Package 'detectRUNS'

October 13, 2022

```
Type Package
Title Detect Runs of Homozygosity and Runs of Heterozygosity in
     Diploid Genomes
Version 0.9.6
Date 2019-10-24
Description Detection of runs of homozygosity and of heterozygosity
     in diploid genomes using two methods: sliding windows (Purcell et al (2007)
     <doi:10.1086/519795>) and consecutive runs (Marras et al (2015)
     <doi:10.1111/age.12259>).
Depends R (>= 3.0.0)
License GPL-3
LazyData TRUE
Encoding UTF-8
Imports plyr, iterators, itertools, ggplot2, reshape2, Rcpp,
     gridExtra, data.table
RoxygenNote 6.1.1
Suggests testthat, knitr, rmarkdown, prettydoc
LinkingTo Rcpp
VignetteBuilder knitr
URL https://github.com/bioinformatics-ptp/detectRUNS/tree/master/detectRUNS
BugReports https://github.com/bioinformatics-ptp/detectRUNS/issues
NeedsCompilation yes
Author Filippo Biscarini [aut, cre],
     Paolo Cozzi [aut],
     Giustino Gaspa [aut],
     Gabriele Marras [aut]
Maintainer Filippo Biscarini <filippo.biscarini@gmail.com>
Repository CRAN
Date/Publication 2019-10-24 13:40:02 UTC
```

2 consecutiveRUNS.run

R topics documented:

consecutiveRUNS.run		2
consecutiveRunsCpp		4
createRUNdf		5
findOppositeAndMissing		6
Froh_inbreeding		6
Froh_inbreedingClass		7
genoConvertCpp		8
heteroZygotTest		9
heteroZygotTestCpp		9
homoZygotTest		10
homoZygotTestCpp		11
pedConvertCpp		11
plot_DistributionRuns		12
plot_InbreedingChr		13
plot_manhattanRuns		14
plot_PatternRuns		15
plot_Runs		16
plot_SnpsInRuns		17
plot_StackedRuns		18
plot_ViolinRuns		19
readExternalRuns		20
readPOPCpp		21
reorderDF		21
slidingRUNS.run		22
slidingWindow		23
slidingWindowCpp		24
snpInRun		25
snpInRunCpp		25
snpInsideRuns		26
snpInsideRunsCpp		27
summaryRuns		27
tableRuns		28
writeRUN		29
		31

consecutiveRUNS.run

Main function to detect genomic RUNS (ROHom/ROHet) using the consecutive method

Description

Index

This is the main detectRUNS function to scan the genome for runs (of homozygosity or heterozygosity) using the consecutive method (Marras et al. 2015, Animal Genetics 46(2):110-121). All parameters to detect runs (e.g. minimum n. of SNP, max n. of missing genotypes, max n. of opposite genotypes etc.) are specified here. Input data are in the ped/map Plink format (https://www.coggenomics.org/plink/1.9/input#ped)

consecutiveRUNS.run 3

Usage

```
consecutiveRUNS.run(genotypeFile, mapFile, ROHet = FALSE,
  maxOppRun = 0, maxMissRun = 0, minSNP = 15, minLengthBps = 1000,
  maxGap = 10^6)
```

Arguments

genotypeFile genotype (.ped) file path mapFile map file (.map) file path

ROHet should we look for ROHet or ROHom? (default = FALSE)

maxOppRun max n. of opposite genotype SNPs in the run (default = 0)

maxMissRun max n. of missing SNPs in the run (default = 0) minSNP minimum n. of SNP in a RUN (default = 15)

minLengthBps minimum length of run in bps (defaults to 1000 bps = 1 kbps)

maxGap max distance between consecutive SNP in a window to be still considered a

potential run (defaults to 10^6)

Details

This function scans the genome (diploid) for runs using the consecutive method. This is a wrapper function for many component functions that handle the input data (ped/map files), performs internal conversions, accepts parameters specifications, selects the statistical method to detect runs (sliding windows, consecutive loci) and whether runs of homozygosity (RoHom) or of heterozygosity (RoHet) are looked for.

In the ped file, the groups samples belong to can be specified (first column). This is important if comparisons between human ethnic groups or between animal breeds or plant varieties or biological populations are to be performed. Also, if cases and controls are to be compared, this is the place where this information needs to be specified.

This function returns a data frame with all runs detected in the dataset. This data frame can then be written out to a csv file. The data frame is, in turn, the input for other functions of the detectRUNS package that create plots and produce statistics of the results (see plot and statistic functions in this manual, and/or refer to the vignette of detectRUNS).

