Package 'CORE'

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Title Co	ores of Recurrent Events	
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CORE	Cores of Recurrent Events	
Descript	tion	
	en a collection of intervals $s_1,,s_N$, find K intervals $c_1,,c_K$ which approximately minite Sum_i Prod_k (1-E(s_i,c_k)), where E(s_i,c_k) is a geometric measure of association between	

s_i and c_k. Perform permutation tests to estimate the significance of finding.

CORE CORE

Usage

```
CORE(dataIn, keep = NULL, startcol = "start", endcol = "end",
chromcol = "chrom", weightcol = "weight", maxmark = 1, minscore = 0,
pow = 1, assoc = c("I", "J", "P"), nshuffle = 0, boundaries = NULL,
seedme = sample(1e+08, 1), shufflemethod = c("SIMPLE", "RESCALE"),
tiny = -1, distrib = c("vanilla", "Rparallel", "Grid"), njobs = 1, qmem=NA)
```

Arguments

dataIn A matrix, a data frame or an object of class "CORE". If dataIn is a matrix or a

data frame, it should have columns with names specified by the startcol and

endcol arguments, otherwise the function exits with an error.

keep A character vector. If dataIn is of class "CORE", keep specifies the names of

items of dataIn to be kept at their input values. These values take precedence over the corresponding argument values as specified in the function call. keep

is ignored if dataIn is not of class "CORE".

startcol A character string. If dataIn is a matrix or a data frame, startcol specifies the

name of the column containing start coordinates of the input intervals. Other-

wise startcol is ignored.

endcol A character string. If dataIn is a matrix or a data frame, endcol specifies the

name of the column containing end coordinates of the input intervals. Otherwise

endcol is ignored.

chromcol A character string. If dataIn is a matrix or a data frame, chromcol specifies

the name of the column containing chromosome numbers of the input intervals.

Otherwise chromcol is ignored.

weightcol A character string. If dataIn is a matrix or a data frame, weightcol specifies the

name of the column containing initial weights of the input intervals. Otherwise

weightcol is ignored.

maxmark An integer for the maximal number of cores to be computed. The actual number

of cores to be computed is the smaller of maxmark and the number of cores with

scores exceeding minscore.

minscore A single numeric value for the minimal allowed score of the cores to be reported.

pow A single numeric value of at least 1 for the power parameter used in computing

the association measure beween the cores and the input intervals (see Details).

assoc A character specifying the type of association measure to be used (see Details).

nshuffle An integer specifying the number of randomizations to be performed for esti-

mating significance.

boundaries A matrix or a data frame that must have three columns whose names are given

by chromcol, startcol and endcol. These specify the chromosome numbers $\,$

and their start and end positions (see Details).

seedme An integer specifying the random number generator seed (see Details).

shufflemethod A character string specifying the event randomization method used for estima-

tion of significance. If "SIMPLE" (default), each event is placed at random with equal probability for any position where it can fit within chromosome boundaries. If "RESCALE", each event is placed at random in a randomly chosen

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chromosome, and the event length is multiplied by the length ratio of the new to

the original chromosome.

tiny A single numeric value specifying the weight below which events are removed

from the input event set.

distrib A character string specifying the method of distributed computing used for es-

timation of significance. If "vanilla" (default), no distributed computing is performed. If "Rparallel", parallel computation with the local machine is performed using functions from CRAN core package parallel, with the number of worker processes being the smaller number of njobs, and nshuffle. If "Grid", parallel computation with grid engine is performed. The number of submitted array jobs, or cores that are distributed, is the smaller number of njobs, and nshuffle. When using "Grid", make sure you have write premission to the current work

space.

njobs If distributed computing is used for estimation of significance, a single integer

specifying the desired number of worker processes.

qmem A character string that can customize grid engine qsub command. The com-

mand decides memory size per core(each job). The default substring is "-1 vir-

tual free=2G".

Details

The three measures of association specified by assoc are defined as follows (\parallel denotes the length of an interval). For "I" (inclusion) $E(s_i,c_k) = (|c_k|/|s_i|)^p$ ow if c_k is contained in s_i and 0 otherwise. For "J" (Jaccard) $E(s_i,c_k) = J(s_i,c_k)^p$ ow, where J is the Jaccard index. For "P" (piercing) $E(s_i,c_k) = 1$ if c_k is contained and 0 otherwise. In all cases the left (right) boundary of an optimal c_k is one of the left (right) boundaries in the set of input interval events. In addition, there are no event interval boundaries in the interior of an optimal c_k in case "P".

