# Package 'dStruct'

September 19, 2024

Type Package

**Title** Identifying differentially reactive regions from RNA structurome profiling data

Version 1.11.0

**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

URL https://github.com/dataMaster-Kris/dStruct

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

**Encoding** UTF-8

LazyData true

RoxygenNote 7.1.1

Suggests BiocStyle, knitr, rmarkdown, tidyverse, testthat (>= 3.0.0)

VignetteBuilder knitr

#### **Config/testthat/edition** 3

git\_url https://git.bioconductor.org/packages/dStruct

git\_branch devel

git\_last\_commit d23992a

git\_last\_commit\_date 2024-04-30

**Repository** Bioconductor 3.20

Date/Publication 2024-09-18

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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

calcDis(x)

# Arguments

Х

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

# References

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

Choudhary K, Shih NP, Deng F, Ledda M, Li B, Aviran S. Metrics for rapid quality control in RNA structure probing experiments. *Bioinformatics*. 2016; 32(23):3575–3583.

#### dCombs

#### Examples

```
#Lower standard deviation of reactivites 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

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

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

#### Examples

```
#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

# 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 heteregenous 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

```
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.   |  |

#### dStruct

| 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_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.  |  |
| proximity_assisted       |  |  |
|                          | Logical, if TRUE, proximally located regions are tested together.  |  |
| proximity                | Maximum distance between constructed regions for them to be considered prox-<br>imal.  |  |
| proximity_defined_length |  |  |
|                          | If performing a "proximity-assisted" test, minimum end-to-end length of a re-<br>gion 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

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

# Examples

```
#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
```

# 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 heteregenous 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

```
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.   |
| 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.

#### dStructome

#### Author(s)

Krishna Choudhary

#### References

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

# 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_st | -  |
|                 | 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_assis | ted  |
|                 | Logical, if TRUE, proximally located regions are tested together.  |
| proximity       | Maximum distance between constructed regions for them to be considered proximal.   |
| proximity_defin | -  |
|                 | 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

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

#### getCombs

## 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.
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)
#Run dStruct in guide discovery mode for all the transcript regions in this data in one step.
dStructome(wan2014, reps_A = 2, reps_B = 1, method = "guided", processes = 1)
```

*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  | 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.

#### Author(s)

Krishna Choudhary

#### References

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

# Examples

#Get heterogeneous and homogeneous set combinations of samples when there are 2 samples of group A and 1 of group 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

| х   | 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 continguous regions.

#### Author(s)

Krishna Choudhary

# Examples

#Convert an integer vector of nucleotide positions to an IRanges object containing the coordinates of contiguous 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)

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getRegions

# 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_st | trength   |
|                 | 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

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

lai2019Saccharomyces 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

Lai et al. (2019) Genetics, Vol. 212, 153–174 (Genetics)

#### 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

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

# Arguments

raw.estimates A vector of raw reactivities.

#### plotDStructurome

# Value

The normalization factor.

# Author(s)

Krishna Choudhary

# References

Low JT, Weeks KM. SHAPE-directed RNA secondary structure prediction. Methods. 2010; 52(2):150-8.

Sloma MF, Mathews DH, Chen SJ, Burke-Aguero DH. Chapter four – improving RNA secondary structure prediction with structure mapping data. In: *Methods in Enzymology*, vol. 553. Cambridge: Academic Press: 2015. p. 91–114.

Choudhary K, Deng F, Aviran S. Comparative and integrative analysis of RNA structural profiling data: current practices and emerging questions. *Quant Biol.* 2017; 5(1):3–24.

#### Examples

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. |

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

#### Author(s)

Krishna Choudhary

#### References

Choudhary, K., Lai, Y. H., Tran, E. J., & Aviran, S. (2019). dStruct: identifying differentially reactive regions from RNA structurome profiling data. *Genome biology*, 20(1), 1-26.

#### 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 = 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.

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#### wan2014

#### Author(s)

Krishna Choudhary

#### References

Low JT, Weeks KM. SHAPE-directed RNA secondary structure prediction. Methods. 2010; 52(2):150-8.

Sloma MF, Mathews DH, Chen SJ, Burke-Aguero DH. Chapter four – improving RNA secondary structure prediction with structure mapping data. In: *Methods in Enzymology*, vol. 553. Cambridge: Academic Press: 2015. p. 91–114.

Choudhary K, Deng F, Aviran S. Comparative and integrative analysis of RNA structural profiling data: current practices and emerging questions. *Quant Biol.* 2017; 5(1):3–24.

#### 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

Wan et al., Nature, 505, 706–709 (2014) (Nature)

## Examples

data(wan2014)

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