Value

A dataframe with RUNs of Homozygosity or Heterozygosity in the analysed dataset. The returned dataframe contains the following seven columns: "group", "id", "chrom", "nSNP", "from", "to", "lengthBps" (group: population, breed, case/control etc.; id: individual identifier; chrom: chromosome on which the run is located; nSNP: number of SNPs in the run; from: starting position of the run, in bps; to: end position of the run, in bps; size of the run)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")</pre>
```

```
# calculating runs with consecutive run approach
## Not run:
# skipping runs calculation
runs <- consecutiveRUNS.run(genotypeFile, mapFile, minSNP = 15, ROHet = FALSE,
maxOppRun = 0, maxMissRun = 0, maxGap=10^6,
minLengthBps = 100000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.consecutive.csv", package="detectRUNS")
colClasses <- c(rep("character", 3), rep("numeric", 4) )
runs <- read.csv2(runsFile, header = TRUE, stringsAsFactors = FALSE,
colClasses = colClasses)</pre>
```

consecutiveRunsCpp

Function to detect consecutive runs in a vector (individual's genotypes)

Description

This is a core function. It implements the consecutive method for detection of runs in diploid genomes (see Marras et al. 2015)

Usage

```
consecutiveRunsCpp(indGeno, individual, mapFile, ROHet = TRUE,
    minSNP = 3L, maxOppositeGenotype = 1L, maxMiss = 1L,
    minLengthBps = 1000L, maxGap = 1000000L)
```

Arguments

indGeno vector of 0/1/NAs of individual genotypes (0: homozygote; 1: heterozygote) individual list of group (breed, population, case/control etc.) and ID of individual sample

mapFile Plink map file (for SNP position)

ROHet shall we detect ROHet or ROHom?

minSNP minimum number of SNP in a run

maxOppositeGenotype

max n. of homozygous/heterozygous SNP

maxMiss max. n. of missing SNP minLengthBps min length of a run in bps

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

createRUNdf 5

Details

The consecutive method detect runs by consecutively scanning SNP loci along the genome. No sliding windows are used. Checks on minimum n. of SNP, max n. of opposite and missing genotypes, max gap between adjacent loci and minimum length of the run are implemented (as in the sliding window method). Both runs of homozygosity (RoHom) and of heterozygosity (RoHet) can be search for (option ROHet: TRUE/FALSE)

Value

A data frame of runs per individual sample

createRUNdf Function to create a dataframe of RUNS per individual animal Re-

quires a map file (other filename to read or R object) Parameters on maximum number of missing and opposite genotypes in the run (not

the window) are implemented here

Description

Function to create a dataframe of RUNS per individual animal Requires a map file (other filename to read or R object) Parameters on maximum number of missing and opposite genotypes in the run (not the window) are implemented here

Usage

```
createRUNdf(snpRun, mapFile, minSNP = 3, minLengthBps = 1000,
  minDensity = 1/10, oppositeAndMissingSNP, maxOppRun = NULL,
  maxMissRun = NULL)
```

Arguments

snpRun vector of TRUE/FALSE (is the SNP in a RUN?)

mapFile Plink-like map file (data.frame)
minSNP minimum n. of SNP to call a RUN

minLengthBps minimum length of run in bps (defaults to 1000 bps = 1 kbps)

minDensity minimum n. of SNP per kbps (defaults to 0.1 = 1 SNP every 10 kbps)

oppositeAndMissingSNP

indexed array of missing and opposite genotypes (SNP order in the genome is

the index)

max OppRun max n. of opposite genotype SNPs in the run (not in the window!)

maxMissRun max n. of missing SNPs in the run (not in the window!)

Value

a data.frame with RUNS per animal

Froh_inbreeding

findOppositeAndMissing

Function to calculate oppositeAndMissingGenotypes array

Description

This is an helper function, this will be called by another function

Usage

```
findOppositeAndMissing(data, ROHet = TRUE)
```

Arguments

data vector of 0/1/2 genotypes

ROHet TRUE in ROHet evaluation, FALSE for ROHom

Value

character array; names will be index in which opposite and missing snps are found in data array

Froh_inbreeding

Function to calculated Froh genome-wide or chromosome-wide

Description

This function calculates the individual inbreeding coefficients based on runs of homozygosity (ROH), either per-chromosome (chromosome-wide) or based on the entire genome (genome-wide). See details of calculations below

Usage

```
Froh_inbreeding(runs, mapFile, genome_wide = TRUE)
```

Arguments

runs R object (dataframe) with results on runs
mapFile Plink map file (to retrieve SNP position)

genome_wide vector of TRUE/FALSE (genome-wide or chromosome-wide; defaults to TRUE/genome-

wide)

Froh_inbreedingClass

Details

Froh is calculated as:

$$F_{ROH} = \frac{\sum_{ROH_{length}}}{Length_{genome}}$$

Depending on whether genome-wide or chromosome-wide calculations are required, the terms in the numerator and denominator will refer to the entire genome or will be restricted to specific chromosomes.

7

Value

A data frame with the inbreeding coefficients of each individual sample

Examples

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

Froh_inbreeding(runs = runsDF, mapFile = mapFile)
Froh_inbreeding(runs = runsDF, mapFile = mapFile, genome_wide=FALSE)</pre>
```

Froh_inbreedingClass Function to calculated Froh using a ROH-class

Description

This function calculates the individual inbreeding coefficients based on runs of homozygosity (ROH) using only ROH of specific size classes. The parameter class specify the size interval to split up calculations. For example, if class = 2 Froh based on ROH 0-2, 2-4, 4-8, 80-16, >16 Mbps long will be calculated.

