How to use MiRaGE Package

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1 Introduction

This document describes briefly how to use MiRaGE package in order to infer the target gene regulation by miRNA, based upon target gene expression.

MiRaGE is based upon the algorithm proposed in [1, 2, 3]. Basically, its function is the same as the MiRaGE Server. In order to infer the target gene regulation by miRNAs, we made use of target gene (mRNA) expression. Suppose x_{gs} is the expression of the gth gene in the sth sample. Then *P*-value to measure the amount of target gene regulation by the mth miRNAs is computed by several statistical test. More detailed and comprehensive explanations can be found in [4].

2 Background

miRNA is short non-coding RNA (ncRNA) which is believed to degrade target genes. Target genes are believed to be decided by seed match between 7-mer at 5' untranslated region (UTR) of miRNA and 3' UTR of target mRNAs. However because of huge number of miRNAs (c.a. 1000) and the huge number of target genes (c.a. hundreds) of each miRNA, it is not easy to experimentally decide which miRNA regulates target genes.

MiRaGE infers target gene regulation from target gene expression and computationally predicted target gene table. It gives the rejection probability to reject null hypothesis that target genes of a specific miRNA are equally regulated as other genes.

When t-test is employed, P-value is

$$P(S_m^{ss'} > S_m'^{ss'})$$

or

$$P(S_m^{ss'} < S_m'^{ss'})$$

where P is the rejection probability of null hypothesis $S_m = S'_m$ when the alternative hypothesis is either $S_m > S'_m$ or $S_m < S'm$. S_m and S'_m are the test variable to measure the target gene regulation by the *m*th miRNA.

When *P*-values are computed via *t*-test, S_m is the mean gene expression logarithmic ratio of the *m*th miRNA's target genes, i.e.,

$$S_m^{ss'} = \frac{1}{N(G_m)} \sum_{g \in G_m} \log \frac{x_{gs}}{x_{gs'}}$$

where G_m is the set of the *m*th miRNA's target genes and $N(G_m)$ is the total number of genes in G_m . S'_m is the mean expression logarithmic ratio of genes not targeted by the *m*th miRNA but any other miRNAs and is defined as

$$S'_{m}^{'ss'} = \frac{1}{N(G'_{m})} \sum_{g \in G'_{m}} \log \frac{x_{gs}}{x_{gs'}}$$

where G'_m is the set of the *m*th miRNA's target genes and $N(G'_m)$ is the total number of genes in G'_m .

On the other hands, if P-values are computed by Wilcoxon rank sum test, they are

$$P\left(U_m^{ss'} > \frac{N(G_m)N(G_m')}{2}\right)$$

or

$$P\left(U_m^{ss'} < \frac{N(G_m)N(G_m')}{2}\right)$$

which are the rejection probabilities of null hypothesis $U = \frac{N(G_m)N(G'_m)}{2}$ when the alternative hypothesis is either $U > \frac{N(G_m)N(G'_m)}{2}$ or $U < \frac{N(G_m)N(G'_m)}{2}$. Here $U^{ss'}$ is the test variable,

$$U_m^{ss'} = R_m^{ss'} - \frac{N(G_m)(N(G_m) + 1)}{2}$$

Here

$$R_m^{ss'} = \sum_{g \in G_m} R^{ss'} \left(\log \frac{x_{gs}}{x_{gs'}} \right)$$

where $R^{ss'}(\ldots)$ is the rank order of the *m*th genes logarithmic ratio among all of considered genes.

Alternatively, Kolmogorov-Smirnov test can be employed. In this case, test variable is

$$D_m^{ss'} = \sup_g \left(F_m(x_{gs}) - F'_m(x_{gs}) \right)$$

or

$$D'_{m}^{ss'} = \sup_{g} \left(F'_{m}(x_{gs}) - F_{m}(x_{gs}) \right)$$

where

$$F_m^{ss'}(x_{gs}) = \frac{1}{N(G_m)} \sum_{g' \in G_m} \Theta\left(\log\frac{x_{gs}}{x_{gs'}} - \log\frac{x_{g's}}{x_{g's'}}\right)$$

and

$$F'_{m}^{ss'}(x_{gs}) = \frac{1}{N(G'_{m})} \sum_{g' \in G'_{m}} \Theta\left(\log \frac{x_{gs}}{x_{gs'}} - \log \frac{x_{g's}}{x_{g's'}}\right)$$

where $\Theta(x)$ is the step function,

$$\Theta(x) = \begin{cases} 1 & x > 0 \\ 0 & x < 0 \end{cases}$$

Then *P*-value is computed via $P(D_m^{ss'} > 0)$ or $P(D'_m^{ss'} > 0)$ under the null hypothesis $D_m^{ss'} = 0$ or $D'_m^{ss'} = 0$.

