Package 'inveRsion'

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Description

This package scans the whole genome in search of inversion events. Input data can be genotypes or phased haplotypes. It computes regions where inversions are probed by trial segments of fixed length. An inversion model is fit at each trial segment and significance measures, like Bayes Information Criterion, give evidence of segments belonging to the inversion event. Methods are implemented to identify the complete inversion segment and to classify the chromosomes in the sample as inverted or not.

Details

Package: inveRsion Type: Package Version: 1.0

Date: 2010-11-12

License: GPL version 2 or newer

LazyLoad: yes

Depends: methods, haplo.stats

The package is designed as a stream analysis of a sequence of procedures: setUpGenoDatFile loads the genotype data onto R; HaploCode performs local haplotyping around the candidate brake points of the inversion; sanInv takes trial segments of fixed window size and fits the inversion model, sweeping the whole genome; and listInv summarizes the inversion events.

Author(s)

Alejandro Caceres Maintainer: Alejandro Caceres <acaceres@creal.cat>

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References

A Caceres el al. Detection of genetic inversions with SNP-array data, manuscript in preparation.

SS Sindi and BJ Raphael, Identification and frequency estimation of inversion polymorphisms from haplotype data, J Comput Biol. 2010 Mar;17(3):517-31.

PF O'Reilly et al., invertFREGENE: software for simulating inversions in population genetic data, Bioinformatics. 2010 Mar 15;26(6):838-40. Epub 2010 Jan 26.

Examples

```
#vignette("inveRsionMan")
```

ac

Sample data set of class accuracy

Description

Illustrative output of function accBic. It stores the accuracy of classification of each chromosome into the inverted population and the frequency of the inversion as a for a range of Bic thresholds.

Usage

```
data(hapCode)
```

Format

The format is: Formal class '"accuracy"' [package "inveRsion"] with 1 slots ..@ out: num bicInt prob ac [1,] 0.0000 0.3180 0.9045 [2,] 142.4209 0.3745 0.9610 [3,] 284.8419 0.3945 0.9810 [4,] 427.2628 0.4130 0.9995 [5,] 569.6838 0.4130 0.9995

Examples

```
data(ac)
ac
```

accBic

accBic computes "accuracy" from "inversionList"

Description

accBic computes the accuracy of the classification of chromosomes into previously known inverted and non-inverted populations. The classification is obtained from a majority vote of the classifications produced by the trial segment models whose BIC is greater than a given threshold.

Usage

```
accBic(object, mem, classFile, nsub, npoints, geno, wROI)
```

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Arguments

object of class inversionList

mem vector with the numbering of chromosomes known to have the inversion

classFile an alternative to mem, it passes the file name containing the numbering of chro-

mosomes known to have the inversion.

nsub total number of subjects (= 2* total number of chromosomes)

npoints number of BIC threshold between 0 and max (BIC) for which the accuracy is to

be computed

geno whether the accuracy is assessed for inversion genotype or inversion allele (phased

data).

wROI integer indicating the ROI number to be used. The total number of ROIs are the

total number of components in the object list.

Value

accuracy object of class accuracy

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

```
inversionList, accuracy
```

Examples

```
data(invList)
memFile <- system.file("extdata", "mem.txt", package = "inveRsion")
ac <- accBic(invList, classFile = memFile, nsub = 1000, npoints = 10)
plot(ac)</pre>
```

accuracy-class

Class "accuracy"

Description

These objects hold the accuracy computation for the classification of chromosomes into inverted or non-inverted population for increasing levels of BIC thresholds.

Objects from the Class

accuracy is generated by calls to accBic(inversionList, mem, classFile, nsub, npoints, geno, wROI), which is a method for the class inversionList

codeHaplo 5

Slots

out: Object of class "matrix" matrix with three columns storing the range of BIC thresholds, probability of inversion within the population and accuracy of the classification.

Methods

```
plot signature(x = "accuracy", w="character"): plots accuracy Vs BIC for w="a", or fre-
quency of inversion Vs BIC for w="f"
```

Note

version R 2.10.1

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

accBic, inversionList

Examples

data(ac)
ac
plot(ac)

codeHaplo

Codes haplotypes into decimal integers

Description

The function labels the haplotypes of size blockSize around each candidate brake-point. For labeling genotype data, the function takes objects of class genoDat as main argument. For phased data, this argument should be ignored and a file name passed instead.

