$.

**Value**

Returns a vector $P(m|n_0)$ for $m=1,...,m_{\text{max}}$.

**Author(s)**

Torben Tvedebrink

**References**

Examples

```r
## Simulate some allele frequencies:
freqs <- structure(replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g)},
    simplify=FALSE), .Names=paste("locus",1:10,sep=""))
m <- 2
n0 <- unlist(lapply(freqs, function(x){
    length(unique(sample(length(x), size=2*m, replace=TRUE, prob=x))))})
## Compute P(m|n0) for m=1,...,4 and the sampled n0
pContrib(n0=n0, probs=freqs, m.max=4)
```

Description

Compute a matrix of posterior probabilities $P(m|n_0)$ where $m$ ranges from 1 to $m_{\text{max}}$, and $n_0$ is $0,...,2m_{\text{max}}$. This is done by evaluating $P(m|n_0) = P(n_0|m)P(m)/P(n)$, where $P(n_0|m)$ is evaluated by `pNoA`.

Usage

```r
pContrib.locus(prob=NULL, m.prior=NULL, m.max=8, pnoa.locus=NULL, theta=0)
```

Arguments

- `prob`: Vectors with allele probabilities for the specific locus
- `m.prior`: A vector with prior probabilities (summing to 1), where the length of `m.prior` determines the plausible range of `m`
- `m.max`: Derived from the length of `m.prior`, and if `m.prior=NULL` a uniform prior is specified by `m.max`: `m.prior = rep(1/m.max, m.max)`.
- `pnoa.locus`: A named vector of locus specific probabilities $P(N(m) = n), n = 1,...,2m$.
- `theta`: The coancestry coefficient

Details

Computes a matrix of $P(m|n_0)$ values for a specific locus.

Value

Returns a matrix $[P(m|n_0)]$ for $m=1,...,m_{\text{max}}$ and $n_0=1,...,2m_{\text{max}}$.

Author(s)

Torben Tvedebrink
plot.dbcompare

References


Examples

### Simulate some allele frequencies:
```r
freqs <- structure(replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g), simplify=FALSE), .Names=paste("locus",1:10,sep="."))
```

### Compute P(m|nP) for m=1,...,5 and n=1,...,10 for the first locus:
```r
pContrib.locus(prob=freqs[[1]],m.max=5)
```

---

**Description**

Plots the summary matrix with counts on y-axis and classification on x-axis.

**Usage**

```r
## S3 method for class 'dbcompare'
plot(x,log="y",las=3,xlab="Match/Partial",ylab="Counts",...)
```

**Arguments**

- `x` Summary matrix returned from dbcompare
- `log` Specifies whether log(Counts) should be plotted (default)
- `las` Direction of the labels on x-axis. Default is 3 which gives perpendicular labels
- `xlab` Axis label
- `ylab` Axis label
- `...` Other plot options

**Details**

Plots the summary matrix

**Value**

A plot of the summary matrix. The counts are on log10 scale and the x-axis is labeled by appropriate matching/partially-matching levels.

**Author(s)**

James Curran and Torben Tvedebrink
plot.dbOptim

See Also
dbCompare, print.dbcompare

Examples

## Not run:
data(dbExample)
M = dbCompare(dbExample, hit=5)
plot(M)

## End(Not run)

plot.dbOptim

Plots the fitted object function for estimated familial relationships in
the database and theta.

Description

Plots the minimised object function for included values of theta

Usage

## S3 method for class 'dbOptim'
plot(x, type="l", ...)
## S3 method for class 'dbOptim'
points(x, type="p", ...)
## S3 method for class 'dbOptim'
lines(x, type="l", ...)

Arguments

x Object returned by optim.relatedness
type The type of plot character ("l"=line, "p"=points, ...), see 'par' for more details
... Other plot options

Details

Plots the object function

Value

A plot of the object function

Author(s)

James Curran and Torben Tvedebrink
The exact distribution of the number of alleles in a \( m \)-person DNA mixture

Computes the exact distribution of the number of alleles in a \( m \)-person DNA mixture typed with STR loci. For a \( m \)-person DNA mixture it is possible to observe \( 1, \ldots, 2mL \) alleles, where \( L \) is the total number of typed STR loci. The method allows incorporation of the subpopulation correction, the so-called \( \theta \)-correction, to adjust for shared ancestry. If needed, the locus-specific probabilities can be obtained using the \texttt{locuswise} argument.