Since the target genes table is generated by the simple seed match, MiRaGE does not need any other external programs to obtain target gene table. Another advantage is the exclusion of mRNA targeted by no miRNAs. This enables us more accurate prediction of target gene regulation by miRNAs.

3 Quick start

```
> library(MiRaGE)
```

```
> data(gene_exp)
```

```
> library(Biobase)
```

```
> result <- MiRaGE(gene_exp,species="HS")</pre>
```

Then result\$P0 and result\$P1 include *P*-values for upregulation and downregulation of target genes by miRNAs, respectively. The definition "up" or "down" depends upon the order of columns of expression data in gene_exp (see below).

<u>Caution</u> I strongly recommend user to use location="web" option, since it will be most frequently updated. Default setting requires experimental package *miRNATarget* (see Sec. 6.3). Data set on the web can be stored for the later usage, too (see below).

```
> result <- MiRaGE(gene_exp,location="web",species="HS")</pre>
```

4 Data Structure

4.1 Input: target gene expression

In order to execute analysis, you need ExpressionSet objects which stores target gene expression in it. In order to see this, it is easier to see sample data gene_exp as follows.

```
> data(gene_exp)
> gene_exp
ExpressionSet (storageMode: lockedEnvironment)
assayData: 45015 features, 4 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: neg.1 neg.2 day1.1 day1.2
  varLabels: sample_name
  varMetadata: labelDescription
featureData
  featureNames: 1 2 ... 45015 (45015 total)
  fvarLabels: gene_id
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

The above example displays the ExpressionSet object, gene_exp. As you can see, each row corresponds to each gene and each column corresponds to each sample (experiment).

gene_id in featureData must includes gene id. sample_name in phenoData must include sample names. Since MiRaGE package tries to compare two distinct states, you need at least a set of gene expression corresponding to each of them. In gene_exp data, we have two biological replicates of negative control and the results one day after treatment. Thus, the 1st and 2nd columns of expression data are named as neg.1 and neg.2, respectively (This means "negative control 1" and "negative control 2", respectively). The 3rd and 4th columns of expression data corresponds to the two biological replicates one day after the treatment. Thus, they are names as day1.1 and day1.2, respectively.

These column names which express distinct samples keep some flexibilities but must have the form group.n, where group corresponds to either of sample groups and n must be integer starting from 1. This means, it you have N biological replicates for the first group (typically it includes un-treated or negative control samples) names as groupAand M for the second group (typically it includes treated samples) names as groupB, data structure of ExpressioSet which stores target gene expression is,

sample_name: groupA.1 groupA.1 ... groupA.N groupB.1 groupB.2 ... groupB.M gene_id : gene1,gene2,gene3....

gene_id is much easier. It can includes any of gene id which can be treated by MiRaGE. They can be a mixture of the different types of gene ids. In this case, only gene expression having gene id specified when MiRaGE is called are treated as target genes.

The easiest way to generate ExpressionSet which include target gene expression may be importing files including gene expression using standard R function,

> x_gene <- read.csv(system.file("extdata/x_all_7a.csv",package="MiRaGE"),sep="\t") > x_gene[101:103,]

gene neg.1 neg.2 day1.1 day1.2 101 BG167701 7.09 7.63 7.3 8.42 102 NM_001029863 995.00 2090.00 669.0 1370.00 103 NM_001014445 2540.00 6570.00 2070.0 3270.00

As can be seen, the first column includes gene id, which is "refseq" here, and the second to the fifth columns include gene expression. Data frame **x_gene** can be transformed to ExpressionSet objects **gene_exp** as

```
> gene_exp <- new("ExpressionSet",expr=data.matrix(x_gene[,-1]))
> fData(gene_exp)[["gene_id"]] <- x_gene[,1]
> pData(gene_exp)[["sample_name"]] <- colnames(x_gene)[-1]</pre>
```

For users' convenience, we have places a file $x_all_7a.csv$ under csv directory. Please refer to this file for the preparation of files including target gene expression.

4.2 Output: *P*-values

As mentioned in the above, output of MiRaGE is a list which includes two dataframes named as P0 and P1, respectively. P0 includes the rejection probabilities that the target gene expression in the first sample group is less than that in the second group. This means, smaller P-values indicate the target gene expression in the first sample group is more likely less than the second sample groups. Inversely, P1 includes the rejection probabilities that the target gene expression in the second sample is less than that in the first group. Thus, smaller P-values indicate target gene expression in the second group is more likely less than the first groups.

