Package 'IntramiRExploreR'

May 19, 2017

```
Title Predicting Targets for Drosophila Intragenic miRNAs
Version 1.00.0
Author Surajit Bhattacharya and Daniel Cox
Maintainer Surajit Bhattacharya <sbhattacharya3@student.gsu.edu>
Description Intra-miR-ExploreR, an integrative miRNA target prediction
      bioinformatics tool, identifies targets combining expression and biophysical
      interactions of a given microRNA (miR). Using the tool, we have identified
      targets for 92 intragenic miRs in D. melanogaster, using available microarray
      expression data, from Affymetrix 1 and Affymetrix2 microarray array platforms,
      providing a global perspective of intragenic miR targets in Drosophila.
      Predicted targets are grouped according to biological functions using the DAVID
      Gene Ontology tool and are ranked based on a biologically relevant scoring
      system, enabling the user to identify functionally relevant targets for a given
      miR.
Depends R (>= 3.4)
Imports igraph (>= 1.0.1),
      FGNet (>= 3.0.7),
      knitr (>= 1.12.3),
      stats.
      utils,
      grDevices,
     graphics
Suggests RDAVIDWebService,
      gProfileR,
      topGO,
      KEGGprofile,
      org.Dm.eg.db,
      rmarkdown,
      testthat
VignetteBuilder knitr
License GPL-2
biocViews Software, Microarray, GeneTarget, StatisticalMethod, GeneExpression, GenePrediction
Encoding UTF-8
LazyData true
RoxygenNote 5.0.1
URL https://github.com/sbhattacharya3/IntramiRExploreR/
```

BugReports https://github.com/sbhattacharya3/IntramiRExploreR/issues

R topics documented:

	Affy1_Distance_Final
	Affy1_Pearson_Final
	Affy2_Distance_Final
	Affy2_Pearson_Final
	extract_HostGene
	extract_intragenic_miR
	genes_Stat
	Gene_Visualisation
	GetGOS_ALL
	IntramiRExploreR
	miRNA_ID_to_Function
	miRNA_summary_DB
	miRTargets_Stat
	Visualisation
Index	11

Affy1_Distance_Final Targets for the microRNA analyzed from Affy1 plaform using Distance.

Description

A precomputed dataset containing the targets, scores and other attributes of 83 intragenic microR-NAs using Distance Correlation for plaform Affymetrix 1.

Usage

Affy1_Distance_Final

Format

A data frame with 53399 rows and 8 variables:

miRNA miRNA name, miRNA symbol

GeneSymbol Gene name, in Gene Symbol

FBGN Gene name, in FlybaseID

CGID Gene name, in CGID

Score Computed Score, in float

GeneFunction Gene Functions, from Flybase

experiments Experiments, from ArrayExpress

TargetDatabases Target Database Name, from TargetDatabases

Affy1_Pearson_Final 3

Affy1_Pearson_Final

Targets for the microRNA analyzed from Affy1 plaform using Pearson.

Description

A precomputed dataset containing the targets, scores and other attributes of 83 intragenic microR-NAs using Pearson Correlation for plaform Affymetrix 1.

Usage

Affy1_Pearson_Final

Format

A data frame with 41845 rows and 8 variables:

miRNA miRNA name, miRNA symbol

GeneSymbol Gene name, in Gene Symbol

FBGN Gene name, in FlybaseID

CGID Gene name, in CGID

Score Computed Score, in float

GeneFunction Gene Functions, from Flybase

experiments Experiments, from ArrayExpress

TargetDatabases Target Database Name, from TargetDatabases

Affy2_Distance_Final Targets for the microRNA analyzed from Affy2 plaform using Distance.

Description

A precomputed dataset containing the targets, scores and other attributes of 83 intragenic microR-NAs using Distance Correlation for plaform Affymetrix 1.

Usage

Affy2_Distance_Final

Format

A data frame with 73374 rows and 8 variables:

miRNA miRNA name, miRNA symbol

GeneSymbol Gene name, in Gene Symbol

FBGN Gene name, in FlybaseID

CGID Gene name, in CGID

Score Computed Score, in float

GeneFunction Gene Functions, from Flybase

experiments Experiments, from ArrayExpress

TargetDatabases Target Database Name, from TargetDatabases

4 extract_HostGene

Affy2_Pearson_Final

Targets for the microRNA analyzed from Affy2 plaform using Pearson.

Description

A precomputed dataset containing the targets, scores and other attributes of 83 intragenic microR-NAs using Pearson Correlation for plaform Affymetrix 1.

