Package 'sonicLength'

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Type Package

Title Estimating Abundance of Clones from DNA Fragmentation Data

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Description Estimate the abundance of cell clones from the distribution of lengths of DNA fragments (as created by sonication, whence `sonicLength'). The algorithm in "Estimating abundances of retroviral insertion sites from DNA fragment length data" by Berry CC, Gillet NA, Melamed A, Gormley N, Bangham CR, Bushman FD. Bioinformatics; 2012 Mar 15;28(6):755-62 is implemented. The experimental setting and estimation details are described in detail there. Briefly, integration of new DNA in a host genome (due to retroviral infection or gene therapy) can be tracked using DNA sequencing, potentially allowing characterization of the abundance of individual cell clones bearing distinct integration sites. The locations of integration sites can be determined by fragmenting the host DNA (via sonication or fragmentase), breaking the newly integrated DNA at a known sequence, amplifying the fragments containing both host and integrated DNA, sequencing those amplicons, then mapping the host sequences to positions on the reference genome. The relative number of fragments containing a given position in the host genome estimates the relative abundance of cells hosting the corresponding integration site, but that number is not available and the count of amplicons per fragment varies widely. However, the expected number of distinct fragment lengths is a function of the abundance of cells hosting an integration site at a given position and a certain nuisance parameter. The algorithm

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implicitly estimates that function to estimate the relative abundance.

License GPL (>= 2)

LazyLoad yes

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Description

Estimate the number of fragments produced by sonication of integration sites in DNA (as arise in gene therapy or in retroviral infections) from tallies of the number of distinct lengths. See the Berry et al. reference where the experimental setting and algorithm are spelled out in detail.

Details

Package: sonicLength Type: Package

License: see file LICENSE distributed with this software

LazyLoad: yes

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This package has the necessary utilities to provide a function that will do calibration of the number of distinct lengths (for an integration site recovered via sonication) to estimate the number of distinct sonication fragments that underlie the observed number of lengths.

Author(s)

Charles C. Berry

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References

The reference to the algorithm is:

Estimating abundances of retroviral insertion sites from DNA fragment length data. Berry CC, Gillet NA, Melamed A, Gormley N, Bangham CR, Bushman FD. Bioinformatics; 2012 Mar 15;28(6):755-62.

Some applications that illustrate use of this method are found here:

Gillet, Nicolas A., et al. "The host genomic environment of the provirus determines the abundance of HTLV-1 infected T-cell clones." Blood 117.11 (2011): 3113-3122.

Maldarelli, F., et al. "Specific HIV integration sites are linked to clonal expansion and persistence of infected cells." Science 345.6193 (2014): 179-183.

Α1

HTLV-1 Fragment Length Data

Description

A sample from a patient with HTLV-1 was analyzed as described in Gillet et al (2001).

Usage

data(A1)

Format

A data frame with 6176 observations on the following 3 variables.

locations a factor with levels 01 F 101480450 ... X R 99972891 indicating the genomic sites at which insertions were detected

lengths a numeric vector of fragment lengths

replicates a numeric vector giving replicate IDs

Details

For each replicate, the unique combinations of the genome location at which an insertion site was detected and the length(s) of the fragment(s) sequenced were noted. There is one such row in he data.frame for each such combination.

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Source

See Gillet et al 2011 for more details.

References

Gillet, N.A., Malani, N., Melamed, A., Gormley, N., Carter, R., Bentley, D., Berry, C., Bushman, F.D., Taylor, G.P., and Bangham, C.R. (2011). The host genomic environment of the provirus determines the abundance of HTLV-1-infected T-cell clones. *Blood* 117, 3113-3122.

Examples

```
data(A1)
str(A1)
```

estAbund

estimate Abundances from sonicated samples

Description

When one or more replicate samples are sonicated and the lengths for each integration site are recorded, it is of interest to estimate actual number of sonicants allowing for coincidentally equal lengths. A likelihood approach is used here.

Usage

```
estAbund(locations, lengths, replicates = NULL, jackknife = F, kmax=0,
min.length = 20, ...)
```

Arguments

| locations | a vector of IDs for distinct locations |
|------------|--|
| lengths | a vector of the corresponding lengths |
| replicates | an optional vector of the replicate sample ID |
| jackknife | Whether to do leave-one-out jackknife samples over the replicates |
| kmax | highest count to bother with when computing mass function for counts. All higher values are globbed together in the result. If kmax==0 don't compute |
| min.length | Least length expected for a fragment. Shorter lengths will not be permitted |
| | Other arguments that may be passed along to functions that do the real work. |

Details

This is a wrapper function for maxEM so study that function to see what is going on.

