Package 'miRNAtap'

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Type Package Title miRNAtap: microRNA Targets - Aggregated Predictions Version 1.42.0 Date 2016-10-03 Author Maciej Pajak, T. Ian Simpson Maintainer T. Ian Simpson <ian.simpson@ed.ac.uk> **Description** The package facilitates implementation of workflows requiring miRNA predictions, it allows to integrate ranked miRNA target predictions from multiple sources available online and aggregate them with various methods which improves quality of predictions above any of the single sources. Currently predictions are available for Homo sapiens, Mus musculus and Rattus norvegicus (the last one through homology translation). License GPL-2 Depends R (>= 3.3.0), AnnotationDbi Imports DBI, RSQLite, stringr, sqldf, plyr, methods Suggests topGO, org.Hs.eg.db, miRNAtap.db, testthat biocViews Software, Classification, Microarray, Sequencing, miRNA **Roxygen** list(wrap = FALSE) RoxygenNote 5.0.1 git_url https://git.bioconductor.org/packages/miRNAtap git_branch RELEASE_3_21 git_last_commit a81e69a git_last_commit_date 2025-04-15 **Repository** Bioconductor 3.21 Date/Publication 2025-07-16

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aggregateRanks Aggreagate ranks from multiple sources with various methods

Description

This function performs aggregation phase of target prediction for getPredictedTargets. Consensus ranking is derived from multiple individual rankings. Available methods include minimum, maximum and geometric mean with further tuning parameters which promote true positives at the top of the final ranking

Usage

aggregateRanks(ranks, n_valid_srcs, min_src, method = "geom",
 promote = TRUE)

Arguments

ranks	data.frame with ordered scores
n_valid_srcs	number of valid sources in the dataset
min_src	minimum acceptable number fo sources
method	'min','max',or 'geom', default 'geom'
promote	add weights to improve accuracy of the method, default TRUE

Value

data.frame object with ranks per source and aggregate ranks

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getPredictedTargets

Examples

getPredictedTargets Get aggregated ordered list of predicted targets for miRNA

Description

This method performs aggregation of target lists from multiple sources. Aggregated list is more accurate than any list from a single source. Multiple aggregation methods are available.Direct target data from five sources for Human and Mouse is supplied through miRNAtap.db package, for Rat targets are derived through homology translations whenever direct ones are not available.

Usage

```
getPredictedTargets(mirna, sources = c("pictar", "diana",
"targetscan", "miranda", "mirdb"), species = "mmu", min_src = 2,
method = "geom", promote = TRUE, synonyms = TRUE, both_strands = FALSE, ...)
```

Arguments

mirna	miRNA in a standard format
sources	a list of sources to use for aggregation, default is all five sources, i.e. c('pictar', 'diana', 'targetscan
species	species in a standard three-letter acronym, 'mmu' and 'hsa' available as direct targets, 'rno' as homology translations, default 'mmu'
min_src	minimum number of sources required for a target to be considered, default 2
method	method of aggregation - choose from 'min', 'max', and 'geom'; 'min' is a minimum of ranks, 'max' is a maximum of ranks, and default 'geom' is based on geometric mean of the ranks which proves to be the most accurate method.
promote	add weights to improve accuracy of the method, default TRUE
synonyms	when searching for -3p miRNA automatically also searches for miRNA with the same name but ending with * (some databases list -3p miRNA this way) and other way around, similarly for -5p miRNA, default TRUE
both_strands	overrides synonyms and searches for targets of both -5p and -3p strands together
	any optional arguments

Details

Tuning min_src parameter is an easy way of prioritising precision at the top of the list (high values) or total recall (low values). For the five default input sources, recommended values are 2, 3, or 4.

Value

data.frame object where row names are entrez IDs of target genes, ranks from individual sources and aggregated rank are shown in columns. If no targets are found in any of the sources NULL and a warning are returned.

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References

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Wong N and Wang X (2015) miRDB: an online resource for microRNA target prediction and functional annotations. Nucleic Acids Research. 43(D1):D146-152.