Usage

Affy2_Pearson_Final

Format

A data frame with 52913 rows and 8 variables:

miRNA miRNA name, miRNA symbol

GeneSymbol Gene name, in Gene Symbol

FBGN Gene name, in FlybaseID

CGID Gene name, in CGID

Score Computed Score, in float

GeneFunction Gene Functions, from Flybase

experiments Experiments, from ArrayExpress

TargetDatabases Target Database Name, from TargetDatabases

extract_HostGene

Extract Host Gene for a given Intragenic miRNA.

Description

Extract Host Gene for a given Intragenic miRNA.

Usage

```
extract_HostGene(miRNA)
```

Arguments

miRNA

A String containing the miRNA name.

Value

genf, a character string or vector containing Host gene for the Intragenic miRNA.

```
miRNA="dme-miR-12"
extract_HostGene(miRNA)
```

extract_intragenic_miR 5

```
extract_intragenic_miR
```

Extract Intragenic miRNA for a given Host gene.

Description

Extract Intragenic miRNA for a given Host gene.

Usage

```
extract_intragenic_miR(gene)
```

Arguments

gene character. Gene Symbol.

Value

miRf, a character string or vector containing Intragenic miRNA for the Host Gene.

Examples

```
gene="Gmap"
extract_intragenic_miR(gene)
```

genes_Stat

Extracting miRNAs that target a query gene.

Description

Extracting miRNAs that target a query gene.

Usage

```
genes_Stat(gene, geneIDType = c("GeneSymbol", "FBGN", "CGID"),
method = c("Pearson", "Distance", "Both", "BothIntersect"),
Platform = c("Affy1", "Affy2"), Text = FALSE, outpath = tempdir())
```

Arguments

gene character. gene Identifier.

geneIDType character. GeneIDtype choices are 'GeneSymbol', 'FBGN', 'CGID' method character. Choices are 'Pearson', 'Distance', 'Both' and 'BothIntersected'

Platform character. Choices are 'Affy1', 'Affy2'.

Text logical . To choose between storing the data as text file. Default is FALSE. outpath character. The path where the data is stored if TEXT=TRUE. Default is

6 Gene_Visualisation

Value

Outputs the miRNA information, Target Prediction Score, miRNA miRNA function and Target Database that predicts the interaction in a dataframe. Depending upon the ouput choice data is stored in the path specified. Default option prints output to the console.

Examples

Gene_Visualisation

Visualises the targetGene:miRNA network using Cytoscape and igraph

.

Description

Visualises the targetGene:miRNA network using Cytoscape and igraph .

Usage

```
Gene_Visualisation(mRNA, mRNA_type = c("GeneSymbol", "FBGN", "CGID"),
  method = c("Pearson", "Distance", "Both"), platform = c("Affy1", "Affy2"),
  visualisation = c("igraph", "Cytoscape", "text", "console"),
  path = tempdir(), layout = c("kamadakawai", "reingold.tilford",
  "fruchterman.reingold", "interactive"))
```

Arguments

mRNA character. gene Identifier.

mRNA_type character. mRNA id type. The choices are 'GeneSymbol','FBID' and 'CGID'.

method character. Statistical Methods. Choices are 'Pearson', 'Distance', 'Both'

platform character. Affymetrix Platforms. Choices are 'Affy1', 'Affy2'.

visualisation character. Visualisation type. Choices are 'igraph', 'Cytoscape', 'text' and "con-

sole"

path character. Path where data.frame is saved when visualisation is text. Default is

tempdir().

layout character. Network choices. Choices are 'kamadakawai', 'reingold.tilford', 'fruchterman.reingold'

and 'interactive'.

Value

Depending upon the ouput choice network image or dataframe containg miRNAs that target the query gene are ouput.

GetGOS_ALL 7

Description

Gene ontology for Target Genes.

Usage

```
GetGOS_ALL(gene, GO = c("DAVID", "topGO"), term = c("GOTERM_BP_ALL",
   "GOTERM_MF_ALL", "GOTERM_CC_ALL"), geneIdType = "ALIAS", email,
   path = tempdir(), ontology = c("GO_BP", "GO_MF", "GO_CC"), filename)
```

Arguments

gene	List A String or vector containing the Gene names.
GO	A String depicting the chosen GO tool. Choices are "David" and "topGO"
term	A String depicting the chosen term. Choices are "GOTERM_BP_ALL", "GOTERM_MF_ALL", "GOTERM_CC_ALL".
geneIdType	Type of gene Id given as input. Default "ALIAS"
email	Email Id to connect to David.
path	String. The path where the data is stored if TEXT=TRUE.
ontology	Ontology selection for topGO. Choices are "GO_BP", "GO_MF", "GO_CC".
filename	Name of the file to store Gene Ontology.

Value

Depending upon the ouput choice data is stored in the path specified. Default option prints output to the console.