Usage

```
Froh_inbreedingClass(runs, mapFile, Class = 2)
```

8 genoConvertCpp

Arguments

runs R object (dataframe) with ROH results mapFile Plink map file (for SNP position)

Class base ROH-length interval (in Mbps) (default: 0-2, 2-4, 4-8, 8-16, >16)

Value

A data frame with individual inbreeding coefficients based on ROH-length of specific size. The sum of ROH-length of specific size in each individual is reported alongside

Examples

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

Froh_inbreedingClass(runs = runsDF, mapFile = mapFile, Class = 2)</pre>
```

genoConvertCpp

Convert 0/1/2 genotypes to 0/1

Description

This is a utility function, that convert 0/1/2 genotypes (AA/AB/BB) into 0/1 (either homozygous/heterozygous)

Usage

```
genoConvertCpp(genotype)
```

Arguments

genotype vector of 0/1/2 genotypes

Value

converted vector of genotypes (0/1)

heteroZygotTest 9

heteroZygotTest	Function to check whether a window is (loosely) heterozygous or not

Description

This is a core function within the sliding-window workflow. Parameters on how to consider a window heterozygous are here (maxHom, maxMiss)

Usage

```
heteroZygotTest(x, gaps, maxHom, maxMiss, maxGap, i, windowSize)
```

Arguments

x	vector of 0/1 genotypes (from genoConvert())
gaps	vector of differences between consecutive positions (gaps) in bps
maxHom	max n. of homozygous SNP in a heterozygous window
maxMiss	max n. of missing in a window
maxGap	max distance between consecutive SNP in a window to be still considered a potential run
i	index along the genome (genome-vector for each individual)
windowSize	size of window (n. of SNP)

Value

a list: i) TRUE/FALSE (whether a window is heterozygous or NOT); ii) indexes of "opposite and missing" genotype

heteroZygotTestCpp $F\iota$	unction to check whether a window is	(loosely) heterozygous or not
-----------------------------	--------------------------------------	-------------------------------

Description

This is a core function. Parameters on how to consider a window heterozygous are here (maxHom, maxMiss)

Usage

```
heteroZygotTestCpp(x, gaps, maxHom, maxMiss, maxGap)
```

10 homoZygotTest

Arguments

x vector of 0/1 genotypes (from genoConvert())

gaps vector of differences between consecutive positions (gaps) in bps

max n. of homozygous SNP in a heterozygous window

maxMiss max n. of missing in a window

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

Value

TRUE/FALSE (whether a window is heterozygous or NOT)

homoZygotTest Function to check whether a window is (loosely) homozygous or not

Description

This is a core function. Parameters on how to consider a window homozygous are here (maxHet, maxMiss)

Usage

homoZygotTest(x, gaps, maxHet, maxMiss, maxGap, i, windowSize)

Arguments

x vector of 0/1 genotypes (from genoConvert())

gaps vector of differences between consecutive positions (gaps) in bps

max n. of heterozygous SNP in a homozygous window

maxMiss max n. of missing in a window

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

i index along the genome (genome-vector for each individual)

windowSize size of window (n. of SNP)

Value

a list: i) TRUE/FALSE (whether a window is heterozygous or NOT); ii) indexes of "opposite and missing" genotype

homoZygotTestCpp 11

homoZygotTestCpp	Function to check whether a window is (loosely) homozygous or not

Description

This is a core function. Parameters on how to consider a window homozygous are here (maxHet, maxMiss)

Usage

```
homoZygotTestCpp(x, gaps, maxHet, maxMiss, maxGap)
```

Arguments

x vector of 0/1 genotypes (from genoConvert())

gaps vector of differences between consecutive positions (gaps) in bps

maxHet max n. of heterozygous SNP in a homozygous window

maxMiss max n. of missing in a window

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

Value

TRUE/FALSE (whether a window is homozygous or NOT)

Description

This is a utility function, that convert ped genotypes (AA/AB/BB) into 0/1 (either homozygous/heterozygous)

Usage