Examples

```
targets <- getPredictedTargets('let-7a',species='hsa', method = 'min')
head(targets) #top of the list with minimum aggregation
targets2 <- getPredictedTargets('let-7a',species='hsa', method='geom')
head(targets2) #top of the list with geometric mean aggregation</pre>
```

getTargetsFromSource Get target list from a single source

Description

This function queries precompiled annotation SQLite database which contains miRNA - target gene associations with their respective scores.

Usage

```
getTargetsFromSource(mirna, species = "mmu", source = "diana",
    synonyms = TRUE, both_strands = FALSE)
```

Arguments

mirna	miRNA in a standard format
species	species in a standard three-letter acronym, default 'mmu'
source	a source target prediction algorithm table to query, default 'diana', other pos- sible values are 'miranda', 'targetscan', and 'pictar'.
synonyms	when searching for -3p miRNA automatically also searches for miRNA with the same name but ending with * (some databases list -3p miRNA this way) and other way around, similarly for -5p miRNA, default TRUE
both strands	overrides synonyms and searches for targets of both -5p and -3p strands together

Value

data.frame object with entrez IDs of target genes and their scores, if there are no targets found for a given miRNA in a given table then an empty

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References

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Examples

```
targets <- getTargetsFromSource('let-7a', species='hsa', source='targetscan')
head(targets)
#top of the listof human targets of let-7a from TargetScan only</pre>
```

MirnaDb-class

Description

object of MirnaDb class holds the sqlite database connection, and extends AnnotationDb class from AnnotationDbi package. columns, keys, keytypes and select methods allow access to database tables and retrieval of miRNA target information.

select is the most important method, allows querying the database for predictions from a specific source and species for a given miRNA

Usage

```
columns(x)
keytypes(x)
keys(x, keytype, ...)
select(x, keys, columns, keytype, ...)
## S4 method for signature 'MirnaDb'
columns(x)
## S4 method for signature 'MirnaDb'
keytypes(x)
## S4 method for signature 'MirnaDb'
keys(x, keytype, ...)
## S4 method for signature 'MirnaDb'
```

```
select(x, keys, columns, keytype, ...)
```

Arguments

х	the MirnaDb object
keytype	the keytype that matches the keys used; the table in which the search should be performed.
	any optional arguments
keys	the key to select records for from the database - miRNA name; all possible keys (miRNAs) are returned by using the keys method.
columns	in this case same as keytype, the table in which the search should be performed, this value specifies the source of predictions as well as species; as with keys, all possible columns are returned by using the columns method.

Value

string vectors, for select a data.frame with target genes and scores

miRNAtap

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Examples

```
#first load the annotations
require(miRNAtap.db)
#see all available tables
keytypes(miRNAtap.db)
```

miRNAtap

miRNAtap: microRNA Targets - Aggregated Predictions.

Description

It is a package with tools to facilitate implementation of workflows requiring miRNA prediction through access to multiple prediction results (DIANA, Targetscan, PicTar, Miranda, and miRDB) and their aggregation. Three aggregation methods are available: minimum, maximum and geometric mean, additional parameters provide further tuning of the results. Predictions are available for Homo sapiens, Mus musculus and Rattus norvegicus (the last one through homology translation).

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Examples

```
#direct targets in mouse aggregated from all sources:
targets_mouse <- getPredictedTargets('let-7a', species='mmu', method='geom')
#homology-translated targets in rat aggregated from all sources
targets_rat <- getPredictedTargets('let-7a', species='rno', method='geom')</pre>
```

translate

Homology transfer for miRNAtap

Description

This function maps gene entrez ID between species using homology information from Homologene.

Usage

```
translate(entrezes, from = "mmu", to = "rno", ...)
```

translate

Arguments

entrezes	data.frame with entrez Gene IDs and their scores
from	origin species, default 'mmu', Mus musculus
to	target species, default
	any optional arguments

Value

data.frame object with orthologous genes' entrez IDs and corresponding scores

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