Package 'TargetScoreData'

October 31, 2024

Description Precompiled and processed miRNA-overexpression fold-changes from 84 Gene Expres-

Title TargetScoreData

Maintainer Yue Li <yueli@cs.toronto.edu>

Version 1.42.0 **Author** Yue Li

sion Omnibus (GEO) series corresponding to 6 platforms, 77 human cells or tissues, and 113 distinct miRNAs. Accompanied with the data, we also included in this package the sequence feature scores from TargetScanHuman 6.1 including the context+ score and the probabilities of conserved targeting for each miRNA-mRNA interaction. Thus, the user can use these static sequence-based scores together with user-supplied tissue/cell-specific fold-change due to miRNA overexpression to predict miRNA targets using the package TargetScore (download separately)
biocViews ExperimentData, RNASeqData, miRNAData
License GPL-2
Suggests TargetScore, gplots
git_url https://git.bioconductor.org/packages/TargetScoreData
git_branch RELEASE_3_20
git_last_commit 59ea064
git_last_commit_date 2024-10-29
Repository Bioconductor 3.20
Date/Publication 2024-10-31
Contents
TargetScoreData-package
get_miRNA_family_info
get_miRNA_transfection_data
get_precomputed_logFC
get_precomputed_targetScores
get_TargetScanHuman_contextScore
get_TargetScanHuman_PCT
get_validated_targets
Index 6

TargetScoreData-package

Processed human microRNA perturbation data from GEO, and sequence information from TargetScan, and targetScore from TargetScore

Description

To automate the pipeline of calculating targetScore using targetScore, we compiled, processed and generated miRNA-overexpression fold-changes from 84 Gene Expression Omnibus (GEO) series corresponding to 6 platforms, 77 human cells or tissues, and 113 distinct miRNAs. To our knowledge, this is by far the largest miRNA-pertubation data compendium. Accompanied with the data, we also included in this package the sequence feature scores from TargetScanHuman 6.1 including the context+ score and the probabilities of conserved targeting for each miRNA-mRNA interaction. Thus, the user can use these static sequence-based scores together with user-supplied tissue/cell-specific fold-change due to miRNA overexpression to predict miRNA targets using targetScore.

Details

Package: TargetScoreData

Type: Package Version: 0.99.4 Date: 2013-07-13 License: GPL-2

get_TargetScanHuman_contextScore: TargetScan context score for all of the predicted sites (including conserved and nonconserved sites) downloaded from TargetScan 6 website (http://www.targetscan.org/cgibin/targetscan/data_download.cgi?db=vert_61)

get_TargetScanHuman_PCT: TargetScan PCT for all of the predicted sites (including conserved and nonconserved sites) downloaded from TargetScan 6 website (http://www.targetscan.org/cgibin/targetscan/data_download.cgi?db=vert_61)

get_miRNA_family_info: Get miRNA family information obtained TargetScanHuman 6

get_precomputed_targetScores: Pre-calculated targetScores for 112 miRNAs using logFC and sequence scores from TargetScan context score and PCT from the RData files above.

get_precomputed_logFC: We combined all of the logFC data columns into a single N x M matrix for all of the N = 19177 RefSeq mRNAs (NM $_*$ obtained from UCSC) and M = 286 datasets. Missing data (logFC) for some genes across studies were imputed using impute from impute.knn.

get_miRNA_transfection_data: Get log fold-changes (logFC) from each study organized in a list with each item corresponding to a miRNA. Notably, some miRNAs (e.g., hsa-miR-1) appear more than once in the list corresponding to different studies.

get_validated_targets: Get validated targets of human miRNA obtained from mirTarBase v3.5

Author(s)

Yue Li

Maintainer: Yue Li <yueli@cs.toronto.edu>

References

miRTarBase: a database curates experimentally validated microRNA-target interactions. (2011). miRTarBase: a database curates experimentally validated microRNA-target interactions., 39(Database issue), D163-9. doi:10.1093/nar/gkq1107