```
pedConvertCpp(genotype)
```

Arguments

genotype vector of pair of genotypes (01, AA, AG)

Value

converted vector of genotypes (0/1)

12 plot_DistributionRuns

Description

This function the distribution of runs per group. The average run length per size-class, the average run length per chromosome (and group), the percent distribution of runs per size-class and group, and the proportion of runs per chromosome are plotted. With style="All" all three plots are produced.

Usage

```
plot_DistributionRuns(runs, mapFile, groupSplit = TRUE,
   style = c("MeanClass", "MeanChr", "RunsPCT", "RunsPCT_Chr", "All"),
   savePlots = FALSE, outputName = NULL, plotTitle = NULL,
   Class = 2)
```

Arguments

R object (dataframe) with results on detected runs runs Plink map file (for SNP position) mapFile plots split by group (defaults to TRUE) groupSplit type of plot: MeanClass, MeanChr, RunsPCT, RunsPCT_Chr, All (all plots) style should plots be saved out to files or plotted in the graphical terminal (default)? savePlots outputName title prefix (the base name of graph, if savePlots is TRUE)#' plotTitle title in plot (default NULL) Class group of length (in Mbps) by class (default: 0-2, 2-4, 4-8, 8-16, >16)

Value

plot Distribution Runs

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data</pre>
```

plot_InbreedingChr 13

```
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')
plot_InbreedingChr(runs = runsDF, mapFile = mapFile, style='All')</pre>
```

plot_InbreedingChr

Plot Froh-based inbreeding coefficients by group

Description

The function plots the distribution of inbreeding/consanguinity coefficients per chromosome and/or group. Three types of plots can be produces: barplots, boxplots, violin plots. With style="All" all three plots are produced.

Usage

```
plot_InbreedingChr(runs, mapFile, groupSplit = TRUE,
   style = c("ChrBarPlot", "ChrBoxPlot", "FrohBoxPlot", "All"),
   outputName = NULL, plotTitle = NULL, savePlots = FALSE)
```

Arguments

runs R object (dataframe) with results on detected runs

mapFile Plink map file (for SNP position)

groupSplit plots split by group (defaults to TRUE)

style type of plot: ChrBarPlot, ChrBoxPlot, FrohBoxPlot, All (all plots)

outputName title prefix (the base name of graph, if savePlots is TRUE)

plotTitle title in plot (default NULL)

savePlots should plots be saved out to files or plotted in the graphical terminal (default)?

Value

plots of the distribution of inbreeding by chromosome and group

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)</pre>
```

14 plot_manhattanRuns

```
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')
plot_InbreedingChr(runs = runsDF, mapFile = mapFile, style='All')</pre>
```

plot_manhattanRuns $Plot \ the \ proportion \ of \ times \ SNPs \ are \ inside \ runs - MANHATTAN \ PLOT$

Description

Function to plot the proportion of times/percentage each SNP in inside a run (population-specific signals) against SNP position in all chromosomes together Proportions on the y-axis, bps on the x-axis for all analysed chromosomes This is similar to the familiar GWAS Manhattan plot

Usage

```
plot_manhattanRuns(runs, genotypeFile, mapFile, savePlots = FALSE,
  outputName = NULL, plotTitle = NULL)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length)
genotypeFile genotype (.ped) file path
mapFile map file (.map) file path
savePlots should plots be saved out in files (default) or plotted in the graphical terminal?
outputName title prefix (the base name of graph, if savePlots is TRUE)

plotTitle title in plot (default)

Value

Manhattan plots of proportion of times SNPs are inside runs, per population (pdf files)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)</pre>
```

plot_PatternRuns 15

```
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

# plot runs per animal (interactive)
plot_manhattanRuns(runs = runsDF, genotypeFile = genotypeFile, mapFile = mapFile,
savePlots = FALSE, plotTitle = "ROHom")</pre>
```

plot_PatternRuns

Plot sum of run-lengths (or average run-lengths) against the number of runs per individual

Description

Function to plot the sum of run lengths (or the average run length) per individual against the average number of runs per individual. Points can be differentially coloured by group/population. This plot can be useful to identify patterns in the distribution of runs in different groups (e.g. few long runs vs many short runs)

Usage

```
plot_PatternRuns(runs, mapFile, method = c("sum", "mean"),
  outputName = NULL, savePlots = FALSE, plotTitle = NULL)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length)
mapFile map file (.map) file path
method "sum" or "mean" of run lengths per individual sample
outputName title prefix (the base name of graph, if savePlots is TRUE)#'
savePlots should plots be saved out to files or plotted in the graphical terminal (default)?

plotTitle title in plot (default NULL)

Value

plot of number of runs vs run-length sum/mean per individual sample

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")
# calculating runs of Homozygosity
## Not run:</pre>
```

plot_Runs

