# Package 'nearBynding'

November 12, 2025

Type Package

Title Discern RNA structure proximal to protein binding

Version 1.20.0

Description Provides a pipeline to discern RNA structure at and proximal to the site of protein binding within regions of the transcriptome defined by the user. CLIP protein-binding data can be input as either aligned BAM or peak-called bedGraph files. RNA structure can either be predicted internally from sequence or users have the option to input their own RNA structure data. RNA structure binding profiles can be visually and quantitatively compared across multiple formats.

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**biocViews** Visualization, MotifDiscovery, DataRepresentation, StructuralPrediction, Clustering, MultipleComparison

**Encoding UTF-8** 

LazyData true

**Depends** R (>= 4.0)

Imports R.utils, matrixStats, plyranges, transport, Rsamtools, S4Vectors, grDevices, graphics, rtracklayer, dplyr, Seqinfo, methods, GenomicRanges, utils, stats, magrittr, TxDb.Hsapiens.UCSC.hg19.knownGene, TxDb.Hsapiens.UCSC.hg38.knownGene, ggplot2, gplots, BiocGenerics, rlang

Suggests knitr, rmarkdown

**SystemRequirements** bedtools (>= 2.28.0), Stereogene (>= v2.22), CapR (>= 1.1.1)

VignetteBuilder knitr

Collate 'assessGrouping.R' 'bindingContextDistance.R' 'bindingContextDistanceCapR.R' 'CleanBAMtoBG.R' 'CleanBEDtoBG.R' 'ExtractTranscriptomeSequence.R' 'GenomeMappingToChainFile.R' 'get\_outfiles.R' 'liftOverToExomicBG.R' 'processCapRout.R' 'runCapR.R' 'runStereogene.R' 'runStereogeneOnCapR.R' 'visualizeCapRStereogene.R' 'visualizeStereogene.R' 'write\_config.R' 'write\_fasta.R' 'getChainChrSize.R' 'utilities.R' 'symmetryCapR.R' 'symmetryContext.R'

RoxygenNote 7.1.1

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Author Veronica Busa [cre]
Maintainer Veronica Busa < vbusa 1@ihmi .edu>

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# Description

Assess grouping of samples assigned to the same category relative to random.

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#### Usage

```
assessGrouping(
  distances,
  annotations,
  measurement = "mean",
  output = "KS.pvalue",
  ctrl_iterations = 10000
)
```

#### **Arguments**

distances Data frame object with at least three columns where the first three columns are

sample 1 name, sample 2 name, and the distance between them.

annotations Data frame object with at least two columns where the first two columns are

sample name and the category of the sample for grouping. Sample names must

match sample 1 and sample 2 names in distances data frame.

measurement The measurement for comparison between cases and controls and statistical

analysis ("mean", "max", or "min). Default "mean"

output A string denoting what information will be returned: either a list of test and con-

trol measurement distances ("measurements"), the p-value of the Kolmogorov-Smirnov test comparing test and control distributions ("KS.pvalue"), or a ggplot object plotting the test and control distributions ("plot"). Default "KS.pvalue"

ctrl\_iterations

The number of iterations to test for the control distribution; an integer. Default

10000.

#### Value

bindingContextDistance

bindingContextDistance

# **Description**

Calculate the Wasserstein distance between two replicates' or two proteins' binding contexts for CapR-generated RNA contexts.

#### Usage

```
bindingContextDistance(
    dir_stereogene_output = ".",
    RNA_context,
    protein_file,
    protein_file_input = NULL,
    dir_stereogene_output_2 = NULL,
    RNA_context_2 = NULL,
    protein_file_2 = NULL,
    protein_file_input_2 = NULL,
    range = c(-200, 200)
)
```

#### **Arguments**

dir\_stereogene\_output

Directory of Stereogene output for first protein. Default current directory.

RNA\_context Name of the RNA context file input to Stereogene. File names must exclude extensions such as ".bedGraph". Requred

protein\_file A vector of at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Optional.

dir\_stereogene\_output\_2

Directory of Stereogene output for second protein. Default dir\_stereogene\_output.

RNA\_context\_2 Name of the RNA context file input to Stereogene. File names must exclude extensions such as ".bedGraph". Default same as RNA\_context.

protein\_file\_2 Similar to protein\_file. A second vector of at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates.