Usage

codeHaplo(objectGenoDat, blockSize, minAllele, saveRes = TRUE, file = NULL, ROI, intSNP=FALSE, phasing="

Arguments

objectGenoDat Genodat object; if phased data then provide file instead

blockSize numeric. number of SNPs flanking each side of each candidate brake-point.

Default value 3

minAllele numeric. minimum allele frequency for each probe to be considered as a candi-

date brake-point. Default value 0.1

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saveRes logical. Whether results should be saved in file hapCode.RData

file character. File name with phased data

ROI numeric. 2-vector specification passes a chromosome segment to be encoded.

4-vector specification passes the region of interest for the left brake-point (ROI[1]

and ROI[2]) and the right brake-point (ROI[3] and ROI[4])

intSNP logical. whether build flanking blocks of uniform SNP density across the

genome.

phasing character. Either c("ivenrsion&BP", "forward", "forward&inverted"), defines

the phasing strategy between all four flanking blocks of each break poin. ivenr-sion&BP phases the internal flanking block first, then the external blocks to the internal bloks and match them. "forward" uses the blocks in the sequence assumming the forward population; in the inversion model, the backward population is obtained inverting the internal blocks. "forward&inverted" phases the

forward and inverted population independently.

Details

When setUpGenodat is passed, the coding first computes the local haplotypes for each candidate-brake point form the genotype data and then encodes each haplotype into a decimal integer. The local haplotypes are computed with haplo.em form haplo.stats and assigns those with highest posterior probability to each chromosome. In the case of phased data, passed through file, no local haplotyping is needed and only the labeling is performed.

Value

Object of class HaploCode

Author(s)

Alejandro Caceres <acaceres@creal.cat>

References

http://mayoresearch.mayo.edu/mayo/research/schaid lab/software.cfm

See Also

GenoDat, HaploCode,

```
data(gDat)
hapCode <-codeHaplo(gDat,blockSize=3,minAllele=0.3,saveRes=FALSE)
hapCode</pre>
```

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gDat

Sample data of class genoDat

Description

Data set used to illustrate local haplotype coding performed with codeHaplo

Usage

```
data(gDat)
```

Format

```
The format is: Formal class 'GenoDat' [package "inveRsion"] with 4 slots ..@ genoDat : int [1:9, 1:10] 0 1 0 1 0 0 1 0 0 1 ... ... - attr(*, "dimnames")=List of 2 ... ... $: chr [1:9] "V1" "V2" "V3" "V4" ... ... ... $: NULL ..@ lociPos : Named int [1:10] 959 1268 1393 1467 1531 1761 1847 1987 2006 2030 ... - attr(*, "names")= chr [1:10] "V1" "V2" "V3" "V4" ... ..@ alleleSum : num [1:10, 1] 3 1 8 3 9 1 1 9 1 1 ..@ noMissCount: num [1:10, 1] 9 9 9 9 9 9 9 9 9
```

Examples

```
data(gDat)
gDat
plot(gDat)
```

GenoDat-class

Class "GenoDat"

Description

Object that handles genotype data

Objects from the Class

Objects can be created by calls of the form setUpGenoDatFile or setUpGenoDatSNPmat.

Slots

```
genoDat: "matrix" . Genotypes
lociPos: "numeric". Genomic coordinates
alleleSum: "matrix". Total number of variant alleles in the population per SNP
noMissCount: "matrix". Total number of subjects with no-missing values
```

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Methods

```
plot signature(x = "GenoDat"): Plots minor allele frequency
show signature(object = "GenoDat"): shows data summary
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

```
setUpGenoDatFile, setUpGenoDatSNPmat
```

Examples

```
data(gDat)
gDat
```

GenoDatROI-class

Class "GenoDatROI"

Description

GenoDat defined within an region of interest

Usage

```
ROIGenoDat(objectGenoDat, ROI)
```

Arguments

objectGenoDat GenoDat

ROI numeric. Region of interest. 2-component vector that defines the limits of a

chromosome segments where an inversions is thought to occur.

Details

ROIGenoDat is the constructor of the class.

Objects from the Class

object are created with calls to ROIGenoDat(objectGenoDat,ROI)

Slots

```
genoDat: "matrix" . Genotypes
lociPos: "numeric". Genomic coordinates
alleleSum: "matrix" . Total number of variant alleles in the population per SNP
noMissCount: "matrix" . Total number of subjects with no-missing values
ROI: "numeric" . Region of interest.
```

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Extends

```
Class "GenoDat", directly.
```

Methods

```
initialize signature(.Object = "GenoDatROI")
show signature(object = "GenoDatROI")
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

GenoDat

Examples

```
data(gDat)
gDatROI<-ROIGenoDat(gDat,ROI=c(1268,1847))
gDatROI</pre>
```

 ${\tt getClassif-methods}$

Overall classification

Description

Classifies into inverted or non-inverted populations each chromosome in the sample.