```
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

plot_PatternRuns(runs = runsDF, mapFile = mapFile, method = 'sum')
plot_PatternRuns(runs = runsDF, mapFile = mapFile, method = 'mean')</pre>
```

plot_Runs

Function to plot runs per individual

Description

Function to plot runs per individual (see Williams et al. 2016, Animal Genetics, for an example with animal data) Individual IDs on the y-axis, bps on the x-axis (position along the chromosome)

Usage

```
plot_Runs(runs, suppressInds = FALSE, savePlots = FALSE,
    separatePlots = FALSE, outputName = NULL)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length)
suppressInds shall we suppress individual IDs on the y-axis? (defaults to FALSE)
savePlots should plots be saved out to files (one pdf file for all chromosomes) or plotted in the graphical terminal (default)?
separatePlots should plots for each chromosome be saved out to separate files?
outputName title prefix (the base name of graph, if savePlots is TRUE)

Value

plot of runs by chromosome

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation</pre>
```

plot_SnpsInRuns 17

```
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

# plot runs per animal (interactive)
plot_Runs(runs = runsDF, suppressInds = FALSE, savePlots = FALSE, outputName = "ROHom")</pre>
```

plot_SnpsInRuns

Plot the number of times each SNP falls inside runs

Description

Function to plot the number of times/percentage each SNP is inside a run (population-specific signals) against the SNP positions in the genome. Proportions on the y-axis, bps on the x-axis

Usage

```
plot_SnpsInRuns(runs, genotypeFile, mapFile, savePlots = FALSE,
    separatePlots = FALSE, outputName = NULL)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length)

genotypeFile genotype (.ped) file path mapFile map file (.map) file path

savePlots should plots be saved out in files (default) or plotted in the graphical terminal?

separatePlots should plots for each chromosome be saved out to separate files?

outputName title prefix (the base name of graph, if savePlots is TRUE)

Value

plot number of times a SNP is in a run by chromosome and population (pdf files)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
# skipping runs calculation
## Not run:
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,</pre>
```

18 plot_StackedRuns

```
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')
# plot runs per animal (interactive)
plot_SnpsInRuns(runs = runsDF, genotypeFile = genotypeFile, mapFile = mapFile,
savePlots = FALSE, outputName = "ROHom")</pre>
```

plot_StackedRuns

Plot stacked runs

Description

Function to plot stacked runs along the chromosome (signaling presence of large numbers of runs in specific regions of a chromosome) Counts on the y-axis, bps on the x-axis (position along the chromosome)

Usage

```
plot_StackedRuns(runs, savePlots = FALSE, separatePlots = FALSE,
  outputName = NULL)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length) savePlots should plots be saved out in files (default) or plotted in the graphical terminal?

separatePlots should plots for chromosomes be saved out to separate files? outputName title prefix (the base name of graph, if savePlots is TRUE)

Value

plot of stacked runs by population and by chromosome (pdf files)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)</pre>
```

plot_ViolinRuns 19

```
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')
# plot runs per animal (interactive)
plot_StackedRuns(runs = runsDF, savePlots = FALSE, outputName = "ROHom")</pre>
```

plot_ViolinRuns

Violin plot of run length per individual (either sum or mean)

Description

Function to produce violin plots of the distribution of runs lengths per group The sum of run lengths, or its average, per individual sample is used to characterize the distribution of runs

Usage

```
plot_ViolinRuns(runs, method = c("sum", "mean"), outputName = NULL,
  plotTitle = NULL, savePlots = FALSE)
```

Arguments

runs a data.frame with runs per individual (group, id, chrom, nSNP, start, end, length)

method "sum" or "mean" of run lengths per individual samples outputName title prefix (the base name of graph, if savePlots is TRUE)

plotTitle title in plot (default NULL)

savePlots should plots be saved out to files or plotted in the graphical terminal (default)?

Value

Violin plot of the distribution of runs-lengths (sum or mean)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)
## End(Not run)</pre>
```

20 readExternalRuns

```
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

plot_ViolinRuns(runs = runsDF, method = "sum" , savePlots = FALSE)
plot_ViolinRuns(runs = runsDF, method = "mean" , savePlots = FALSE)</pre>
```

readExternalRuns

Read runs from external files

Description

Function to read in, from external files, the output of software for ROH:

- 1. detectRUNS: output saved out to a file (e.g. write.table)
- 2. Plink: output from the --homozyg option (.hom files)
- 3. BCFtools: output from the roh option

Usage

```
readExternalRuns(inputFile = NULL, program = c("plink", "BCFtools",
   "detectRUNS"))
```

Arguments

```
inputFile name of (path to) external file

program source program that produced the ROH file (one of detectRUNS, Plink, BCFtools)
```

Value

dataframe in the correct format to be used with plots and statistics functions from detectRUNS

```
# getting map and ped paths
## Not run:
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxMissRun = 1, maxMissWindow = 1, minLengthBps = 100000, minDensity = 1/10000)

write.table(x= runs,file = 'RunsFileTest.txt', quote=F, row.names = F)
newData=readRunsFromFile(runsFile = 'RunsFileTest.txt', program = 'detectRUNS')

## End(Not run)</pre>
```

readPOPCpp 21

readPOPCpp

Function to return a dataframe of population (POP, ID)