Default same as protein\_file

protein\_file\_input\_2

Similar to protein\_file\_input. A second protein file name of background input to be subtracted from protein\_file\_2 signal. File name must exclude extension. Only one input file is permitted. Optional.

range

A vector of two integers denoting the range upstream and downstream of the center of protein binding to consider in the comparison. Ranges that are too small miss the holistic binding context, while large ranges amplify distal noise in the binding data. Cannot exceed wSize/2 from write\_config. Default c(-200, 200)

#### Value

Wasserstein distance between the two protein file sets provided for the RNA structure context specified, minus the input binding signal if applicable

#### Note

Either RNA\_context\_2 or protein\_file\_2 must be input. Otherwise, the distance would be calculated between the same file and equal 0.

Wasserstein distance calculations are reciprocal, so it does not matter which protein is first or second so long as replicates and input files correspond to one another.

# **Examples**

bindingContextDistanceCapR

bindingContextDistanceCapR

# **Description**

Calculate the Wasserstein distance between two replicates' or two proteins' binding contexts.

#### Usage

```
bindingContextDistanceCapR(
   dir_stereogene_output = ".",
   CapR_prefix = "",
   protein_file,
   protein_file_input = NULL,
   dir_stereogene_output_2 = NULL,
   CapR_prefix_2 = "",
   protein_file_2,
   protein_file_input_2 = NULL,
```

```
context = "all",
  range = c(-200, 200)
)
```

# **Arguments**

dir\_stereogene\_output

Directory of Stereogene output for first protein. Default current directory.

CapR\_prefix The prefix common to CapR output files of protein\_file, if applicable. Equivalent to output\_prefix from runStereogeneOnCapR. Default ""

protein\_file A vector of strings with at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Optional.

dir\_stereogene\_output\_2

Directory of Stereogene output for second protein. Default current directory.

CapR\_prefix\_2 The prefix common to CapR output files of protein\_file\_2, if applicable.Equivalent to output\_prefix from r unStereogeneOnCapR. Default ""

protein\_file\_2 Similar to protein\_file. A second vector of at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates. Required.

protein\_file\_input\_2

context

range

Similar to protein\_file\_input. A second protein file name of background input to be subtracted from protein\_file\_2 signal. File name must exclude extension. Only one input file is permitted. Optional.

The RNA structure context being compared for the two protein file sets. Acceptable contexts include "all", which sums the distance of all six contexts, or any of the contexts individually ("bulge", "hairpin", "stem", "exterior", "multibranch", or "internal"). Default "all"

A vector of two integers denoting the range upstream and downstream of the center of protein binding to consider in the comparison. Ranges that are too small miss the holistic binding context, while large ranges amplify distal noise in the binding data. Cannot exceed wSize/2 from write\_config. Default c(-200,

200)

#### Value

Wasserstein distance between the two protein file sets provided for the RNA structure context specified, minus the input binding signal if applicable

# Note

Wasserstein distance calculations are reciprocal, so it does not matter which protein is first or second so long as replicates and input files correspond to one another.

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#### **Examples**

CleanBAMtoBG

CleanBAMtoBG

# **Description**

Writes a script to convert a BAM file to a clean bedGraph file.

#### Usage

```
CleanBAMtoBG(in_bam, out_bedGraph = NA, unwanted_chromosomes = NULL)
```

# **Arguments**

in\_bam Name of sorted BAM file to be converted to a bedGraph file. Required.

prefix.

 $unwanted\_chromosomes$ 

A vector of unwanted chromosomes that are present in the BAM file.

# Value

deposits bedGraph from BAM in same directory

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CleanBEDtoBG

Clean BED to BG

# **Description**

Writes a script to convert a BED file to a clean bedGraph file.

# Usage

```
CleanBEDtoBG(
  in_bed,
  out_bedGraph = NA,
  unwanted_chromosomes = NULL,
  alignment = "hg19"
)
```

# **Arguments**

in\_bed Name of sorted BAM file to be converted to a bedGraph file. Required.

out\_bedGraph 
Name of bedGraph output file, including full directory path; a string. Default

in\_bam prefix.

unwanted\_chromosomes

A vector of unwanted chromosomes that are present in the BAM file.

alignment The human genome alignment used, either "hg19" or "hg38". Default "hg19"

# Value

deposits bedGraph from BED in same directory

```
bam <- system.file("extdata/chr4and5.bam", package="nearBynding")
out_bed <- "bamto.bed"
## convert BAM to BED
if(suppressWarnings(system2("bedtools", "--version",
stdout = NULL, stderr = NULL)) == 0){
    system2("bedtools", paste0("bamtobed -i ", bam, " > ", out_bed))
}
CleanBEDtoBG(in_bed = out_bed,
    alignment = "hg38")
```

ExtractTranscriptomeSequence

 ${\it ExtractTranscriptomeSequence}$ 

# **Description**

Writes a FASTA file of transcript sequences from a list of transcripts.

