Package 'DegNorm'

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Type Package Title DegNorm: degradation normalization for RNA-seq data Date 2024-03-26 Author Bin Xiong and Ji-Ping Wang Maintainer Ji-Ping Wang < jzwang@northwestern.edu> biocViews RNASeq, Normalization, GeneExpression, Alignment, Coverage, DifferentialExpression, BatchEffect,Software,Sequencing, ImmunoOncology, QualityControl, DataImport **Description** This package performs degradation normalization in bulk RNAseq data to improve differential expression analysis accuracy. **License** LGPL (>= 3) **Depends** R (>= 4.0.0), methods **Imports** Rcpp (>= 1.0.2), Genomic Features, txdbmaker, parallel, for each, S4Vectors, doParallel, Rsamtools (>= 1.31.2), GenomicAlignments, heatmaply, data.table, stats, ggplot2, GenomicRanges, IRanges, plyr, plotly, utils, viridis LinkingTo Rcpp, RcppArmadillo,S4Vectors,IRanges **NeedsCompilation** yes Suggests knitr,rmarkdown,formatR VignetteBuilder knitr BugReports https://github.com/jipingw/DegNorm/issues git_url https://git.bioconductor.org/packages/DegNorm git_branch RELEASE_3_21 git_last_commit 0d4cbd3 git_last_commit_date 2025-04-15 Repository Bioconductor 3.21

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Description

DegNorm is an R package for degradation normalication for bulk RNA-seq data.DegNorm, short for degradation normalization, is a bioinformatics pipeline designed to correct for bias due to the heterogeneous patterns of transcript degradation in RNA-seq data.

Details

DegNorm is a data-driven approach for RNA-Seq normalization resulting in the adjusted read count matrix. This adjustment applies to each gene within each sample, accounting for sample- and gene-specific degradation bias while simultaneously controlling for the sequencing depth. The algorithm at the center of DegNorm is the rank-one over-approximation of a gene's coverage score matrix, which is comprised of the different samples' coverage score curves along the transcript for each gene. For each gene, DegNorm estimates (1) an envelope function representing the ideal shape of the gene's coverage curve when no degradation is present, and (2) scale factors for each sample (for said gene) that indicates the relative abundance of the gene within the sample.

functions: read_coverage_batch,degnorm,plot_coverage,plot_heatmap,plot_corr,plot_boxplot

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References

DegNorm reference:

Xiong, B., Yang, Y., Fineis, F. Wang, J.-P., DegNorm: normalization of generalized transcript degradation improves accuracy in RNA-seq analysis, Genome Biology, 2019,20:75

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| CoverageClass data |
|--------------------|
|--------------------|

Description

Example of CoverageClass data from DegNorm package. It is the output from read_coverage_batch function for human chromosome 21.

Usage

```
data(coverage_res_chr21)
```

Format

A coverageClass list of the following

coverage a list of converage matrices for all genes within each sample counts a data.frame of read counts for all genes within each sample.

Examples

```
data(coverage_res_chr21)
summary_CoverageClass(coverage_res_chr21)
```

degnorm

Main function to perform degradation normalization.

Description

degnorm calcualtes the degradation index score for each gene within each sample and return the degradation-normalized read counts.

Usage

```
degnorm(read_coverage,counts,iteration,loop,down_sampling=1,grid_size=10,
cores=1)
```

Arguments

read_coverage a list of converage matrices, one per gene

counts dataframe of read counts, each row for one gene, and column for sample. The

order and number of genes must match the order in read_coverage matrices.

iteration iteration number for degnorm algorithm. 5 is sufficient.

loop iteration number inside of nonnegative matrix factorization-over approximation.

Default is 100.

down_sampling 1 for yes (default) and 0 for no. If yes, average coverage score is calcualted on a

grid of size specified by grid_size argument. The new coverage matrix formed by the grid average score will be used for baseline selection. This increases the

efficiency of algorithm while maintaining comparable accuracy.

grid_size default size is 10 bp.

cores number of cores. Default number if 1. Users should input the maximum possible

number of cores for efficiency.

Value

degnorm outputs a list of following objects:

counts a data.drame of read counts for each gene within each sample.

counts_normed a data.drame of degradation-normalized read counts for each gene within each

sample.

DI a matrix of degradation index scores for each gene within each sample.

K normalizing scale factor for each gene within each sample after accounting for

degradation normalization.

convergence convergence tag; 0 = degnorm was not done on this gene because smaller counts

or too short length.1 = degnorm was done with baseline selection. 2 = degnorm done without baseline selection because gene length (after filtering out low count regions)<200 bp. 3= baseline was found, but DI score is too large. 4 = baseline

selection didn't coverge.

envelop list of the envelop curves for all genes.

Examples

DegNorm-plot-functions

Degradation index (DI) score plot functions

Description

DegNorm provides three functions for visualization gene-/sample-wise degradation.

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Usage

```
plot_corr(DI)
plot_heatmap(DI)
plot_boxplot(DI)
```

Arguments

DΙ

a matrix or data.frame of degradation index (DI) scores with each row corresponding to one gene and each column for a sample.

Details

plot_corr plots the correlation matrix of DI scores between samples. plot_heatmap plots the heatmap of DI scores. Left is plotted in descending order of average DI scores of genes where each row corresponds to one gene. In the right plot, DI scores were sorted within each sample and plotted in descending order. plot_boxplotplots the boxplot of DI scores by samples.