Description

This is a core function. Read PED file and returns a data frame with the first two columns

Usage

```
readPOPCpp(genotypeFile)
```

Arguments

```
genotypeFile genotype (.ped) file location
```

Value

a dataframe of POP, ID

reorderDF

Function to reorder data frames by CHROMOSOME

Description

The data frame will be reordered according to chromosome: from 1 to n, then X, Y, XY, MT The data frame needs to have a column with name "CHROMOSOME"

Usage

```
reorderDF(dfx)
```

Arguments

dfx

data frame to be reordered (with column "CHROMOSOME")

Details

Reorder results based on chromosome

Value

A reordered data frame by chromosome

22 slidingRUNS.run

slidingRUNS.run	Main function to detect RUNS (ROHom/ROHet) using sliding windows (a la Plink)
-----------------	---

Description

This is one of the main function of detectRUNS and is used to detect runs (of homozygosity or heterozygosity) in the genome (diploid) with the sliding-window method. All parameters to detect runs (e.g. minimum n. of SNP, max n. of missing genotypes, max n. of opposite genotypes etc.) are specified here. Input data are in the ped/map Plink format (https://www.coggenomics.org/plink/1.9/input#ped)

Usage

```
slidingRUNS.run(genotypeFile, mapFile, windowSize = 15,
   threshold = 0.05, minSNP = 3, ROHet = FALSE, maxOppWindow = 1,
   maxMissWindow = 1, maxGap = 10^6, minLengthBps = 1000,
   minDensity = 1/1000, maxOppRun = NULL, maxMissRun = NULL)
```

Arguments

genotypeFile	genotype (.ped) file path
mapFile	map file (.map) file path
windowSize	the size of sliding window (number of SNP loci) (default = 15)
threshold	the threshold of overlapping windows of the same state (homozygous/heterozygous) to call a SNP in a RUN (default = 0.05)
minSNP	minimum n. of SNP in a RUN (default = 3)
ROHet	should we look for ROHet or ROHom? (default = FALSE)
maxOppWindow	\max n. of homozygous/heterozygous SNP in the sliding window (default = 1)
maxMissWindow	max. n. of missing SNP in the sliding window (default = 1)
maxGap	max distance between consecutive SNP to be still considered a potential run (default = 10^6 bps)
minLengthBps	minimum length of run in bps (defaults to 1000 bps = 1 kbps)
minDensity	minimum n. of SNP per kbps (defaults to 0.1 = 1 SNP every 10 kbps)
maxOppRun	max n. of opposite genotype SNPs in the run (optional)
maxMissRun	max n. of missing SNPs in the run (optional)

Details

This function scans the genome (diploid) for runs using the sliding-window method. This is a wrapper function for many component functions that handle the input data (ped/map files), perform internal conversions, accept parameters specifications, select whether runs of homozygosity (RoHom) or of heterozygosity (RoHet) are looked for.

slidingWindow 23

In the ped file, the groups samples belong to can be specified (first column). This is important if comparisons between human ethnic groups or between animal breeds or plant varieties or biological populations are to be performed. Also, if cases and controls are to be compared, this is the place where this information needs to be specified.

This function returns a data frame with all runs detected in the dataset. This data frame can then be written out to a csv file. The data frame is, in turn, the input for other functions of the detectRUNS package that create plots and produce statistics from the results (see plots and statistics functions in this manual, and/or refer to the detectRUNS vignette).

Value

A dataframe with RUNs of Homozygosity or Heterozygosity in the analysed dataset. The returned dataframe contains the following seven columns: "group", "id", "chrom", "nSNP", "from", "to", "lengthBps" (group: population, breed, case/control etc.; id: individual identifier; chrom: chromosome on which the run is located; nSNP: number of SNPs in the run; from: starting position of the run, in bps; to: end position of the run, in bps; size of the run)

Examples

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")
# calculating runs with sliding window approach
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1,
minSNP = 15, ROHet = FALSE, maxOppWindow = 1, maxMissWindow = 1, maxGap=10^6,
minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
colClasses <- c(rep("character", 3), rep("numeric", 4) )
runs <- read.csv2(runsFile, header = TRUE, stringsAsFactors = FALSE,
colClasses = colClasses)</pre>
```

slidingWindow

Function to slide a window over a vector (individual's genotypes)

Description

This is a core function. The functions to detect RUNS are slid over the genome

Usage