# Usage

```
ExtractTranscriptomeSequence(
  transcript_list,
  ref_genome,
  genome_gtf,
  RNA_fragment = "exon",
  exome_prefix = "exome"
)
```

#### **Arguments**

transcript\_list

A vector of transcript names that represent the most expressed isoform of their

respective genes and correspond to GTF annotation names. Required

ref\_genome The name of the reference genome FASTA from which exome sequences will

be derived; a string. Required

genome\_gtf The name of the GTF/GFF file that contains all exome annotations; a string.

Coordinates must match the file input for the ref\_genome parameter. Required

RNA\_fragment A string of RNA component of interest. Options depend on the gtf file but often

include "gene", "transcript", "exon", "CDS", "five\_prime\_utr", and/or "three\_prime\_utr".

Default "exon" for the whole exome.

exome\_prefix A string to add to the prefix for all output files. Default "exome"

# Value

writes FASTA file of transcriptome sequences into directory

### Note

transcript\_list, genome\_gtf, and RNA\_fragment arguments should be the same as GenomeMappingToChainFile function arguments

```
RNA_fragment = "three_prime_utr",
exome_prefix = "chr4and5_3UTR")
```

GenomeMappingToChainFile

Genome Mapping To Chain File

### **Description**

Writes a chain file mapped from a genome annotation file.

# Usage

```
GenomeMappingToChainFile(
  genome_gtf,
  out_chain_name,
  RNA_fragment = "exon",
  transcript_list,
  chrom_suffix = "exome",
  verbose = FALSE,
  alignment = "hg19",
  check_overwrite = FALSE
)
```

#### **Arguments**

genome\_gtf The name of the GTF/GFF file that will be converted to an exome chain file.

Required

out\_chain\_name The name of the chain file to be created. Required

RNA\_fragment RNA component of interest. Options depend on the gtf file but often include

"gene", "transcript", "exon", "CDS", "five\_prime\_utr", and/or "three\_prime\_utr".

Default "exon" for the whole exome.

transcript\_list

A vector of transcript names that represent the most expressed isoform of their respective genes and correspond to gtf annotation names. Isoforms cannot over-

lap. Required

chrom\_suffix 
The suffix to be appended to all chromosome names created in the chain file.

Default "exome"

verbose Output updates while the function is running. Default FALSE

alignment The human genome alignment used, either "hg19" or "hg38". Default "hg19"

check\_overwrite

Check for file wth the same out\_chain\_name before writing new file. Default

FALSE.

# Value

writes a chain file into directory

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#### **Examples**

getChainChrSize

getChainChrSize

#### **Description**

Output a table of mapped chromosome names and lengths from a chain file.

# Usage

```
getChainChrSize(chain, out_chr)
```

# **Arguments**

chain The name of the chain file for which chromosome sizes should be determined

and output; a string. Required.

out\_chr Name of the chromosome names and lengths table file; a string. Required.

# Value

writes a two-column tab-delineated file without a header containing chromosome names and lengths for a given chain file

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# **Description**

Copy files necessary to complete the vignette onto the local machine in cases where Stereogene, CapR, or bedtools are not available.

# Usage

```
get_outfiles(dir = ".")
```

# **Arguments**

dir

Directory into which files ought to be stored. Default current work directory.

#### Value

deposits six \*.dist StereoGene output files into the selected directory

# **Examples**

```
## pull example StereoGene output files
get_outfiles()
```

liftOverToExomicBG

liftOverToExomicBG

# Description

Lifts features such as CLIP-seq reads or RNA structure annotations from genome to transcriptome.