Usage

\[
\texttt{pNoA(probs, } m=2, \texttt{ theta } = 0, \texttt{ noa.tabs=\texttt{NULL}, locuswise=\texttt{FALSE})}
\]

Arguments

- \texttt{probs} List of vectors with allele probabilities for each locus
- \texttt{m} The number of contributors
- \texttt{theta} The coancesterny coefficient
- \texttt{noa.tabs} If \texttt{noa.tabs} has been computed by the \texttt{noaTabs} function for \( m \), then this will speed up the computations.
- \texttt{locuswise} Logical. If \texttt{TRUE} the locuswise probabilities will be returned. Otherwise, the probability over all loci is returned.

Details

Computes the exact distribution of the number of alleles for a \( m \)-person DNA mixture.
Value

Returns a vector of probabilities, or a list of locuswise probability vectors.

Author(s)

Torben Tvedebrink

References


Examples

```r
## Simulate some allele frequencies:
freqs <- structure(replicate(10, { g = rgamma(n=10, scale=4, shape=3); g/sum(g)}, simplify=FALSE), .Names=paste("locus",1:10, sep="."))
## Compute alpha_2
noa.tab3 <- noaTabs(m=2)
## Compute P(N(m=3)=n), n=1,...,2*L*m, where L=10 here
pNoA(freqs,m=2,theta=0,noa.tabs=noa.tab3)
## Same, but locuswise results
pNoA(freqs,m=2,theta=0,noa.tabs=noa.tab3,locuswise=TRUE)
```

print.dbcompare  

Prints the summary matrix

Description

Prints the summary matrix and possible "big hits".

Usage

```r
## S3 method for class 'dbcompare'
print(x,...)
```

Arguments

- `x` Summary matrix returned from dbcompare

Details

Prints the summary matrix

Value

Prints the summary matrix and data frame with "big hits"
Author(s)
James Curran and Torben Tvedebrink

See Also
dbCompare, plot.dbcompare

Examples
## Not run:
data(dbExample)
M = dbCompare(dbExample, hit=5)
M
## End(Not run)

print.dbOptim                  Prints the results from optim.relatedness()

Description
Prints the evaluated functions for the object function, best estimate of alpha and possibly list of variances.

Usage
## S3 method for class 'dbOptim'
print(x, var.list=FALSE, ...)

Arguments
x                  Object returned by optim.relatedness()
var.list           Logical. Whether the (long) list of variance components should be printed to the screen.
...                 ...

Details
Prints the summary details of the fit

Value
A dataframe with [theta, value] and a vector of fitted alpha parameters

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See Also

optim.relatedness

Examples

## Not run:
## Simulate some allele frequencies:
freqs <- replicate(10, { g = rgamma(n=10,scale=4,shape=3); g/sum(g)}, simplify=FALSE)
## Load the sample database:
data(dbExample)
obs <- dbCompare(dbExample,trace=FALSE)$m
C3 <- optim.relatedness(obs,theta0=0.0,theta1=0.03,probs=freqs, objFunction="C3",max.bisect=30,trace=TRUE)
print(C3)

## End(Not run)
Index

* Topic **Forensic genetics**
  DNAtools-package, 2

  dbCollapse, 3
  dbCompare, 4
  dbExample, 5
  dbExpect, 6
  dbSimulate, 7
  dbVariance, 9
  DNAtools (DNAtools-package), 2
  DNAtools-package, 2

  estimatePD, 10

  freqEst, 11

  lines.dbOptim(plot.dbOptim), 18

  noaTabs, 12

  optim.relatedness, 13

  p.numberofalleles (pNoA), 19
  pContribution, 15
  pContribution.locus, 16
  plot.dbcompare, 17
  plot.dbOptim, 18
  pNoA, 19
  points.dbOptim (plot.dbOptim), 18
  print.dbcompare, 20
  print.dbOptim, 21