```
> result$P1[1:3,]
```

```
Refseqmixed1hsa-let-7a-5p8.678355e-262hsa-let-7b-5p8.678355e-263hsa-let-7c-5p8.678355e-26
```

In the above, we have shown the first three lines in the dataframe result\$P1. Since these are small, target genes of these three miRNAs is possibly expressive in the second group. In the first column of result\$P0 and result\$P1, names of considered miRNAs are listed. The number of miRNAs considered varys dependent upon the argument conv of MiRaGE. The second column includes *P*-values attributed to each miRNA. Dependent upon argument method, the number of columns which store *P*-values may change (see below).

5 Example

5.1 Example1: non-differentiated vs differentiated ES cell

In this section, we demonstrate how to infer target gene regulation via MiRaGE. First we import data set from experimental package humanStemCell

```
> require(humanStemCell)
> data(fhesc)
```

In this data set, human stem cells were assayed using Affymetrix 133plus 2 arrays. There were six arrays, three were biological replicates for undifferentiated cells, the other three were biological replicates for differentiated cells. In order to analyze this set, we modify ExpressionSet **fhesc** as

```
> pData(fhesc)[["sample_name"]] <- c("neg.1","neg.2","neg.3",</pre>
```

```
+ "pos.1","pos.2","pos.3")
```

```
> fData(fhesc)[["gene_id"]] <-rownames(exprs(fhesc))</pre>
```

Then, first three are designated as non-differentiated ES cell and the later three are differentiated ES cell. Obtaining P-values is easy,

```
> require(MiRaGE)
> result <- MiRaGE(fhesc,species="HS",ID="affy_hg_u133a_2")</pre>
```

Using the results, we can list miRNAs whose target genes are upregulated in the later (i.e., differentiated ES cell) group with P-values.

```
> result$P0[order(result$P0[,2])[1:5],]
```

```
Refseqmixed9hsa-miR-17-5p1.022976e-2113hsa-miR-20a-5p1.022976e-2128hsa-miR-93-5p1.022976e-2136hsa-miR-106a-5p1.022976e-21113hsa-miR-106b-5p1.022976e-21
```

Since miRNAs are believed to suppress target genes, these miRNAs are supposed to be upregulated in the former (i.e., non-differentiated ES cell) group.

5.2 Example 2: Universal Human Reference RNA vs brain

In this section, we demonstrate how to infer target gene regulation via MiRaGE using another example.

First we import data set from experimental package beadarrayExampleData

> require(beadarrayExampleData)

```
> data(exampleBLData)
```

> data(exampleSummaryData)

The data in this package are a subset of the MAQC bead-level data available in the beadarrayUseCases package. Bead-level refers to the availability of intensity and location information for each bead on each BeadArray in an experiment. In this dataset, BeadArrays were hybridized with either Universal Human Reference RNA (UHRR, Stratagene) or Brain Reference RNA (Ambion) as used in MAQC project. This object is a representation of the bead-level data for 2 arrays and was created by the beadarray package.

Since this is two color array, and the number of columns of expression must be the number of columns of expression data MUST be the length of sample_name, we omit later half of samples and employ only the first twelve samples, for simplicity.

```
> vv <- exampleSummaryData[,1:12]</pre>
```

```
> fData(vv)[["gene_id"]] <- fData(exampleSummaryData)[["IlluminaID"]]</pre>
```

```
> pData(vv)[["sample_name"]] <- c("neg.1","neg.2","neg.3","neg.4",</pre>
```

```
+ "neg.5", "neg.6", "brain.1", "brain.2", "brain.3", "brain.4", "brain.5", "brain.6")
```

```
> result <- MiRaGE(vv,species="HS",ID="illumina_humanwg_6_v3")</pre>
```

Then we can list miRNAs whose target genes are upregulated in negative control, i.e., miRNAs which are expected to be upregulated in brain as follows.