Usage

```
getClassif(object, thBic, wROI, bin,geno,id,nmod)
```

Arguments

object	inversionList. List of inversions obtained from a chromosome scan.
thBic	numeric. BIC threshold above which significant segments are chosen for the final classification
wROI	numeric. ROI number from the list to select classification
bin	logic. Whether binary or continous classification is retrieved
geno	logic. Whether inversion genotype is retrieved
id	character. Vector of subject IDs
nmod	numeric. number of trial segments to be used in the classification counting from the trial segments with highest BIC downwards.

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Details

The overall classification of chromosomes into inverted and non-inverted populations is given by the weighted average of the classifications obtained for each trial segment in the ROI, with BIC greater than thBic.

Value

numeric. Vector with values between 0 and 1 representing membership to the non-inverted and inverted population respectively.

Methods

signature(object = "inversionList") for each of the inversions of the list, it returns the classification of each chromosome.

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

inversionList

Examples

```
data(invList)
r<-getClassif(invList)
head(r)</pre>
```

getInv-methods

gets "scan" into a matrix

Description

Each row of the matrix represents a trial segment of fixed window size, for which the inversion model has been fit. It lists the left and right brake-points and output of the fitting: Log-likelihood ratio, probability of inversion, entropy, BIC (Bayes Information Criterion) and number of haplotypes.

Usage

```
getInv(object, thBic, rnd, Like)
```

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Arguments

1		1.0 ' '		C C 1 ' 1
obiect	scan. Cromosome scanne	ed for inversions	with trial segment	s of fixed window

size.

thBic numeric. BIC threshold above which data is retrieved.

rnd logic. Whether round matrix elements.

Like numeric. Log-likelihood ratio threshold above which data is retrieved.

Details

Matrix with output of scanInv. Each row corresponds to a trail segments with given brake points and significance measures for the inversion model.

Value

matrix.

Methods

signature(object = "scan") returns matrix with output of inversion model for each trial segment

Examples

```
data(scanRes)
a<-getInv(scanRes,thBic=2500)
a</pre>
```

getROIs-methods

Extracts regions of possible inversion events from "scan"

Description

lists the regions of interest in a matrix, given by the overlapping of significant trial segments (of fixed window seize) that may be part of an inversion event.

Usage

```
getROIs(object, thBic)
```

Arguments

object scan. Cromosome scanned for inversions with trial segments of fixed window

size.

thBic BIC threshold above which overlapping segments are considered for the defini-

tion of each ROI

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Details

ROIs are defined as overlapping trial segments with BIC greater than thBIC. The output is a matrix for which each row is one ROI. The first two columns give intervals defining the left brake-points and the two subsequent columns are the intervals for the right brake-points. ROIs are given in mega-basis.

Value

matrix

Methods

```
signature(object = "scan") list of regions of interest
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

scan

Examples

```
data(scanRes)
ROI <- getROIs(scanRes, thBic = 0)
ROI</pre>
```

hapCode

Sample data set of class HaploCode

Description

Illustrative data set, with local haplotypes encoded, to be used as input of scanInv.

Usage

```
data(hapCode)
```

Format

```
The format is: Formal class 'HaploCode' [package "inveRsion"] with 3 slots ..@ haploCode: num [1:2000, 1:583] 8 8 37 37 8 37 8 37 37 ... ... ..- attr(*, "dimnames")=List of 2 .. ... .$: NULL ... ...$: chr [1:583] "0.602976-0.604061" "0.604061-0.605972" "0.60602-0.608417" "0.608668-0.608855" ... ..@ blockSize: num 3 ..@ minAllele: num 0.3
```