```
slidingWindow(data, gaps, windowSize, step, maxGap, ROHet = TRUE,
  maxOppositeGenotype = 1, maxMiss = 1)
```

24 slidingWindowCpp

Arguments

data vector of 0/1/2 genotypes

gaps vector of differences between consecutive positions (gaps) in bps

windowSize size of window (n. of SNP)

step by which (how many SNP) is the window slid

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

ROHet shall we detect ROHet or ROHom?

maxOppositeGenotype

max n. of homozygous/heterozygous SNP

maxMiss max. n. of missing SNP

Value

vector of TRUE/FALSE (whether a window is homozygous or NOT)

slidingWindowCpp Function to slide a window over a vector (individual's genotypes)

Description

This is a core function. The functions to detect RUNS are slid over the genome

Usage

```
slidingWindowCpp(data, gaps, windowSize, step, maxGap, ROHet = TRUE,
   maxOppositeGenotype = 1L, maxMiss = 1L)
```

Arguments

data vector of 0/1/2 genotypes

gaps vector of differences between consecutive positions (gaps) in bps

windowSize size of window (n. of SNP)

step by which (how many SNP) is the window slid

maxGap max distance between consecutive SNP in a window to be still considered a

potential run

ROHet shall we detect ROHet or ROHom?

maxOppositeGenotype

max n. of homozygous/heterozygous SNP

maxMiss max. n. of missing SNP

Value

vector of TRUE/FALSE (whether a window is homozygous or NOT)

snpInRun 25

snpInRun	Function to return a vector of T/F for whether a SNP is or not in a RUN

Description

This is a core function. The function to determine whether a SNP is or not in a RUN. The ratio between homozygous/heterozygous windows and total n. of windows is computed here

Usage

```
snpInRun(RunVector, windowSize, threshold)
```

Arguments

RunVector vector of TRUE/FALSE (is a window homozygous/heterozygous?)

windowSize size of window (n. of SNP)

threshold to call a SNP in a RUN

Value

vector of TRUE/FALSE (whether a SNP is in a RUN or NOT)

snpInRunCpp	Function to return a vector of T/F for whether a SNP is or not in a RUN

Description

This is a core function. The function to determine whether a SNP is or not in a RUN. The ratio between homozygous/heterozygous windows and total n. of windows is computed here

Usage

```
snpInRunCpp(RunVector, windowSize, threshold)
```

Arguments

RunVector vector of TRUE/FALSE (is a window homozygous/heterozygous?)

windowSize size of window (n. of SNP)
threshold threshold to call a SNP in a RUN

Value

vector of TRUE/FALSE (whether a SNP is in a RUN or NOT)

26 snpInsideRuns

snpInsideRuns	Function to count number of times a SNP is in a RUN
Shpinsidendis	1 unction to count number of times a 5111 is in a Ref.

Description

Function to count number of times a SNP is in a RUN

Usage

```
snpInsideRuns(runsChrom, mapChrom, genotypeFile)
```

Arguments

runsChrom R object (dataframe) with results per chromosome (column names: "POPULATION", "IND", "CHROMOS

mapChrom R object (dataframe) with SNP name and position per chromosome (map file)

(column names: "CHR", "SNP_NAME", "x", "POSITION")

genotypeFile genotype (.ped) file location

Value

dataframe with counts per SNP in runs (per population)

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")</pre>
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")</pre>
# defining mapChrom
mappa <- data.table::fread(mapFile, header = FALSE)</pre>
names(mappa) <- c("CHR", "SNP_NAME", "x", "POSITION")</pre>
mappa$x <- NULL
chrom <- "24"
mapChrom <- mappa[mappa$CHR==chrom, ]</pre>
# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)
## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
colClasses <- c(rep("character", 3), rep("numeric", 4) )</pre>
runs <- read.csv2(runsFile, header = TRUE, stringsAsFactors = FALSE,
colClasses = colClasses)
# fix column names and define runsChrom
```

snpInsideRunsCpp 27

```
names(runs) <- c("POPULATION","IND","CHROMOSOME","COUNT","START","END","LENGTH")
runsChrom <- runs[runs$CHROMOSOME==chrom, ]
snpInsideRuns(runsChrom, mapChrom, genotypeFile)</pre>
```

snpInsideRunsCpp

Function to count number of times a SNP is in a RUN

Description

Function to count number of times a SNP is in a RUN

Usage

```
snpInsideRunsCpp(runsChrom, mapChrom, genotypeFile)
```

Arguments

runsChrom R object (dataframe) with results per chromosome

mapChrom R map object with SNP per chromosome

genotypeFile genotype (.ped) file location

Value

dataframe with counts per SNP in runs (per population)

summaryRuns

Summary statistics on detected runs

Description

This function processes the results from slidingRUNS.run and consecutiveRUNS.run and produces a number of interesting descriptives statistics on results.