# Usage

```
liftOverToExomicBG(input, chain, chrom_size, output_bg, format = "bedGraph")
```

# Arguments

input	A single input file name or a vector of input file names in the format of c(forward_reads, reverse_reads) for strand-separated alignments. Files must be BED or bedGraph format. Required
chain	The name of the chain file to be used for liftOver. Format should be like chain files derived from getChainChrSize function. Required
chrom_size	Name of chromosome size file. File must be in two-column format without a header where first column is chromosome name and second column is chromosome length, as from liftOverToExomicBG. Required.
output_bg	The name of the lifted-over output bedGraph file. Required.
format	File type of input file(s). Recommended "BED" or "bedGraph". Default "bed-Graph"

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#### Value

writes lifted-over bedGraph file

# **Examples**

```
## first, get chain file
load(system.file("extdata/transcript_list.Rda", package="nearBynding"))
gtf<-system.file("extdata/Homo_sapiens.GRCh38.chr4&5.gtf",</pre>
                package="nearBynding")
GenomeMappingToChainFile(genome_gtf = gtf,
                        out_chain_name = "test.chain",
                        RNA_fragment = "three_prime_utr",
                         transcript_list = transcript_list,
                        alignment = "hg38")
## and chain file chromosome sizes
getChainChrSize(chain = "test.chain"
               out_chr = "chr4and5_3UTR.size")
## get bedGraph file
chr4and5_sorted.bedGraph<-system.file("extdata/chr4and5_sorted.bedGraph",</pre>
                                      package="nearBynding")
liftOverToExomicBG(input = chr4and5_sorted.bedGraph,
                  chain = "test.chain",
                  chrom_size = "chr4and5_3UTR.size",
                  output_bg = "chr4and5_liftOver.bedGraph")
```

nearBynding

Discern RNA structure proximal to protein binding

# **Description**

nearBynding is a package designed to discern annotated RNA structures at and proximal to the site of protein binding. It allows users to annotate RNA structure contexts via CapR or input their own annotations in BED/bedGraph format and it accommodates protein binding information from CLIP-seq experiments as either aligned CLIP-seq reads or peak-called intervals.

# **Details**

Package: nearBynding Type: Package

Title: nearBynding package

Version: 1.3.3

Date: June 1, 2021' License: Artistic-2.0

LazyLoad: yes

URL: http://github.com/vbusa1/nearBynding

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#### Author(s)

Veronica Busa <vbusa1@jhmi.edu>

#### References

StereoGene: Stavrovskaya, Elena D., Tejasvi Niranjan, Elana J. Fertig, Sarah J. Wheelan, Alexander V. Favorov, and An CapR: Tsukasa Fukunaga, Haruka Ozaki, Goro Terai, Kiyoshi Asai, Wataru Iwasaki, and Hisanori Kiryu. "CapR:

#### See Also

See the nearBynding package vignette.

```
## Not run:
library(nearBynding)
library(Rsamtools)
# get transcript list
load(system.file("extdata/transcript_list.Rda", package="nearBynding"))
# get GTF file
gtf<-system.file("extdata/Homo_sapiens.GRCh38.chr4&5.gtf",</pre>
                package="nearBynding")
# make chain file
GenomeMappingToChainFile(genome_gtf = gtf,
                        out_chain_name = "test.chain",
                        RNA_fragment = "three_prime_utr",
                        transcript_list = transcript_list,
                        alignment = "hg38")
# get size of chromosomes of chain file
getChainChrSize(chain = "test.chain",
                out_chr = "chr4and5_3UTR.size")
# get transcript sequences
ExtractTranscriptomeSequence(transcript_list = transcript_list,
                    ref_genome = "Homo_sapiens.GRCh38.dna.primary_assembly.fa",
                    genome_gtf = gtf,
                    RNA_fragment = "three_prime_utr",
                    exome_prefix = "chr4and5_3UTR")
# run CapR on extracted sequences
runCapR(in_file = "chr4and5_3UTR.fa")
# get BAM file of protein binding
bam <- system.file("extdata/chr4and5.bam", package="nearBynding")</pre>
# sort it and convert to bedGraph format
sorted_bam<-sortBam(bam, "chr4and5_sorted")</pre>
CleanBAMtoBG(in_bam = sorted_bam)
# lift over protein binding and RNA structure to chain
liftOverToExomicBG(input = "chr4and5_sorted.bedGraph",
                    chain = "test.chain",
                    chrom_size = "chr4and5_3UTR.size",
```

