Package ‘dStruct’

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Type Package

Title Identifying differentially reactive regions from RNA structurome profiling data

Version 1.10.0

Depends R (>= 4.1)

Description dStruct identifies differentially reactive regions from RNA structurome profiling data. dStruct is compatible with a broad range of structurome profiling technologies, e.g., SHAPE-MaP, DMS-MaPseq, Structure-Seq, SHAPE-Seq, etc. See Choudhary et al., Genome Biology, 2019 for the underlying method.

Imports zoo, ggplot2, purrr, reshape2, parallel, IRanges, S4Vectors, rlang, grDevices, stats, utils

License GPL (>= 2)

biocViews StatisticalMethod, StructuralPrediction, Sequencing, Software

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BugReports https://github.com/dataMaster-Kris/dStruct/issues

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**calcDis**

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#### calcDis

*Calculates d score.*

**Description**

*d* score of a nucleotide is a measure of dissimilarity of its normalized reactivity scores. Consider a transcript and its reactivity profiles from a group of samples. Then, the *d* score of a nucleotide is \((2/\pi)\) times the arc-tangent of the ratio of the sample standard deviation of its reactivities to their mean.

**Usage**

```r
calcDis(x)
```

**Arguments**

- **x**  
  A numeric vector or matrix.

**Value**

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

**Author(s)**

Krishna Choudhary
**dCombs**

**References**


**Examples**

```r
# Lower standard deviation of reactivities results in lower d-score.
calcDis(rnorm(10, 1, 0.2))
calcDis(rnorm(10, 1, 0.6))
```

---

**dCombs**

**Assesses within-group or between-group variation.**

**Description**

Given the reactivity profiles for a transcript from multiple samples, and a list of sample identifiers, this function computes the dissimilarity of reactivity scores between the specified samples. These are returned as a sequence of nucleotide-wise $d$ scores.

**Usage**

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

**Arguments**

- `rdf` : Data.frame of reactivities for each sample.
- `combs` : Data.frame with each column containing groupings of samples.

**Value**

Nucleotide-wise $d$ scores.

**Author(s)**

Krishna Choudhary

**References**

Examples

```r
# Example of a data frame with reactivities.
reacs <- data.frame(matrix(runif(30, 0, 10), 10, 3))

# The columns of data frame with must indicate sample grouping and id.
colnames(reacs) <- c("A1", "A2", "B1")

# Get nucleotide-wise dissimilarity scores for a set of samples.
dCombs(rdf = reacs, combs = data.frame(c("A1", "B1")))
```

---

dStruct

**Performs de novo discovery of differentially reactive regions.**

Description

This function takes reactivity profiles for samples of two groups as input and identifies differentially reactive regions in three steps (see Choudhary et al., *Genome Biology*, 2019 for details). First, it regroups the samples into homogeneous and heterogeneous sub-groups, which are used to compute the within-group and between-group nucleotide-wise $d$ scores. Second, smoothed between- and within-group $d$ score profiles are compared to construct candidate differential regions. Finally, unsmoothed between- and within-group $d$ scores are compared using the Wilcoxon signed-rank test. The resulting p-values quantify the significance of difference in reactivity patterns between the two input groups.

Usage

```r
dStruct(
  rdf,
  reps_A,
  reps_B,
  batches = FALSE,
  min_length = 11,
  check_signal_strength = TRUE,
  check_nucs = TRUE,
  check_quality = TRUE,
  quality = "auto",
  evidence = 0,
  signal_strength = 0.1,
  within_combs = NULL,
  between_combs = NULL,
  ind_regions = TRUE,
  gap = 1,
  get_FDR = TRUE,
  proximity_assisted = FALSE,
  proximity = 10,
  proximity_defined_length = 30
)
```
**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. If TRUE, a heterogeneous/homogeneous subset may not have multiple samples from the same batch.
- `min_length`: Minimum length of constructed regions.
- `check_signal_strength`: Logical, if TRUE, construction of regions must be based on nucleotides that have a minimum absolute value of reactivity.
- `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 increase in variation from within-group comparisons to between-group comparisons for a region to be tested.
- `signal_strength`: Threshold for minimum signal strength.
- `within_combs`: Dataframe 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.
- `proximity_assisted`: Logical, if TRUE, proximally located regions are tested together.
- `proximity`: Maximum distance between constructed regions for them to be considered proximal.
- `proximityDefinedLength`: If performing a "proximity-assisted" test, minimum end-to-end length of a region to be tested.

**Value**

Constructs regions, reports p-value and median difference of between-group and within-group d-scores for each region, and FDR for them.

