Package 'fcScan'

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Type Package

Title fcScan for detecting clusters of coordinates with user defined options

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Author Abdallah El-

Kurdi <ak161@aub.edu.lb>Ghiwa khalil <gk39@aub.edu.lb>Georges Khazen <gkhazen@lau.edu.lb>Pierre Khou

Maintainer Pierre Khoueiry <pk17@aub.edu.lb> Abdallah El-Kurdi <ak161@aub.edu.lb>

Description This package is used to detect combination of genomic coordi-

nates falling within a user defined window size along with user defined overlap between identified neighboring clusters. It can be used for genomic data where the clusters are built on a specific chromosome or specific strand. Clustering can be performed with a ``greedy" option allowing thus the presence of additional sites within the allowed window size.

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Encoding UTF-8

Imports stats, plyr, VariantAnnotation, SummarizedExperiment, rtracklayer, GenomicRanges, methods, IRanges

VignetteBuilder knitr

Suggests RUnit, BiocGenerics, BiocStyle, knitr, rmarkdown

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getCluster

Description

Given a data frame or a GRanges object or a list of bed/vcf files, this function will look for clusters of genomic features found within a user defined window size and satisfying user defined categorical conditions.

Usage

```
getCluster(x, w, c, overlap = 0, greedy = FALSE, seqnames = NULL, s = "*",
order = NULL, sites_orientation = NULL, verbose = FALSE)
```

Arguments

x	Vector of file names or a data frame	
W	Value for the desired cluster size (numeric)	
с	A named vector for condition relative to each site name	
overlap	numeric; If negative, it will correspond to the maximum overlap allowed be- tween consecutive clusters, if positive, it will correspond to the minimum gap allowed between consecutive clusters. The default is set to 0	
greedy	logical; If FALSE, the formed clusters will contain the exact combination of sites defined in "c". If TRUE, clusters containing the condition and more will be labeled as TRUE as long as window size is below or equal the defined value. The default is set to FALSE	
seqnames	character; seqname/chromosome to cluster on. The default is set to NULL so the clustering will be performed on all seqnames	
S	character; Strand to cluster on. Can be "+", "-" or "*". The default is set to "*" which indicate both clusters	
order	vector; A vector containing the order of sites the user wants the clusters to con- tain. By default it is set to NULL	
sites_orientation		
	vector; A vector containing the orientation of the sites specified by the user. By default it is set to NULL	
verbose	logical; A If FALSE, only clusters that passed combination condition are shown. used for debugging purposes. By default it is set to FALSE	

Details

The function *getCluster* will cluster coordinates based on a user defined number of sites and window size. The user needs to specify the condition for clustering, which is the sites and the number of sites required in each cluster. The user can exclude sites from clusters. this should be specified also in the condition for clustering. the user can also set the distance required between consective clusters using the overlap argument. If overlap is a negative, this will represent the maximum overlap allowed between clusters. If overlap is positive, this means that the clusters should have a minimum gap of the given value.

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The user can also choose to cluster on a specific strand by using the strand option, and specific seqname by specifying the seqnames as an argument.

x is a vector of files or a data frame. When data frame is given as input, it should have 5 columns *seqnames start end strand site. start* and *end* column need to be numeric or integer and the rest of the columns are of type character.

The Greedy option controls if more sites are allowed in the cluster. When set to TRUE, getCluster will return clusters containing the required number of sites and more as long as the window size condition is satisfied. When set to FALSE, getCluster will return clusters containing the exact number of the required sites. Default is set to FALSE.

order allows the user to choose whether the clusters should contain sites ordered in a certain way. When set to NULL, sites order is not important. When defined, the order of the sites in the cluster must abide by the user defined order. Default is NULL.

Value

The returned value is a GRanges object containing the clusters.

Author(s)

Abdallah El-Kurdi Ghiwa Khalil Georges Khazen Pierre Khoueiry

Examples

```
x = data.frame(seqnames = rep("chr1", times = 16),
start = c(10,17,25,27,32,41,47,60,70,87,94,99,107,113,121,132),
end = c(15,20,30,35,40,48,55,68,75,93,100,105,113,120,130,135),
strand = rep("+", 16),
site = c("s1","s2","s2","s1","s2","s1","s1","s2",
"s1","s2","s2","s1","s2","s1","s1","s2"))
```

```
clusters = getCluster(x, w = 25, c = c("s1"=1,"s2"=2),
greedy = TRUE, overlap = -5, s = "+", order = c("s1","s2","s1"))
```

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