Package 'dStruct'

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туре гаскаде
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R topics documented:
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2 dCombs

calcDis

Calculates d score.

Description

Calculates d score.

Usage

calcDis(x)

Arguments

Χ

A numeric vector or matrix.

Value

If input is a numeric vector, a number is returned. For a matrix, a numeric vector is returned.

dCombs

Assesses within-group variation.

Description

Assesses within-group variation.

Usage

```
dCombs(rdf, combs)
```

Arguments

rdf Dataframe of reactivities for each sample.

combs Data.frame with each column containing groupings of samples.

Value

Nucleotide-wise d score.

dStruct 3

dStruct	Performs de novo discovery of differentially reactive regions.	

Description

Performs de novo discovery of differentially reactive regions.

Usage

```
dStruct(rdf, reps_A, reps_B, batches = F, min_length = 11,
  check_signal_strength = T, check_nucs = T, check_quality = T,
  quality = "auto", evidence = 0, signal_strength = 0.1,
  within_combs = NULL, between_combs = NULL, ind_regions = T, gap = 1,
  get_FDR = T)
```

Arguments

rdf	Dataframe of reactivities for each sample.
reps_A	Number of replicates of group A.
reps_B	Number of replicates of group B.
batches	Logical suggesting if replicates of group A and B were performed in batches and are labelled accordingly.
min_length	Minimum length of constructed regions.
check_signal_s	trength
	Logical, if TRUE, constructed regions must have a minimum signal strength.
check_nucs	Logical, if TRUE, constructed regions must have a minimum number of nucleotides participating in Wilcoxon signed rank test.
check_quality	Logical, if TRUE, check constructed regions for quality.
quality	Worst allowed quality for a region to be tested.
evidence	Minimum evidence of between-group variation for a region to be tested.
signal_strengt	h
	Threshold for minimum signal strength.
within_combs	Data.frame with each column containing groupings of replicates of groups A or B, which will be used to assess within-group variation.
between_combs	Dataframe with each column containing groupings of replicates of groups A and B, which will be used to assess between-group variation.
ind_regions	Logical, if TRUE, test each region found in the transcript separately.
gap	Integer. Join regions if they are separated by these many nucleotides.
get_FDR	Logical, if FALSE, FDR is not reported.

Value

Constructs regions, reports p-values and FDR for them.

dStructome

dStruct.guided	Performs guided discovery of differentially reactive regions.

Description

Performs guided discovery of differentially reactive regions.

Usage

```
dStruct.guided(rdf, reps_A, reps_B, batches = F, within_combs = NULL,
  between_combs = NULL, check_quality = TRUE, quality = "auto",
  evidence = 0)
```

Arguments

rdf	Dataframe of reactivities for each sample.
reps_A	Number of replicates of group A.
reps_B	Number of replicates of group B.
batches	Logical suggesting if replicates of group A and B were performed in batches and are labelled accordingly.
within_combs	Data.frame with each column containing groupings of replicates of groups A or B, which will be used to assess within-group variation.
between_combs	Dataframe with each column containing groupings of replicates of groups A and B, which will be used to assess between-group variation.
check_quality	Logical, if TRUE, check regions for quality.
quality	Worst allowed quality for a region to be tested.
evidence	Minimum evidence of between-group variation for a region to be tested.

Value

p-value for the tested region, estimated using one-sided Wilcoxon signed rank test.

transcriptome-wide data.	dStr	uctome	v			of	differentially	reactive	regions	for
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Description

Performs de novo discovery of differentially reactive regions for transcriptome-wide data.

```
dStructome(rl, reps_A, reps_B, batches = F, min_length = 11,
   check_signal_strength = T, check_nucs = T, check_quality = T,
   quality = "auto", evidence = 0, signal_strength = 0.1,
   within_combs = NULL, between_combs = NULL, ind_regions = T, gap = 1,
   processes = "auto", method = "denovo")
```