```
data(hapCode)
hapCode
```

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HaploCode-class

Class "HaploCode"

Description

The object stores the haplotype coding for each of the candidate brake-points. It is typically generated by calling the function codeHaplo on a GenoDat object.

Objects from the Class

```
Objects can be created by calls of codeHaplo(objectGenoDat,blockSize,minAllele,saveRes = TRUE,file = NULL,ROI)
```

Slots

haploCode: Object of class "matrix" Haplotype coding into decimal integers

blockSize: Object of class "numeric" block size of SNP used to identify the haplotype of each candidate brake-point.

minAllele: Object of class "numeric" minimum allele above which candidate brake-points are considered

Methods

```
initialize signature(.Object = "HaploCode"): Internal, users should use codeHaplo
show signature(object = "HaploCode"): shows HaploCode
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

```
codeHaplo, GenoDat
```

```
data(gDat)
hapCode <-codeHaplo(gDat,blockSize=3,minAllele=0.3,saveRes=FALSE)
hapCode</pre>
```

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inversion-class

Class "inversion"

Description

Internal class (not to be called by the user) that retrieves the output on the inversion model run in an ROI. inversionList is a list of objects of class inversion

Objects from the Class

Objects are created by calls to listInv(object, hapCode, geno, ROI, saveRes, thBic, all)

Slots

classification: Object of class "vector" overall classification (majority vote) of each chromosome for all the trail segments in the ROI

leftBP: Object of class "vector" left brake-points for each of the trial segments

rightBP: Object of class "vector" right brake-points for each of the trial segments

bic: Object of class "vector" BIC for the models on each trail segment

intLeftBP: Object of class "vector" interval for the left brake-point, in the ROI

intRightBP: Object of class "vector" interval for the right-brake point in the ROI

invFreq: Object of class "numeric" overall inversion frequency

RR: Object of class "list" classification of given by each of the trail segments.

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

listInv, inversionList

inversionList-class

Class "inversionList"

Description

Lists output of the inversion model for each region of interest specified.

Objects from the Class

Objects can be created by calls of the form listInv(object, hapCode, geno, ROI, saveRes, thBic, all) and are lists of inversion objects.

invList 15

Slots

```
results: Object of class "list" list of inversion objects
```

Methods

```
accBic signature(object = "inversionList"): Computes accuracy for chromosome classifi-
cation when known
getClassif signature(object = "inversionList"): Extracts classification for each chromosome
plot signature(x = "inversionList"): Plots BIC values for trial segments
show signature(object = "inversionList"): shows object
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

```
listInv, HaploCode
```

Examples

```
data(invList)
invList
```

invList

Sample data of class inversionList

Description

list of objects of class inversion, each of which contains the information of overlapping segments that cover candidate inversions within a chromosome.

Usage

```
data(invList)
```

Format

The format is: Formal class 'inversionList' [package "inveRsion"] with 1 slots ..@ results:List of 1\$:Formal class 'inversion' [package "inveRsion"] with 8 slots

Details

The object is constructed with the function listInv

```
data(invList)
invList
```

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|--|

Description

Determines the full inversion sequence from overlapping trial segments of fixed window sized. It computes the limits of the inversion, the population frequency and retrieves the maximum BIC across the trial segments.

Usage

```
listInv(object, hapCode, geno, ROI, saveRes, thBic, all, saveROI)
```

Arguments

object	scan. Results of scanning the genome for an inversion with a fixed window size; output of scanInv.
hapCode	HaploCode. Object with the result of coding haplotypes for each candidate brake-point; output of codeHaplo
geno	logical. Whether original data is genotypes of phased haplotypes.
ROI	numeric. 2-vector specification passes the chromosome segment to be analyzed. 4-vector specification passes the region of interest for the left brake-point (ROI[1] and ROI[2]) and the right brake-point (ROI[3] and ROI[4]). ROI should be specified in mega-basis units.
saveRes	logical.Whether results should be saved into file list Inv.RData
thBic	numeric. BIC threshold above which trial segments are selected.
all	logical. Whether recomputing within the ROI should be done for all possible segment sizes or the window size in scan.
saveR0I	logical. Whether saving the blocks for the candidate break-points for all the ROIs.

Details

listInv is both a method for class scan and constructor of inversionList. It re-runs the inversion model within the ROIs found in the previous scan. However, it is also possible to explicitly pass the ROIs defined by the user. The re-run is done with the same window size of the scan, which is convenient if enough significant trial segments where found within the inversion segment. If a more detailed re-run is needed set all=TRUE. This computes the model for all possible trial segments of any length within the ROI. Note that for high SNP density this can be computational intensive.

Value

inversionList

scan-class 17

Methods

