

Package ‘RTIGER’

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Type Package

Title HMM-Based Model for Genotyping and Cross-Over Identification

Version 1.99.0

Description Our method integrates information from all sequenced samples, thus avoiding loss of alleles due to low coverage. Moreover, it increases the statistical power to uncover sequencing or alignment errors.

Depends R (>= 3.6), GenomicRanges, GenomeInfoDb

License GPL (>= 2)

Encoding UTF-8

LazyData true

LazyDataCompression gzip

Imports methods, e1071, reshape2, ggplot2, TailRank, JuliaCall, IRanges, qpdf, grDevices, graphics, stats, utils

RoxygenNote 7.1.2

VignetteBuilder knitr

Suggests knitr, rmarkdown, markdown, Gviz, rtracklayer

biocViews GenomeAnnotation, HiddenMarkovModel, Sequencing

NeedsCompilation no

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ATseqlengths	<i>The autosome chromosome lengths for Arabidopsis Thaliana.</i>
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Description

The autosome chromosome lengths for Arabidopsis Thaliana.

Author(s)

Rafael Campos-Martin

calcCNumber	<i>Obtain number of Cross-Over events per sample and chromosome.</i>
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Description

Obtain number of Cross-Over events per sample and chromosome.

Usage

```
calcCNumber(object)
```

Arguments

object a RViterbi object.

Value

Matrix $m \times n$. M number of samples and N chromosomes.

#' @return a matrix with n chromosomes and m samples (n x m) and the number of CO events.

Examples

```
data("fittedExample")
co.num = calcC0number(myDat)
```

dev	<i>Function to developers. It runs one EM step</i>
-----	--

Description

Function to developers. It runs one EM step

Usage

```
dev(psi, rigidity = NULL, nstates = 3, transition = NULL, start = NULL)
```

Arguments

psi	list of psi probabilities.
rigidity	Rigidity value.
nstates	Number of states.
transition	transition matrix
start	initial probabilities

Value

List with updates probabilities

fit	<i>Call Julia code to fit the values</i>
-----	--

Description

Call Julia code to fit the values

Usage

```
fit(rtigerobj, max.iter , eps,
trace, all = TRUE, random = FALSE,
specific = FALSE, nsamples = 20,
post.processing = TRUE)
```

Arguments

<code>rtigerobj</code>	an RTIGER object.
<code>max.iter</code>	maximum number of iterations to accomplish by the EM.
<code>eps</code>	differnece threshold to halt the EM.
<code>trace</code>	logical value whether to trace the changes in the parameters along the iterations.
<code>all</code>	logical value whether to use all data to fit the model.
<code>random</code>	if all FALSE use random samples.
<code>specific</code>	if all FALSE use specific samples.
<code>nsamples</code>	if random TRUE, how many samples to use.
<code>post.processing</code>	logical value, whether to run post.processing process.

Value

RTIGER object

Examples

```
## Not run:
data("fittedExample")
sourceJulia()
myfit = fit(myDat, max.iter = 2, eps=0.01,
            trace = TRUE, all = TRUE,
            random = FALSE, specific = FALSE,
            nsamples = 20, post.processing = TRUE)

## End(Not run)
```

`generateObject`

Load data

Description

Load data

Usage

```
generateObject(experimentDesign = NULL, nstates = 3, rigidity=NULL,
               seqlengths = NULL, verbose = TRUE)
```

Arguments

experimentDesign	a data Frame that contains minimum a column with the files direction (name of the column files) and another with a shorter name to be used inside the function.
nstates	the number of states to be fitted in the model. A standard setting would use 3 states (Homozygous1, Heterozygous, and Homozygous2).
rigidity	an integer number specifying the rigidity parameter to be used.
seqlengths	a named vector with the chromosome lengths of the organism that the user is working with.
verbose	logical value. Whether to print info messages.

Value

RTIGER object

Examples

```
data("ATseqlengths")
path = system.file("extdata", package = "RTIGER")
files = list.files(path, full.names = TRUE)
nam = sapply(list.files(path ), function(x) unlist(strsplit(x, split = "[.]"))[1])
expDesign = data.frame(files = files, name = nam)
names(ATseqlengths) = paste0("Chr", 1:5)
myres = generateObject(experimentDesign = expDesign,
                      seqlengths = ATseqlengths,
                      rigidity = 10
                      )
```

myDat

A fitted example using three own samples of Arabidopsis. More information in publication:

Description

A fitted example using three own samples of Arabidopsis. More information in publication:

Author(s)

Rafael Campos-Martin

plotCOs	<i>Obtain number of Cross-Over events per sample and chromosome.</i>
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Description

Obtain number of Cross-Over events per sample and chromosome.