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```
output_bg = "chr4and5_liftOver.bedGraph")
processCapRout(CapR_outfile = "chr4and5_3UTR.out",
                chain = "test.chain",
                output_prefix = "chr4and5_3UTR",
                chrom_size = "chr4and5_3UTR.size",
                genome_gtf = gtf,
                RNA_fragment = "three_prime_utr")
# input to StereoGene
runStereogeneOnCapR(protein_file = "chr4and5_liftOver.bedGraph",
                    chrom_size = "chr4and5_3UTR.size",
                    name_config = "chr4and5_3UTR.cfg",
                    input_prefix = "chr4and5_3UTR")
# visualize protein binding context
visualizeCapRStereogene(CapR_prefix = "chr4and5_3UTR",
                        protein_file = "chr4and5_liftOver",
                        heatmap = T,
                        out_file = "all_contexts_heatmap",
                        x_{lim} = c(-500, 500)
## End(Not run)
```

processCapRout

processCapRout

#### **Description**

Creates context-separated bedGraph files of CapR output for genome and transcriptome alignments.

#### Usage

```
processCapRout(
   CapR_outfile,
   output_prefix,
   chrom_size,
   genome_gtf,
   RNA_fragment,
   chain
)
```

# Arguments

CapR\_outfile Name of CapR output file. Required

output\_prefix Prefix string to be appended to all output files. Required.

Chrom\_size Name of chromosome size file. File must be in two-column format without a header where first column is chromosome name and second column is chromosome length, as from getChainChrSize. Required.

genome\_gtf The name of the GTF/GFF file that contains all exome annotations. Required

RNA\_fragment RNA component of interest. Options depend on the gtf file but often include "gene", "transcript", "exon", "CDS", "five\_prime\_utr", and/or "three\_prime\_utr".

Default "exon" for the whole exome.

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chain

The name of the chain file to be used. Format should be like chain files derived from GRangesMappingToChainFile. Required

#### Value

writes bedGraph files of structure signal for each of the six CapR contexts 1) mapped to the genome and 2) lifted-over to the transcriptome

# **Examples**

```
## make chain file
load(system.file("extdata/transcript_list.Rda", package="nearBynding"))
gtf<-system.file("extdata/Homo_sapiens.GRCh38.chr4&5.gtf",</pre>
                package="nearBynding")
GenomeMappingToChainFile(genome_gtf = gtf,
                        out_chain_name = "test.chain",
                        RNA_fragment = "three_prime_utr",
                        transcript_list = transcript_list,
                        alignment = "hg38")
## get chromosome size file
getChainChrSize(chain = "test.chain",
               out_chr = "chr4and5_3UTR.size")
processCapRout(CapR_outfile = system.file("extdata/chr4and5_3UTR.out",
                                          package="nearBynding"),
              chain = "test.chain",
              output_prefix = "chr4and5_3UTR",
              chrom_size = "chr4and5_3UTR.size",
              genome_gtf = gtf,
              RNA_fragment = "three_prime_utr")
```

runCapR

runCapR

# **Description**

Runs CapR

# Usage

```
runCapR(in_file, out_file = NA, max_dist = 100)
```

# Arguments

in_file	An .fa file like from ExtractTranscriptomeSequence that is a list of fasta sequences to be folded. Required
out_file	Name of the CapR output file of nucleotide folding probabilities. Default is in_file prefix.out
max_dist	Integer of maximum distance between folded nucleotides in sequences. Recommeded between 50 and 100, with higher values yielding potentially more accurate results but dramatically increasing run time. Default 100.

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#### Value

```
generates CapR outfile
```

# **Examples**

runStereogene

runStereogene

# Description

Writes a StereoGene script in the working directory

# Usage

# Arguments

track_files	Vector of at least two track or interval file names to be pairwise-correlated by StereoGene. Required.
name_config	Name of corresponding configuration file; a string. Required
pcorProfile	Name of track file name for partial correlation; a string. More information for this can be found in the StereoGene README. Optional
confounder	Confounder file name; a string. More information for this can be found in the StereoGene README. Optional
nShuffle	Permutations used to estimate error. Default 5000.
get_error	Whether to calculate the standard error of background permutations from nShuf- fle. FALSE will save calculation time. Default FALSE

#### Value

generates StereoGene output files in directory

runStereogeneOnCapR runStereogeneOnCapR

# Description

Writes a configuration file and Stereogene script and runs Stereogene for all CapR tracks

# Usage