```
signature(object = "scan")
```

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

 ${\tt HaploCode, scan, inversionList}$

Examples

```
data(scanRes)
data(hapCode)
invList<-listInv(scanRes,hapCode=hapCode,geno=FALSE,saveRes=FALSE,all=FALSE,thBic=0, saveROI=FALSE)
invList</pre>
```

scan-class

Class "scan"

Description

Results from scanning the genome with the inversion model for trial segments of fixed window size.

Objects from the Class

scan objects are typically generated by callings to the constructor function scanInv

Slots

```
leftBP: Object of class "matrix" Left brake-point coordinates (right SNP)
rightBP: Object of class "matrix" Right brake-point coordinates (right SNP)
leftBP2: Object of class "matrix" Left brake-point coordinates (left SNP)
rightBP2: Object of class "matrix" Right brake-point coordinates (right SNP)
LogLike: Object of class "matrix" Log-likelihood ratio for each trial segment
prob: Object of class "matrix" probability of no-inversion for each trial segment
ent: Object of class "matrix" entropy for each trial segment
entTh: Object of class "matrix" entropy threshold for each trial segment
bic: Object of class "matrix" BIC for each trial segment
window: Object of class "numeric" window size
```

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Methods

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

```
listInv, HaploCode, getInv, getROIs
```

Examples

```
data(scanRes)
scanRes
plot(scanRes, which="bic", thBic=0)
```

scanInv

Inversion scan

Description

This function scans a whole chromosome in search for inversion events. The scan is done by fitting an inversion model to all segments in the chromosome with fixed length size.

Usage

```
scanInv(objectHaploCode, window, maxSteps = 30, geno = FALSE, saveRes = TRUE, saveBlocks=TRUE)
```

Arguments

objectHaploCode

Object of class HaploCode produced by the codeHaplo function.

window numeric, size of the window in mega-basis.

maxSteps numeric, maximum number of iteration in the EM algorithm for the inversion

model

geno logical. Whether the original data is genotypes or phases haplotypes.

saveRes logical. Whether results should be saves into file invRes.RData
saveBlocks logical. Whether save blocks for each candidate break point.

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Details

The function processes the haplotypes coded in objectHaploCode. If subsequent re-runs are requires for different window sizes, this object can be omitted. The function will thus search the local directory for previous results to speed up further scans.

Value

object of class scan

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

HaploCode, scan

Examples

```
data(hapCode)
window<-0.5
scanRes<-scanInv(hapCode,window=window,saveRes=FALSE,geno=FALSE,saveBlocks=FALSE)
scanRes
plot(scanRes)</pre>
```

scanRes

Sample data set of class scan

Description

Sample set that illustrates the result of scanning a chromosome with segments of fix window size. It fits the inversion model for each segment and stores its results.

Usage

```
data(scanRes)
```

Format

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Details

The object is constructed with the function scanInv and used as an input for listInv.

Examples

```
data(scanRes)
plot(scanRes)
```

setUpGenoDatFile

Loads genotype data onto R

Description

Loads onto an R session genotype data from text files or PLINK files.

Usage

```
setUpGenoDatFile(file = "GenoDat.txt", saveRes = FALSE, sortMinor = TRUE)
setUpGenoDatSNPmat(Chr, Geno, Annot, saveRes = FALSE, saveGeno = FALSE)
```

Arguments

file character. File path with genotype information

saveRes logical. Wether results should be saved into file gDat.RData

sortMinor logical. Whether genotypes should be sorted by minor allele frequency.

Chr numeric. Chromosome number

Geno snpMatrix. Matrix of raw with genotype data

Annot numeric. Annotation information read from an . bim file

saveGeno logical. Wether . txt file should be saved with genotype information

Value

GenoDat object

Author(s)

Alejandro Caceres <acaceres@creal.cat>

See Also

GenoDat

```
gen <- system.file("extdata", "genotypes.txt", package = "inveRsion")
gDat <-setUpGenoDatFile(file=gen,sortMinor=TRUE,saveRes=FALSE)
gDat</pre>
```

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