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Value

a list with components

theta estimated abundances
phi estimated bin probabilities

var.theta variances of theta

iter number of iterations till convergence was achieved

call the function call

1frame the data. frame used to construct the estimate of phi

obs the observed counts of distinct lengths

jackknife a list of calls to maxEM with one replicate omitted in each

data a data. frame with columns locations, lengths, and replicates (if supplied)

Author(s)

Charles C Berry <ccberry@users.r-forge.r-project.org>

References

Estimating abundances of retroviral insertion sites from DNA fragment length data. Berry CC, Gillet NA, Melamed A, Gormley N, Bangham CR, Bushman FD. Bioinformatics; 2012 Mar 15;28(6):755-62.

See Also

maxEM phi.update.lframe The package vignette:Estimating Abundances with sonicLength

Ey.given.x

E-Step for Abundance Estimation

Description

Take the E-step of the EM algorithm for estimating the number of sonicants in a sample

Usage

```
Ey.given.x(x, theta, phi)
pr.y.given.x(x, theta, phi, kmax=20 )
```

Arguments

| x a zero-one indicator matrix whose rows correspond to unique length | ths with row- | |
|--|---------------|--|
|--|---------------|--|

names indicating those lengths

theta a vector of the abundance estimates. length(theta) = ncol(x)

phi a vector of the probabilities of sonicant lengths. length(phi)==nrow(x)

kmax highest count to bother with (all higher values are globbed together in the result)

Details

Supposing Poisson sampling of sonicants, Ey.given.x(x,theta,phi)[i,j] gives the expected value of the number of sonicants given that x[i,j] distinct sonicant lengths were observed. pr.y.given.x(x,theta,phi.km gives the probability of k sonicants (censored at kmax+1) of insertion site j

Value

```
Ey.given.x() is a double matrix of the same dimension as x. pr.y.given.x(...,kmax) is a double matrix of dimension c(kmax+1,ncol(x))
```

Author(s)

Charles C. Berry <ccberry@users.r-forge.r-project.org>

Examples

```
mat <- diag(10)
mat[ ,10 ] <- 1.0
phi1 <- prop.table( rep(1,10))
theta1 <- 1:10
Ey.given.x( mat, theta1, phi1 )
pr.mat <- pr.y.given.x( mat, theta1, phi1 )
## Estimate Seen plus Unseen sites
popsize.chao <- function(tab) sum(tab)+tab[1]*(tab[1]-1)/(2*(tab[2]+1))
## evaluate at expected counts
popsize.chao( rowSums(pr.mat) )
## average randomly sampled counts
cnt.samp <- function(x) sample( seq_along(x) , 1 ,pr=x )
mean(replicate(100,popsize.chao( table(apply(pr.mat,2, cnt.samp) ))))</pre>
```

Fragment Distribution Simple Fragment Length Distribution

Description

Density, distribution function, quantile function and random generation for a simple parametric distribution for fragments lengths (given by conditioning a geometric distribution on whether the length can be recovered).

Usage

```
dfrag( x, loc=45, lscale=2.5, rate=0.02, maxx=qgeom(1-1e-7,rate) )
pfrag( q, loc=45, lscale=2.5, rate=0.02, maxx=qgeom(1-1e-7,rate), lower.tail=TRUE )
qfrag( p, loc=45, lscale=2.5, rate=0.02, maxx=qgeom(1-1e-7,rate), lower.tail=TRUE )
rfrag( n, loc=45, lscale=2.5, rate=0.02, maxx=qgeom(1-1e-7,rate) )
```

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Arguments

| x,q | vector of quantile (of lengths). |
|------------|--|
| n | number of lengths to sample. |
| p | vector of probabilities |
| loc | vector of locations of logistic for prob of recovery. |
| lscale | vector of scales of logistic for prob of recovery. |
| rate | probability for geometric probability for fragment lengths. |
| maxx | integer; largest value of x to bother with. |
| lower.tail | logical; if TRUE (default), probabilities are $P[X \le x]$ otherwise, $P[X > x]$. |

Details

The mass function is given by plogis(x,loc,lscale)*dgeom(x,rate) / denom, where denom scales the result to sum to 1.0 in the range 0:maxx. The other functions all depend on this in the obvious manner. If maxx is not large enough a warning may be issued, but even without this warning the results may be slightly innaccurate if pgeom(maxx,rate,lower.tail=FALSE) is non-negligible.

Value

dfrag gives the mass function, pfrag gives the distribution function, qfrag gives the quantile function, and rfrag generates random deviates.