```
runStereogeneOnCapR(
  dir_CapR_bg = ".",
  input_prefix,
  protein_file,
  output_prefix = input_prefix,
  name_config = "config.cfg",
  chrom_size,
  nShuffle = 100,
  get_error = FALSE,
  ...
)
```

# **Arguments**

dir_CapR_bg	Directory of lifted-over CapR bedGraph files. Default current directory
input_prefix	Prefix string appended to input files; same as input_prefix argument in process-CapRout. Required
protein_file	Name of protein file in bedGraph format. Required
output_prefix	Prefix string to be appended to all output files. Default to be same as input_prefix
name_config	Name of output config file. Default config.cfg
chrom_size	Name of chromosome size file. File must be in two-column format without a header where first column is chromosome name and second column is chromosome length, as from getChainChrSize. Required
	includes all other parameters acceptable to write_config and write_stereogene
nShuffle	Permutations used to estimate error. Default 100.
get_error	Whether to calculate the standard error of background permutations from nShuffle. FALSE will save calculation time. Default FALSE

# Value

generates StereoGene output files, including \*.dist files

symmetryCapR 19

yCapR
)

#### **Description**

Calculate the symmetry of a binding context.

# Usage

```
symmetryCapR(
  dir_stereogene_output = ".",
  CapR_prefix = "",
  protein_file,
  protein_file_input = NULL,
  context = "all",
  range = c(-200, 200)
)
```

#### **Arguments**

dir\_stereogene\_output

Directory of Stereogene output for first protein. Default current directory.

CapR\_prefix

The prefix common to CapR output files of protein\_file, if applicable. Equivalent to output\_prefix from runStereogeneOnCapR. Default ""

protein\_file

A vector of strings with at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Optional.

context

The RNA structure context being interrogated. Acceptable contexts include "all", which sums the distance of all six contexts, or any of the contexts individually ("bulge", "hairpin", "stem", "exterior", "multibranch", or "internal"). Default "all"

range

A vector of two integers denoting the range upstream and downstream of the center of protein binding to consider in the comparison. Ranges that are too small miss the holistic binding context, while large ranges amplify distal noise in the binding data. Cannot exceed wSize/2 from write\_config. Default c(-200, 200)

### Value

Wasserstein distance between the two halves of the binding context, with lower values suggesting greater symmetry.

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#### **Examples**

symmetryContext

symmetryContext

# **Description**

Calculate the symmetry of a binding context.

# Usage

```
symmetryContext(
  dir_stereogene_output = ".",
  context_file,
  protein_file,
  protein_file_input = NULL,
  range = c(-200, 200)
)
```

# Arguments

dir\_stereogene\_output

Directory of Stereogene output for protein. Default current directory.

context\_file Name of the RNA context file input to Stereogene. File names must exclude extensions such as ".bedGraph". Required

protein\_file A vector of at least one protein file name to be averaged for calculation of distance. File names must exclude extensions such as ".bedGraph". All files in the list should be experimental/biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Optional.

range

A vector of two integers denoting the range upstream and downstream of the center of protein binding to consider in the comparison. Ranges that are too small miss the holistic binding context, while large ranges amplify distal noise in the binding data. Cannot exceed wSize/2 from write\_config. Default c(-200, 200)

# Value

Wasserstein distance between the two halves of the binding context, with lower values suggesting greater symmetry.

#### **Examples**

visualizeCapRStereogene

visualizeCapRStereogene

#### **Description**

Creates a visual output of all CapR RNA structure contexts relative to protein binding.

#### Usage

```
visualizeCapRStereogene(
   dir_stereogene_output = ".",
   CapR_prefix,
   protein_file,
   protein_file_input = NULL,
   x_lim = c(-100, 100),
   y_lim = NULL,
   error = 1,
   nShuffle = 100,
   out_file = "out_file",
   legend = TRUE,
   heatmap = FALSE
)
```

#### Arguments

dir\_stereogene\_output

Directory of stereogene output. Default working directory.

CapR\_prefix The prefix string common to CapR output files of protein\_file. Required.

protein\_file A vector of at least one protein file name to be averaged for visualization. File names must exclude extensions such as ".bedGraph". All files in the list should

be experimental or biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Op-

tional.

x\_lim A vector of two integers denoting the lower and upper x axis limits. Cannot

exceed wSize/2 from write\_config. Default (-100, 100)

y\_lim A vector of two numbers denoting the lower and upper y axis limits. Optional

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error	A numeric value that determines the number of standard deviations to show in the error bar. Default 1
nShuffle	Relevant if multiple protein files are input and background error has been calculated. It is the number of iterations used to derive background signal error. Should be same for all protein files. Default 100.
out_file	Name of output file, excluding extension. ".pdf" or ".jpeg" will be added as relevant to the output file name. Default "out_file"
legend	Whether a legend should be included with the output graph. Default TRUE
heatmap	Whether the output graph should be in the form of a heatmap (TRUE) or of a line graph (FALSE). Default FALSE

# Value

heatmap (JPEG) or line graph (PDF) image file

# **Examples**

visualizeStereogene visualizeStereogene

# Description

Creates a visual output of a single RNA structure context relative to protein binding.

# Usage