The boundaries argument is used for assessing statistical significance of the solution. If boundaries is not specified, the chromosome boundaries for each chromosome are taken to be the leftmost left and the rightmost right boundaries of all events in the chromosome.

If significance of finding is estimated, the random number generator stream, and hence the resultant estimate, only depends on seedme and is independent of the parallelization option chosen.

Value

An object of class "CORE" with the following items.

input A matrix with four columns called "chrom", "start", "end" and "weight", speci-

fying the input interval events.

call A character string specifying the function call.

coreTable A matrix with columns named "start", "end" and "score", for start and end posi-

tions and CORE scores of the cores found by the algorithm.

seedme If significance estimate was performed, the random number generator seed.

assoc One of "I", "J" or "P", indicating the geometric measure of association used.

shufflemethod One of "SIMPLE" or "RESCALE", indicating the randomization method used.

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p A numeric vector of the length equal to the row dimension of coreTable con-

taining estimated p-values for the cores.

simscores A matrix with the row dimension equal to that of coreTable and nshuffle

columns, containing core scores computed for nshuffle sets of randomized

events.

minscore A single numeric value for the minimal score of the reported cores.

maxmark A single numeric value for the requested maximal number of cores to be com-

puted.

tiny A single numeric value for the weight below which events were removed from

the input set.

pow A single numeric value for the power used in computing the association mea-

sures.

boundaries A matrix with three columns named "chrom", "start" and "end", indicating chro-

mosome numbers and boundary positions used for estimation of significance.

Author(s)

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Examples

```
#Compute 3 cores and perform no randomization
#(meaningless for estimate of significance).
data(testInputCORE)
data(testInputBoundaries)
myCOREobj<-CORE(dataIn=testInputCORE,maxmark=3,nshuffle=0,
boundaries=testInputBoundaries,seedme=123)
## Not run:
#Extend this computation to a much larger number of randomizations,
#using 2 cores of a host computer.
newCOREobj<-CORE(dataIn=myCOREobj,keep=c("maxmark","seedme","boundaries"),</pre>
nshuffle=20,distrib="Rparallel",njobs=2)
#When using "Grid", make sure you have write premission to the current
#work space.
newCOREobj<-CORE(dataIn=myCOREobj,keep=c("maxmark","seedme","boundaries"),</pre>
nshuffle=20,distrib="Grid",njobs=2)
## End(Not run)
```

testInputBoundaries

A table of chromosome boundary positions for DNA copy number analysis

Description

The entire length of the genome was divided into 50009 bins, with chromosomes laid out in the usual genomic order: 1,...,22,X,Y. Each observation in the table provides the start and end bin numbers of each chromosome (chrom).

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Usage

```
data(testInputBoundaries)
```

Format

A data frame with 24 observations on the following 3 variables.

```
chrom a numeric vector
start a numeric vector
end a numeric vector
```

References

Navin N, Kendall J, Troge J, Andrews P, Rodgers L, McIndoo J, Cook K, Stepansky A, Levy D, Esposito D et al. 2011. Tumour evolution inferred by single-cell sequencing. Nature 472(7341): 90-U119.

Examples

```
data(testInputBoundaries)
## maybe str(testInputBoundaries) ; plot(testInputBoundaries) ...
```

testInputCORE

A table of DNA copy number gain events observed in 100 individual tumor cells

Description

Each observation in the table corresponds to a DNA copy number gain event in one of 100 individual breast cancer cells. The entire length of the genome was divided into 50009 bins. An event is an interval in chromosome chrom whose start and end bin numbers are given by start and end.

Usage

```
data(testInputCORE)
```

Format

A data frame with 2490 observations on the following 3 variables.

```
chrom a numeric vector
start a numeric vector
end a numeric vector
```

References

Navin N, Kendall J, Troge J, Andrews P, Rodgers L, McIndoo J, Cook K, Stepansky A, Levy D, Esposito D et al. 2011. Tumour evolution inferred by single-cell sequencing. Nature 472(7341): 90-U119.

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Examples

```
data(testInputCORE)
## maybe str(testInputCORE) ; plot(testInputCORE) ...
```

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