**Author(s)**

Krishna Choudhary
References


Examples

```r
# Load data from Lai et al., 2019
data(lai2019)

# Run dStruct in de novo discovery mode for a transcript with id YAL042W.
dStruct(rdf = lai2019[['YAL042W']], reps_A = 3, reps_B = 2,
batches = TRUE, min_length = 21,
between_combs = data.frame(c("A3", "B1", "B2")),
within_combs = data.frame(c("A1", "A2", "A3")),
ind_regions = TRUE)
```

**dStructGuided**

Performs guided discovery of differentially reactive regions.

**Description**

This function takes as input reactivity profiles for a transcript region from samples of two groups. First, it regroups the samples into homogeneous and heterogeneous sub-groups, which are used to compute the within-group and between-group nucleotide-wise $d$ scores. If the region meets the quality criteria, the between- and within-group $d$ scores are compared using the Wilcoxon signed-rank test. The resulting p-values quantify the significance of difference in reactivity patterns between the two input groups.

**Usage**

```r
dStructGuided(
  rdf, 
  reps_A, 
  reps_B, 
  batches = FALSE, 
  within_combs = NULL, 
  between_combs = NULL, 
  check_quality = TRUE, 
  quality = "auto", 
  evidence = 0
)
```

**Arguments**

- `rdf`: Dataframe of reactivities for each sample. Each column must be labelled as A1, A2, ..., B1, B2, ...
- `reps_A`: Number of replicates of group A.
dStructome

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. If TRUE, a heterogeneous/homogeneous subset may not have multiple samples from the same batch.
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 increase in variation from within-group comparisons to between-group comparisons for a region to be tested.

Value

p-value for the tested region (estimated using one-sided Wilcoxon signed rank test) and the median of nucleotide-wise difference of between-group and within-group d-scores.

Author(s)

Krishna Choudhary

References


Examples

#Load Wan et al., 2014 data
data(wan2014)

#Run dStruct in the guided mode on first region in wan2014.
dStructGuided(wan2014[[1]], reps_A = 2, reps_B = 1)

---

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

Description

This function provides a convenient way to call the dStruct or dStructGuided functions for multiple transcripts simultaneously. By default, the transcripts are processed in using multiple parallel processes if available.
Usage

dStructome(
  rl,
  reps_A,
  reps_B,
  batches = FALSE,
  min_length = 11,
  check_signal_strength = TRUE,
  check_nucs = TRUE,
  check_quality = TRUE,
  quality = "auto",
  evidence = 0,
  signal_strength = 0.1,
  within_combs = NULL,
  between_combs = NULL,
  ind_regions = TRUE,
  gap = 1,
  processes = "auto",
  method = "denovo",
  proximity_assisted = FALSE,
  proximity = 10,
  proximity_defined_length = 30
)

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. If TRUE, a heterogeneous/homogeneous subset may not have multiple samples from the same batch.
min_length Minimum length of constructed regions.
check_signal_strength Logical, if TRUE, construction of regions must be based on nucleotides that have a minimum absolute value of reactivity.
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 increase in variation from within-group comparisons to between-group comparisons for a region to be tested.
signal_strength 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.

processes  Number of parallel processes to use.

method  Character specifying either guided or de novo discovery approach.

proximity_assisted  Logical, if TRUE, proximally located regions are tested together.

proximity  Maximum distance between constructed regions for them to be considered proximal.

proximity_defined_length  If performing a “proximity-assisted” test, minimum end-to-end length of a region to be tested.

Value

Constructs regions, reports p-value and median difference of between-group and within-group d-scores for each region, and FDR for them.

Author(s)

Krishna Choudhary

References


Examples

#Load data from Lai et al., 2019
data(lai2019)

dStructome(lai2019, 3, 2, batches= TRUE, min_length = 21,
between_combs = data.frame(c("A3", "B1", "B2")),
within_combs = data.frame(c("A1", "A2", "A3")),
ind_regions = TRUE, processes = 1)

#Load data from Wan et al., 2014
data(wan2014)

dStructome(wan2014, reps_A = 2, reps_B = 1, method = "guided", processes = 1)
getCombs

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

Description

Regroup all the samples of A and B groups into homogoneous and heterogeneous sub-groups. Each homogenous sub-group contains replicates of either group A only or group B only. Each heterogeneous sub-group has a mix of samples from both the groups A and B.

Usage

getCombs(
  reps_A,
  reps_B,
  batches = FALSE,
  between_combs = NULL,
  within_combs = NULL
)

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. If TRUE, a heterogeneous/homogeneous subset may not have multiple samples from the same batch.
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 Dataframe 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.

Author(s)

Krishna Choudhary

References

getContigRegions

Examples

#Get heterogeneous and homogeneous set combinations of samples when there are 2 samples of group A and 1 of group B.
getCombs(2, 1)

getContigRegions

Identifies contiguous regions from a list of nucleotide indices.

Description

Given a sequence of nucleotide indices, this function returns integer ranges covered by the indices. There is an option to merge ranges if they are separated by less than a user-specified distance.