Usage

```
plotCOs(object, file = NULL)
```

Arguments

object	a RViterbi object.
file	file where to save the plot for CO numbers

Value

a plot

Examples

```
data("fittedExample")  
co.num = calcCOnumber(myDat)
```

RTIGER	<i>Load, Fit, and plot</i>
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Description

Load, Fit, and plot

Usage

```
RTIGER(expDesign, rigidity=NULL, outputdir=NULL, nstates = 3,  
seqlengths = NULL, eps=0.01, max.iter=50, trace = FALSE,  
tiles = 4e5, all = TRUE, random = FALSE, specific = FALSE,  
nsamples = 20, post.processing = TRUE, save.results = TRUE, verbose = TRUE)
```

Arguments

<code>expDesign</code>	a data Frame that contains minimum a column with the files direction (name of the column files) and another with a shorter name to be used inside the function.
<code>rigidity</code>	an integer number specifying the rigidity parameter to be used.
<code>outputdir</code>	a character string that specifies the directory in which to save the results form the function.
<code>nstates</code>	the number of states to be fitted in the model. A standard setting would use 3 states (Homozygous1, Heterozygous, and Homozygous2).
<code>seqlengths</code>	a named vector with the chromosome lengths of the organism that the user is working with.
<code>eps</code>	the threshold of the difference between the parameters value between the previous and actual iteration to stop the EM algorithm.
<code>max.iter</code>	maximum number of iterations of the EM algorithm before to stop in case that <code>eps</code> has not been achieved.
<code>trace</code>	logical value. Whether or not to keep track of the parameters for the HMM along the iterations. Default FALSE
<code>tiles</code>	length of the tiles by which the genome will be segmented in order to compute the ratio of COs in the complete dataset.
<code>all</code>	logical value. Whether to use the complete data set to fit the rHMM. default TRUE.
<code>random</code>	Logical value. Choose randomly a subset of the complete dataset to fit the rHMM. Default FALSE
<code>specific</code>	Logical value to specify which samples to take.
<code>nsamples</code>	if <code>random</code> TRUE, how many samples should be taken randomly.
<code>post.processing</code>	Logical value. Whether to run an extra step that fine maps the segment borders. Default TRUE
<code>save.results</code>	Logical value, whether to generate and save the plots and igv files.
<code>verbose</code>	Logical, whether to print info to console.

Value

Matrix $m \times n$. M number of samples and N chromosomes.

RTIGER object

Examples

```
## Not run:
data("ATseqlengths")
sourceJulia()
path = system.file("extdata", package = "RTIGER")
files = list.files(path, full.names = TRUE)
nam = sapply(list.files(path), function(x) unlist(strsplit(x, split = "[.]"))[1])
expDesign = data.frame(files = files, name = nam)
```

```

names(ATseqlengths) = paste0("Chr", 1:5)
myres = RTIGER(expDesign = expDesign,
               outputdir = "/home/campos/Documents/outputjulia/",
               seqlengths = ATseqlengths,
               rigidity = 4,
               max.iter = 2,
               trace = FALSE,
               save.results = TRUE)

## End(Not run)

```

RTIGER-class	<i>This class is a generic container for RTIGER analysis</i>
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Description

This class is a generic container for RTIGER analysis

Slots

matobs Nested lists. the first level is a list of samples. For each sample there are 5 matrices that contains the allele counts for each position.

params a list with the parameters after training.

info List with phenotypic data of the samples.

Viterbi List of chromosomes with the viterbi path per sample.

Probabilities Computed probabilities for the EM algorithm.

num.iter Number of iterations needed to stop the EM algorithm.

setupJulia	<i>Installs the needed packages in JULIA to run the EM algorithm for rHMM.</i>
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Description

Installs the needed packages in JULIA to run the EM algorithm for rHMM.

Usage

```
setupJulia(JULIA_HOME = NULL)
```

Arguments

JULIA_HOME	the file folder which contains julia binary, if not set, JuliaCall will look at the global option JULIA_HOME, if the global option is not set, JuliaCall will then look at the environmental variable JULIA_HOME, if still not found, JuliaCall will try to use the julia in path.
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Value

empty

sourceJulia	<i>Function needed before using RTIGER() function. It loads the scripts in Julia that fit the rHMM.</i>
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Description

Function needed before using RTIGER() function. It loads the scripts in Julia that fit the rHMM.

Usage

sourceJulia()

Value

empty

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