Value

These functions return a boxplot of DI scores by sample, a heatmap of DIS scores of all genes in all samples and a correlation plot of DI scores between samples respectively.

Examples

```
## res_DegNorm_chr21 is degnorm otuput stored in sysdata.Rda
data(res_DegNorm_chr21)
plot_boxplot(res_DegNorm_chr21$DI)
plot_heatmap(res_DegNorm_chr21$DI)
plot_corr(res_DegNorm_chr21$DI)
```

plot_coverage

Coverage plot functions for DegNorm

Description

plot_coverage plots the before- and after-degradation coverage curves

Usage

```
plot_coverage(gene_name, coverage_output, degnorm_output, group=NULL, samples=NULL)
```

Arguments

```
gene_name the name of the gene whose coverage coverage to be plotted.

coverage_output

CoverageClass object, the output from function coverage_cal_batch.

degnorm_output DegNormClass object, the output from function DegNorm.
```

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group a vector of integers or character strings indicating the biological conditions of

the samples. Coverage curves will be plotted in the same color for the same group. Default is NULL. By default all curves will plotted in different colors.

samples a string vector for the subset of samples to be plotted. NULL means all samples

to be plotted. The length of samples must be of the same length of group if both

specified.

Details

plot_coverageoutputs the coverage curves before- and after-degradation normalization.

Value

The coverage curve before and after degradation normalization.

Examples

```
## gene named "SOD1",plot coverage curves
data(coverage_res_chr21)
data(res_DegNorm_chr21)
plot_coverage(gene_name="SOD1", coverage_output=coverage_res_chr21,
degnorm_output=res_DegNorm_chr21, group=c(0,1,1))
```

read_coverage

Function to calculate read coverage score for one bam file

Description

This function judges whether bam file is single-end and paired-end, and generate bam file index if needed. It calls function paired_end_cov_by_ch or single_end_by_ch. It takes multiple-core structure for parallel computing for efficiency.

Usage

```
read_coverage(bam_file,all_genes,cores)
```

Arguments

bam_file The name of the bam file.

all_genes An GRangesList object. It's the parsed genes annotation file from GTF file.

cores number of cores to use.

Details

This function judges whether bam file is single-end and paired-end, and generate bam file index if needed. It takes multiple-core structure for parallel computing for efficiency.

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Value

This function returns a coverageClass object. It contains a list of: (1) a list of coverage score for each gene in RLE format and (2) a dataframe for read counts

See Also

```
read_coverage_batch
```

Description

This function calls read_coverage to compute read coverage socre and read counts for all genes and samples.

Notes: 1. Coverage score is calcualted per gene, i.e. concatenation of all exons from the same gene.

- 2. We follow HTseq protocol for counting valid read or read pairs for each gene.
- 3. When reading alignment file, isSecondaryAlignment flag is set as FALSE to avoid possible redundant counting.
- 4. For paired-end data, isPaired is set as TRUE. We don't recommend setting isProperPair as TRUE as some fragments length may exceed 200bp.
- 5. User can modify scanBamParam in the R codes below as needed.

Usage

```
read_coverage_batch(bam_file_list,gtf_file,cores=1)
```

Arguments

bam_file_list a character vector of bam file names.

gtf_file that RNA-seq reads were aligned with reference to.

cores number of cores to be used. Default=1.

Value

A list of the following:

coverage a list of converage matrices for all genes within each sample.

counts data.frame of read counts for all genes within each sample.

See Also

read_coverage

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Examples

res_DegNorm_chr21

Example DegNormClass data

Description

Example of DegNormClass data from DegNorm package. It is the output from degnorm function for human chromosome 21.

Usage

```
data("res_DegNorm_chr21")
```

Format

A DegNormClass list of the following items:

counts a data.drame of read counts for each gene within each sample.

counts_normed a data.drame of degradation-normalized read counts for each gene within each sample.

DI a matrix of degradation index scores for each gene within each sample.

K normalizing scale factor for each gene within each sample after accounting for degradation normalization.

convergence convergence tag; 0 = degnorm was not done on this gene because smaller counts or too short length.1 = degnorm was done with baseline selection. 2 = degnorm done without baseline selection because gene length (after filtering out low count regions)<200 bp. 3= baseline was found, but DI score is too large. 4 = baseline selection didn't coverge.

envelop a list of the envelop curves for all genes.

Examples

```
data(res_DegNorm_chr21)
summary_DegNormClass(res_DegNorm_chr21)
```

summary_CoverageClass Summary method for CoverageClass.

Description

It prints a summary of the data objects contained in the list from read_coverage_batch.

Usage

```
summary_CoverageClass(object)
```

Arguments

object

CoverageClass from coderead_coverage_batch.

Value

On-screen plot of summary of CoverageClass object.

Examples

```
## Summary of coverage_cal_batch output (CoverageClass)
data(coverage_res_chr21)
summary_CoverageClass(coverage_res_chr21)
```

summary_DegNormClass Summary method for DegNormClass.

Description

It prints a summary of the data objects contained in the list from degnorm function.

Usage

```
summary_DegNormClass(object)
```

Arguments

object

DegNormClass from degnorm function.

Value

On-screen summary of DegNormClass object.

Examples

```
## Summary of degnorm output (DegNormlass)
data(res_DegNorm_chr21)
summary_DegNormClass(res_DegNorm_chr21)
```

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