```
visualizeStereogene(
  dir_stereogene_output = ".",
  context_file,
  protein_file,
  protein_file_input = NULL,
  x_lim = c(-100, 100),
  y_lim = NULL,
  error = 3,
  nShuffle = 1000,
```

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```
out_file = "out_file",
  legend = TRUE,
  heatmap = FALSE
)
```

#### **Arguments**

dir\_stereogene\_output

Directory of stereogene output. Default working directory.

context\_file A single context file name for visualization with the protein\_file(s). File names must exclude extensions such as ".bedGraph". Required.

protein\_file A vector of at least one protein file name to be averaged for visualization. File names must exclude extensions such as ".bedGraph". All files in the list should

be experimental or biological replicates. Required.

protein\_file\_input

A protein file name of background input to be subtracted from protein\_file signal. File name must exclude extension. Only one input file is permitted. Optional.

x\_lim A vector of two integers denoting the lower and upper x axis limits. Cannot

exceed wSize/2 from write\_config. Default (-100, 100)

y\_lim A vector of two numbers denoting the lower and upper y axis limits. Optional.

error A numeric value that determines the number of standard deviations to show in

the error bar. Default 3

nShuffle Relevant if multiple protein files are input and background error has been cal-

culated. It is the number of iterations used to derive background signal error.

Should be same for all protein files. Default 1000.

out\_file Name of output file, excluding extension. ".pdf" or ".jpeg" will be added as

relevant to the output file name. Default "out\_file"

legend Whether a legend should be included with the output graph. Default TRUE.

heatmap Whether the output graph should be in the form of a heatmap (TRUE) or of a

line graph (FALSE). Default FALSE

#### Value

heatmap (JPEG) or line graph (PDF) image file

24 write\_config

# Description

Writes a configuration file for use by Stereogenes in the working directory.

# Usage

```
write_config(
  name_config = "config.cfg",
  chrom_size,
  Rscript = FALSE,
  silent = TRUE,
  na_noise = FALSE,
  bin = 1,
  threshold = 0,
  cross_width = 200,
  wSize = 10000,
  kernel_width = 1000,
  resPath = "."
)
```

# **Arguments**

name_config	Name of output config file. Default config.cfg
chrom_size	Name of chromosome size file. File must be in two-column format without a header where first column is chromosome name and second column is chromosome length, as from getChainChrSize. Required
Rscript	Write R script for the result presentation. Equivalent to -r argument in Stereo-Gene. Default FALSE
silent	Provides an output when Stereogene is run. Equivalent to -s or -silent argument in StereoGene. Default TRUE
na_noise	Use NA values as unknown and fill them with noise. Equivalent to -NA argument in StereoGene. Default FALSE
bin	Bin size for input averaging; an integer. Default 1
threshold	Threshold for input data to remove small values. An integer between 0 and 250. Default $0$
cross_width	Width of cross-correlation plot output in Rscript; an integer. Default 200.
wSize	Window size; an integer. If windows are too small, cross correlations will have a lot of noise; if they are too large, there may be too few windows for robust statistical assessment. Default 10000
kernel_width	Kernel span in nucleotides; an integer. Equivalent to KernelSigma invStereo-Gene. Default 1000

# Value

writes a configuration file into directory

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#### Note

Not all StereoGene parameters are included in this function so refer to the StereoGene manual and modify the output .cfg file manually if additional parameters are desired.

# **Examples**

write\_fasta

write\_fasta

# **Description**

Writes a FASTA file from a vector of sequences

#### Usage

```
write_fasta(sequences, names, file.out)
```

# **Arguments**

sequences A vector of sequences

names A vector of names corresponding to the sequences

file.out Name of output FASTA file; a string

# Value

writes FASTA file into directory

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