Friedman, R. C., Farh, K. K.-H., Burge, C. B., & Bartel, D. P. (2009). Most mammalian mRNAs are conserved targets of microRNAs. Genome Research, 19(1), 92-105. doi:10.1101/gr.082701.108

Garcia, D. M., Baek, D., Shin, C., Bell, G. W., Grimson, A., & Bartel, D. P. (2011). Weak seed-pairing stability and high target-site abundance decrease the proficiency of lsy-6 and other microR-NAs. Nature structural & molecular biology, 18(10), 1139-1146. doi:10.1038/nsmb.2115

Please follow the GEO series number to find the references for each microRNA-transfection dataset.

See Also

```
get_TargetScanHuman_contextScore, get_TargetScanHuman_PCT, get_validated_targets,
get_miRNA_family_info, get_precomputed_targetScores, get_precomputed_logFC, get_miRNA_transfection_
```

Examples

```
ls("package:TargetScoreData")
```

```
get_miRNA_family_info MicroRNA family information
```

Description

Get human miRNA family information obtained frin TargetScanHuman 6 website

Usage

```
get_miRNA_family_info(datapath = system.file("extdata/miR_Family_Info.txt", package = "TargetScore")
```

Arguments

```
datapath data path
```

Description

Get log fold-changes (logFC) from each study organized in a list with each item corresponding to a miRNA. Notably, some miRNAs (e.g., hsa-miR-1) appear more than once in the list corresponding to different studies.

Usage

```
get_miRNA_transfection_data(datapath = system.file("extdata/miRNA_transfection_data.RData", packa
```

Arguments

datapath data path

get_precomputed_logFC get_precomputed_logFC

Description

We combined all of the logFC data columns into a single N x M matrix for all of the N = 19177 RefSeq mRNAs (NM $_{-}$ * obtained from UCSC) and M = 286 datasets. Missing data (logFC) for some genes across studies were imputed using impute from impute.knn.

Usage

```
get_precomputed_logFC(datapath = system.file("extdata/logFC.RData", package = "TargetScoreData"))
```

Arguments

datapath data path

get_precomputed_targetScores

get_precomputed_targetScores

Description

Pre-calculated targetScores for 112 miRNAs using logFC and sequence scores from TargetScan context score and PCT from the RData files above.

Usage

get_precomputed_targetScores(datapath = system.file("extdata/targetScores.RData", package = "TargetScores")

Arguments

datapath data path

Description

TargetScan context score for all of the predicted sites (including conserved and nonconserved sites) downloaded from TargetScan 6 website (http://www.targetscan.org/cgi-bin/targetscan/data_download.cgi?db=vert_61)

Usage

```
\verb|get_TargetScanHuman_contextScore| (datapath = system.file("extdata/TargetScanHuman_contextScore.RDatapath = system.file("extdata/TargetScore.RDatapath = system.file("extdata/TargetScore.RDatapath = system.file("extdata/TargetScore.RDatapath = system.file("extdata/TargetScore.RDatapath = system.file("extdata/TargetScore.RDatapath = system.file("extdata/TargetScore.RD
```

Arguments

datapath data path

```
\label{eq:confluence} get\_TargetScanHuman\_PCT \\ get\_TargetScanHuman\_PCT
```

Description

TargetScan PCT for all of the predicted sites (including conserved and nonconserved sites) downloaded from TargetScan 6 website (http://www.targetscan.org/cgi-bin/targetscan/data_download.cgi?db=vert_61)

Usage

```
\verb|get_TargetScanHuman_PCT| (datapath = system.file("extdata/TargetScanHuman_PCT.RData", package = "TargetScanHuman_PCT.RData", packag
```

Arguments

datapath data path

```
get_validated_targets
```

Description

Get validated targets of human miRNA obtained from mirTarBase v3.5

Usage

```
get_validated_targets(datapath = system.file("extdata/hsa_MTI.xls", package = "TargetScoreData"))
```

Arguments

datapath data path

Index