 ${\tt IntramiRExploreR}$

IntramiRExploreR:Prediction of targets for Intragenic miRNA in Drosophila.

Description

Prediction of targets for Drosophila Intragenic microRNAs and Integrated approach using Gene Ontology and Networking tools.

Examples

```
## Not run:
gene='Gmap'
extract_intragenic_miR(gene)
## End(Not run)
```

miRNA_ID_to_Function

Contains the miRNA function information from Flybase database.

Description

A dataset containing the function for the intragenic miRNA.

Usage

```
miRNA_ID_to_Function
```

Format

A data frame with 66 rows and 4 variables:

```
miRNA miRNA name, miRNA symbolFBGN target gene name, gene symbolmiRNAFunction miRNA function, from Flybase
```

Source

```
http://flybase.org/
```

miRNA_summary_DB

miRNA_summary_DB

Contains the summary for the intragenic miRNA.

Description

A dataset containing the summary for the intragenic miRNA.

Usage

```
miRNA_summary_DB
```

Format

A data frame with 257 rows and 6 variables:

miRNA miRNA name, miRNA symbol
Intragenic Responsee, in boolean
Intergenic Responsee, in boolean
Gene miRNA name, miRNA symbol

Type.of.HostGene.mRNA.lncRNA. Type of Hostgene

Notes Comments about the miRNA

miRTargets_Stat

Extracting miRNAs that target a query gene.

Description

Extracting miRNAs that target a query gene.

Usage

```
miRTargets_Stat(miR, method = c("Pearson", "Distance", "Both",
   "BothIntersect"), Platform = c("Affy1", "Affy2"), outpath = tempdir(),
   Text = FALSE)
```

Arguments

miR character. miRNA symbol.

method character. Choices are "Pearson", "Distance", "Both" and "BothIntersected"

Platform character. Choices are "Affy1", "Affy2".

outpath character. The path where the data is stored if TEXT=TRUE. Default is tem-

pdir().

Text logical . To choose between storing the data as text file. Default is FALSE.

Value

Outputs the target information, Target Prediction Score, miRNA target function and Target Database that predicts the interaction in a dataframe. Depending upon the ouput choice data is stored in the path specified. Default option prints output to the console.

10 Visualisation

Examples

```
\label{eq:miral} $$\min RNA="dme-miR-12"$ $$\min RTargets\_Stat (miRNA,method=c ("Pearson"),Platform=c ("Affy1"),Text=FALSE)$ $$
```

Visualisation

Visualises the targetGene:miRNA network using Cytoscape and igraph

•

Description

Visualises the targetGene:miRNA network using Cytoscape and igraph.

Usage

```
Visualisation(miRNA, mRNA_type = c("GeneSymbol", "FBID", "CGID"),
  method = c("Pearson", "Distance", "Both"), platform = c("Affy1", "Affy2"),
  thresh = 50, visualisation = c("igraph", "Cytoscape", "Text", "console"),
  path = tempdir(), layout = c("kamadakawai", "reingold.tilford",
  "fruchterman.reingold", "interactive"))
```

Arguments

miRNA character, miRNA Identifier.

mRNA_type character. mRNA id type. The choices are 'GeneSymbol', 'FBID' and 'CGID'.

method character. Statistical Methods. Choices are 'Pearson', 'Distance', 'Both'

platform character. Affymetrix Platforms. Choices are 'Affy1','Affy2'.

thresh integar. Threshold depicting number of rows to show.

visualisation character. Visualisation type. Choices are 'igraph', 'Cytoscape', 'text' and 'con-

sole'.

path character. Path where data.frame is saved when visualisation is text. Default is

tempdir().

layout character. Network choices. Choices are 'kamadakawai', 'reingold.tilford', 'fruchterman.reingold'

and 'interactive'.

Value

Depending upon the ouput choice network image or dataframe containg miRNAs that target the query gene are ouput.

Index

```
*Topic datasets
    Affy1_Distance_Final, 2
    Affy1_Pearson_Final, 3
    Affy2_Distance_Final, 3
    Affy2_Pearson_Final, 4
    {\tt miRNA\_ID\_to\_Function,\,8}
    miRNA_summary_DB, 9
Affy1_Distance_Final, 2
Affy1_Pearson_Final, 3
Affy2_Distance_Final, 3
Affy2_Pearson_Final, 4
extract_HostGene, 4
\verb|extract_intragenic_miR|, 5
Gene_Visualisation, 6
genes_Stat, 5
GetGOS_ALL, 7
IntramiRExploreR, 8
IntramiRExploreR-package
        (IntramiRExploreR), 8
miRNA_ID_to_Function, 8
miRNA_summary_DB, 9
miRTargets_Stat, 9
Visualisation, 10
```