Usage

getContigRegions(x, gap = 0)

Arguments

x

A vector of integers.

gap

Include gaps in the ranges if they are shorter than or equal to this length.

Value

IRanges object storing start and end sites of contiguous regions.

Author(s)

Krishna Choudhary

Examples

#Convert an integer vector of nucleotide positions to an IRanges object containing the coordinates of contiguous regions.
nucleotide_positions <- c(1, 3, 2, 8, 4:7, 11:20)
getContigRegions(nucleotide_positions)

#Merge regions if their end points are within 3 nt of each other.
getContigRegions(nucleotide_positions, gap = 3)
getRegions

Constructs potential differentially reactive regions.

Description

This function takes between- and within-group \( d \) scores for a transcript as input and identifies regions where the former is generally larger. Regions that pass minimum quality and minimum signal criteria are returned.

Usage

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

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, construction of regions must be based on nucleotides that have a minimum absolute value of reactivity.
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 increase in variation from within-group comparisons to between-group comparisons for a region to be tested.
signal_strength  Threshold for minimum signal strength.

Value

Integer vector of nucleotides that constitute potential differentially reactive regions.
Author(s)

Krishna Choudhary

References


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lai2019  
Saccharomyces cerevisiae Structure-seq data

Description

Data from a Structure-seq assay of five samples of *S. cerevisiae*, three of which were wild-type samples and two mutant samples. The data was pre-processed to obtain DMS reactivities as described by Lai et al. (2019).

Usage

data("lai2019")

Format

An object of class "list".

Source

Raw data from Lai et al., 2019 in processed form.

References


Examples

data("lai2019")
normalizer

Returns normalizer for reactivity vector.

Description
Assesses normalization factor for raw reactivities using the 2-8% method. Given a reactivity profile, first, remove 2% of the nucleotides with the highest reactivities. Then, the normalization factor is the mean of reactivities of the 8% of the nucleotides with the next highest reactivities.

Usage

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

Arguments

- `raw.estimates` A vector of raw reactivities.

Value

The normalization factor.

Author(s)

Krishna Choudhary

References


Examples

```r
normalizer(c(NA, rnorm(20, 0.5, 0.3), NA, -999))
```
plotDStructurome

Plots differentially reactive regions.

Description

Given the table of results from dStruct or dStructGuided and the corresponding lists with reactivity scores for all transcripts, this function saves a PDF file with detailed visualizations of reactivities for all differential regions.

Usage

plotDStructurome(
  rl,
  diff_regions,
  outfile,
  fdr = 0.05,
  ylim = c(-0.05, 3),
  del_d_cutoff = 0.01
)

Arguments

rl List of dataframes of reactivities for each sample.

diff_regions Output from dStruct or dStructGuided containing coordinates 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.

del_d_cutoff Minimum effect size for plotted regions specified in terms of median difference of the between-group and within-group d-scores.

Value

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

Author(s)

Krishna Choudhary

References

Examples

# Load data from Lai et al., 2019
data(lai2019)

# Run dStruct in de novo discovery mode for all the transcripts in this data in one step.
res <- dStructome(lai2019, 3, 2, batches = TRUE, min_length = 21,
between_combs = data.frame(c("A3", "B1", "B2")),
within_combs = data.frame(c("A1", "A2", "A3")),
ind_regions = TRUE, processes = 1)

# Plot the significant results and save to a PDF file.
plotDStructurome(rl = lai2019,
diff_regions = res,
outfile = "significantly_differential_regions",
fdr = 0.05,
ylim = c(-0.05, 3))

twoEightNormalize

Normalizes reactivity vector.

Description

Given a reactivity profile, first, remove 2% of the nucleotides with the highest reactivities. Then, the normalization factor is the mean of reactivities of the 8% of the nucleotides with the next highest reactivities. The raw reactivities are divided by the normalization factor to get normalized reactivities. This is called as 2-8% normalization and has been a common way to normalize data from RNA structurome profiling technologies such as SHAPE-Seq, Structure-Seq, etc. (see Low and Weeks, 2010, Sloma et al., 2015, and Choudhary et al., 2017).

Usage

twoEightNormalize(raw.estimates)

Arguments

raw.estimates A vector of raw reactivities.

Value

A vector of normalized reactivities.

Author(s)

Krishna Choudhary
References


Examples

twoEightNormalize(c(NA, rnorm(20, 0.5, 0.3), NA, -999))

---

**wan2014**  
Homo sapiens *PARS data*

---

**Description**

Data from a PARS assay of a family trio of mother, father, and child. The data was pre-processed to obtain PARS scores as described in Choudhary et al. (2019).

**Usage**

data(wan2014)

**Format**

An object of class "list".

**Source**

Counts data from Wan et al., 2014 in processed form.

**References**


**Examples**

data(wan2014)
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