getCombs 5

Arguments

rl	List of dataframes of reactivities for each sample.		
reps_A	Number of replicates of group A.		
reps_B	Number of replicates of group B.		
batches	Logical suggesting if replicates of group A and B were performed in batches and are labelled accordingly.		
min_length	Minimum length of constructed regions.		
check_signal_s	trength		
_	Logical, if TRUE, constructed regions must have a minimum signal strength.		
check_nucs	Logical, if TRUE, constructed regions must have a minimum number of nucleotides participating in Wilcoxon signed rank test.		
check_quality	Logical, if TRUE, check constructed regions for quality.		
quality	Worst allowed quality for a region to be tested.		
evidence	Minimum evidence of between-group variation for a region to be tested.		
signal_strengt	h		
	Threshold for minimum signal strength.		
within_combs	Data.frame with each column containing groupings of replicates of groups A or B, which will be used to assess within-group variation.		
between_combs	Dataframe with each column containing groupings of replicates of groups A and B, which will be used to assess between-group variation.		
ind_regions	Logical, if TRUE, test each region found in the transcript separately.		

Integer. Join regions if they are separated by these many nucleotides.

Character specifying either guided or de novo discovery approach.

Value

gap

processes

method

Constructs regions, reports p-values and FDR for them.

Number of parallel processes to use.

getCombs	Identifies subgroupings of replicates for assessing within-group and between-group variation.

Description

Identifies subgroupings of replicates for assessing within-group and between-group variation.

```
getCombs(reps_A, reps_B, batches = F, between_combs = NULL,
   within_combs = NULL)
```

6 getRegions

Arguments

reps_A	Number of replicates of group A.
reps_B	Number of replicates of group B.
batches	Logical suggesting if replicates of group A and B were performed in batches and are labelled accordingly.
between_combs	Dataframe with each column containing groupings of replicates of groups A and B, which will be used to assess between-group variation.
within_combs	Data.frame with each column containing groupings of replicates of groups A or

B, which will be used to assess within-group variation.

Value

List of two dataframes, containing groupings for within-group and between-group variation.

getContigRegions	Performs guided discovery of differentially reactive regions.	

Description

Performs guided discovery of differentially reactive regions.

Usage

```
getContigRegions(x, gap = 0)
```

Arguments

x A vector of integers.gap Allowed gap to merge regions.

Value

Dataframe storing start and stop sites of continguous regions.

getRegions Constructs potential differentially reactive regions.	ructs potential differentially reactive regions.
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Description

Constructs potential differentially reactive regions.

```
getRegions(d_within, d_spec, rdf, min_length = 11,
  check_signal_strength = T, check_nucs = T, check_quality = T,
  quality = 0.5, evidence = 0, signal_strength = 0.1)
```

normalizer 7

Arguments

d_within Nucleotide-wise d score for within-group variation.d_spec Nucleotide-wise d score for between-group variation.

rdf Dataframe of reactivities for each sample.
min_length Minimum length of constructed regions.

 $check_signal_strength$

Logical, if TRUE, constructed regions must have a minimum signal strength.

check_nucs Logical, if TRUE, constructed regions must have a minimum number of nu-

cleotides participating in Wilcoxon signed rank test.

check_quality Logical, if TRUE, check constructed regions for quality.

quality Worst allowed quality for a region to be tested.

evidence Minimum evidence of between-group variation for a region to be tested.

signal_strength

Threshold for minimum signal strength.

normalizer Returns normalizer for reactivity vector.

Description

Assesses normalization factor for raw reactivities using 2-8 % method.

Usage

```
normalizer(raw.estimates)
```

Arguments

raw.estimates A vector of raw reactivities.

Value

The normalization factor.

Description

Plots differentially reactive regions.

```
plot_dStructurome(rl, diff_regions, outfile, fdr = 0.05, ylim = c(-0.05, 3))
```

8 two.eight.normalize

Arguments

rl List of dataframes of reactivities for each sample.

diff_regions Dataframe of regions with significance of differentially reactivity.

outfile The name for pdf file which will be saved.

fdr FDR threshold for plotted regions.

ylim Y-axis limits for plots.

Value

Saves a PDF for all differentially reactive regions. Returns NULL.

two.eight.normalize Normalizes reactivity vector.

Description

Normalizes raw reactivities using 2-8 % method.

Usage

```
two.eight.normalize(raw.estimates)
```

Arguments

raw.estimates A vector of raw reactivities.

Value

A vector of normalized reactivities.

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