Author(s)

```
Charles C. Berry <ccberry@users.r-forge.r-project.org>
```

See Also

```
pgeom, sample
```

Examples

```
plot( 0:300, table(factor(rfrag(2000),0:300)) )
lines( 0:300, 2000*dfrag(0:300) )
```

8 maxEM

| maxEM maximum likelihood estimates relative abundances | |
|--|--|
|--|--|

Description

From information about the lengths of sonicants of integration sites, infer the relative abundances of different clones and the distributon of sonicant lengths

Usage

Arguments

| slmat | a matrix whose rows correspond to unique lengths with rownames indicating those lengths |
|--------------|--|
| theta.var | logical, return variance of theta estimates? |
| phi.update | a function of an object like slmat that returns estimtes of phi - a default vesrion is invoked if none is provided |
| phi.deriv | function of theta and phi that returns derivatives of phi wrt beta (its parameters) |
| lframe | a data. frame which will be used to estimate phi, the supplied function phi .update must know what to do with this arg or it will be ignored |
| glm.frm | a formula like $y \sim bs(x,knots=c(50,100))$ to fit phi |
| iter.control | a list of default values for iteration control - see maxEM.iter.control |
| | possibly other args to pass to phi.update |

Details

The EM method is used to infer the relative abundances of different sites.

Value

```
theta a vector of the abundance estimates

phi a vector of the probabilities of sonicant lengths

call the call used
```

Author(s)

```
Charles C. Berry <ccberry@users.r-forge.r-project.org>
```

See Also

```
Ey.given.x
```

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Examples

maxEM.iter.control

iteration controls

Description

Set parameters for control of iteration in maxEM

Usage

```
maxEM.iter.control(min.reps = 3, max.reps = 2000, max.abs.le = 0.1,
max.rel.le = 1e-05, phi.min=.Machine$double.eps)
```

Arguments

```
min.reps a positive integer
max.reps a positive integer
max.abs.le a smallish double
max.rel.le a tiny double
phi.min a smallish double
```

Details

Most users should not need to tinker with these settings

Value

```
a list such as formals(maxEM.iter.control)
```

Author(s)

```
Charles C. Berry <ccberry@users.r-forge.r-project.org>
```

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mstep

M-step to maximize theta

Description

This is a utility function for maxEM

Usage

```
mstep(x, theta, phi)
```

Arguments

x a zero-one indicator matrix whose rows correspond to unique lengths with row-

names indicating those lengths

theta a vector of the abundance estimates. length(theta)==ncol(x)

phi a vector of the probabilities of sonicant lengths. length(phi)==nrow(x)

Details

The M-step for theta is computed. Probably, there is no need to use this function directly, but just in case it is here.

Value

a vector like the input theta

Author(s)

Charles C. Berry

See Also

maxEM

Examples

```
mat <- diag(10)
mat[ ,10 ] <- 1.0
phi1 <- prop.table( rep(1,10))
theta1 <- 1:10
sonicLength:::mstep( mat, theta1,phi1)</pre>
```

phi.update.lframe

Description

Estimate phi in a flexible manner as guided by 1frame - usually involving multiple strata, and allowing for formulas other than the default choice

Usage

```
phi.update.lframe(obj, return.fit = FALSE, lframe, glm.frm = NULL, ...)
```

Arguments

| obj | a matrix whose rowSums are used to estimate phi |
|------------|---|
| return.fit | logical - the result of the fit should be returned instead of phi |
| lframe | A data.frame with columns y, x, strata, orig (or matching those in glm.frm) and for which rownames(lframe)[lframe σ] match the rownames of obj. |
| glm.frm | a formula. If omitted $y \sim bs(x,knots=c(50,100))$ will be used. |
| | curently not used |

Details

fitting phi - the probabilities that a sonicant lands in a particular bin - is crucial to estimating theta, then number of sonicants when more than one lans in a bin. bins may be defined by sonicant lengths when there is just one sample. When there are multiple samples, then one may wish to estimate phi separately for each one. lframe\$strata indicates separate sample and the default glm.frm will fit each one separately.

Value

a vector whose sum is 1.0 with one element for each row of obj

Author(s)

```
Charles C. Berry <ccberry@users.r-forge.r-project.org>
```

See Also

maxEM

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simSonic

Simulate Sonicated Data

Description

Sonication and Processing of DNA containing retroviral sequences yields genomic locations of retroviral insertion sites and a set of associated fragment lengths. The simulation depends on the expected number of fragments from each site and the distribution of lengths.

Usage

```
simSonic( theta, phi )
```

Arguments

theta a vector of expected number of fragments from each site. If the vector has

unquee names, these will be used to label the locations

phi a vector of probabilities (or a list containing a vector named phi) whose names

are like 'replen', where 'rep' defines the replicate and 'len' defines the sonicant length. More than one replicate can be used, In any case 1-exp(-phi%o%theta) determines the probability of observations in the different locaitons, of the dif-

ferent lengths, and in different replicates

Details

This object can provide the arguments used by estAbund

Value

a data.frame with columns locations, lengths, and replicates. See estAbund for more details

Author(s)

```
Charles C. Berry <ccberry@users.r-forge.r-project.org>
```

Examples

```
theta <- seq(0.5,20.5,by=0.5)
phi <- prop.table(1:10)
names(phi) <- paste( 1 , 51:60 )
res <- simSonic( theta, phi )
head(res)
tail(res)
summary(res)</pre>
```

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