Usage

```
summaryRuns(runs, mapFile, genotypeFile, Class = 2, snpInRuns = FALSE)
```

Arguments

runs R object (dataframe) with results on detected runs

mapFile Plink map file (for SNP position)
genotypeFile Plink ped file (for SNP position)

Class group of length (in Mbps) by class (default: 0-2, 2-4, 4-8, 8-16, >16)

snpInRuns TRUE/FALSE (default): should the function snpInsideRuns be called to com-

pute the proportion of times each SNP falls inside a run in the group/population?

28 tableRuns

Details

summaryRuns calculates: i) the number of runs per chromosome and group/population; ii) the percent distribution of runs per chromosome and group; iii) the number of runs per size-class and group; iv) the percent distribution of runs per size-class and group; v) the mean length of runs per chromosome and group; vi) the mean length of runs per size-class and group; vii) individual inbreeding coefficient estimated from ROH; viii) individual inbreeding coefficient estimated from ROH per size-class

Value

A list of dataframes containing the most relevant descriptives statistics on detected runs. The list conveniently contains 9 dataframes that can be used for further processing and visualization, or can be written out to text files

Examples

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF <- readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

summaryRuns(runs = runsDF, mapFile = mapFile, genotypeFile = genotypeFile, Class = 2,
snpInRuns = FALSE)</pre>
```

tableRuns

Function to retrieve most common runs in the population

Description

This function takes in input either the run results or the output from the function snpInsideRuns (proportion of times a SNP is inside a run) in the population/group, and returns a subset of the runs most commonly found in the group/population. The parameter threshold controls the definition of most common (e.g. in at least 50%, 70% etc. of the sampled individuals)

Usage

```
tableRuns(runs = NULL, SnpInRuns = NULL, genotypeFile, mapFile,
    threshold = 0.5)
```

writeRUN 29

Arguments

runs R object (dataframe) with results on detected runs

SnpInRuns dataframe with the proportion of times each SNP falls inside a run in the popu-

lation (output from snpInsideRuns)

genotypeFile Plink ped file (for SNP position)
mapFile Plink map file (for SNP position)

threshold value from 0 to 1 (default 0.7) that controls the desired proportion of individuals

carrying that run (e.g. 70%)

Value

A dataframe with the most common runs detected in the sampled individuals (the group/population, start and end position of the run, chromosome and number of SNP included in the run are reported in the output dataframe)

Examples

```
# getting map and ped paths
genotypeFile <- system.file("extdata", "Kijas2016_Sheep_subset.ped", package = "detectRUNS")
mapFile <- system.file("extdata", "Kijas2016_Sheep_subset.map", package = "detectRUNS")

# calculating runs of Homozygosity
## Not run:
# skipping runs calculation
runs <- slidingRUNS.run(genotypeFile, mapFile, windowSize = 15, threshold = 0.1, minSNP = 15,
ROHet = FALSE, maxOppositeGenotype = 1, maxMiss = 1, minLengthBps = 100000, minDensity = 1/10000)

## End(Not run)
# loading pre-calculated data
runsFile <- system.file("extdata", "Kijas2016_Sheep_subset.sliding.csv", package="detectRUNS")
runsDF = readExternalRuns(inputFile = runsFile, program = 'detectRUNS')

tableRuns(runs = runsDF, genotypeFile = genotypeFile, mapFile = mapFile, threshold = 0.5)</pre>
```

writeRUN

Function to write out RUNS per individual animal

Description

Function to write out RUNS per individual animal

Usage

```
writeRUN(ind, dRUN, ROHet = TRUE, group, outputName)
```

30 writeRUN

Arguments

ind ID of animals

dRUN data.frame with RUNS per animal ROHet shall we detect ROHet or ROHom?

group group (factor): population, breed, ethnicity, case/control etc.

outputName output filename

Value

TRUE/FALSE if RUNS are written out or not

Index

```
consecutiveRUNS.run, 2
consecutiveRunsCpp, 4
createRUNdf, 5
findOppositeAndMissing, 6
Froh_inbreeding, 6
Froh_inbreedingClass, 7
genoConvertCpp, 8
heteroZygotTest, 9
heteroZygotTestCpp, 9
homoZygotTest, 10
homoZygotTestCpp, 11
pedConvertCpp, 11
plot_DistributionRuns, 12
plot_InbreedingChr, 13
plot_manhattanRuns, 14
plot_PatternRuns, 15
plot_Runs, 16
plot_SnpsInRuns, 17
plot_StackedRuns, 18
plot_ViolinRuns, 19
readExternalRuns, 20
readPOPCpp, 21
reorderDF, 21
slidingRUNS.run, 22
slidingWindow, 23
slidingWindowCpp, 24
snpInRun, 25
snpInRunCpp, 25
snpInsideRuns, 26
snpInsideRunsCpp, 27
summaryRuns, 27
tableRuns, 28
writeRUN, 29
```