```
> result$P1[order(result$P1[,2])[1:5],]
```

Refseqmixed82hsa-miR-124-3p0.0001196863157hsa-miR-506-3p0.0001196863143hsa-miR-451a0.012822081299hsa-miR-191-5p0.1640405989102hsa-miR-126-3p0.4025617836

6 Rapid use & Off line use

Although the default value of location is "local", when location="web", MiRaGE every time tries to access MiRaGE Server¹ to download target gene tables, gene id conversion table, and miRNA conservation table. It is a time consuming process. Especially, since the target gene table is huge, it may take a few minutes. It may not be often to use MiRaGE iteratively many times, we offer the method to avoid "every time download".

6.1 Suppressing downloading

In MiRaGE, we offer the option to suppress downloading. If you repeatedly use MiRaGE with keeping either species, ID, or conv unchanged, you can suppress time consuming download process by specifying either species_force, ID_force, or conv_force as FALSE (Defaults for these are TRUE).

<u>Caution</u> Do not omit the arguments either species, ID, or conv if they differ from defaults, even if they are not modified during iterative usage and either species_force, ID_force, or conv_force is FALSE. They are used for other purposes than specifying what should be downloaded.

6.2 Save & load tables

More advanced and convenient way is to save the objects storing target gene tables, gene id conversion table, and miRNA conservation table. The names of objects are,

- TBL2 : Target gene tables
- id_conv : Gene id conversion table
- conv_id : MiRNA conservation table

¹http://www.granular.com/DATA2/

Thus, for example TBL2 is saved as

> save(file="TBL2",TBL2)

you can use it later by loading as

```
> load("TBL2")
```

Then you can execute MiRaGE with specifying species_force=F as

```
> result <- MiRaGE(...,species_force=F)</pre>
```

Now, you can skip time consuming download processes for the target gene table. Similar procedures are possible for id_conv and conv_id, too. Execute MiRaGE, save downloaded tables, and use the tables later by loading them when these arguments take same values.

6.3 miRNATarget package

One can also install experimental package miRNATarget instead of the usage of web. Once you install experimental package miRNATarget, you will never be required to access to internet.

```
> library(MiRaGE)
```

```
> data(gene_exp)
```

```
> library(Biobase)
```

```
> result <- MiRaGE(gene_exp,species="HS")</pre>
```

6.4 Generation of tables from scratch

I have also prepared functions which generate TBL2, id_conv and conv_id from scratch. Usually, user do not need them since prepared tables can be obtained from the web or as experimental package as mentioned above.

TBL2_HS can be saved in the current directly by executing

```
> TBL2_HS_gen()
```

and TBL2_MM can saved in the current directly by executing

```
> TBL2_MM_gen()
```

id_conv for mouse can saved in the current directly by executing

and id_conv for human can saved in the current directly by executing

> id_conv_gen(SP="HS")

HS_conv_id can saved in the current directly by executing

> HS_conv_id()

and MM_conv_id can saved in the current directly by executing

> MM_conv_id()

However, basically, execution of some of them are very time consuming. It is highly discouraged to build tables from scratch. It is much better to use prepared tables.

7 Multiple comparison correction

Obtained *P*-values are definitely underestimated, i.e., even if P < 0.05, this does not mean the rejection probability is less than 0.05. If one prefers to use adjusted *P*-values, we recommend to use p.adjust with parameter of BH, as

> p.adjust(result\$P1[,2],method="BH")

```
[1] 1.467431e-24 1.467431e-24 1.467431e-24 1.467431e-24 1.467431e-24
 [6] 1.467431e-24 7.420103e-01 7.420103e-01 9.953000e-01 9.953000e-01
[11] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[16] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[21] 9.953000e-01 9.953000e-01 9.953000e-01 3.114959e-01 9.953000e-01
[26] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 1.467431e-24
[31] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[36] 9.953000e-01 9.953000e-01 9.953000e-01 4.124658e-09 9.953000e-01
[41] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[46] 1.875348e-01 7.037504e-01 9.364901e-01 9.364901e-01 9.953000e-01
[51] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.364901e-01
[56] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[61] 9.953000e-01 9.953000e-01 9.953000e-01 9.364901e-01 9.953000e-01
[66] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[71] 9.953000e-01 9.953000e-01 9.953000e-01 1.467431e-24 1.467431e-24
[76] 9.953000e-01 7.420103e-01 9.953000e-01 9.953000e-01 9.953000e-01
[81] 9.953000e-01 7.591141e-01 9.953000e-01 9.953000e-01 9.953000e-01
[86] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[91] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[96] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[101] 9.953000e-01 9.953000e-01 9.953000e-01 6.556531e-01 8.396576e-01
[106] 9.953000e-01 9.953000e-01 5.944977e-01 7.420103e-01 9.953000e-01
[111] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[116] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[121] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
```

```
[126] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[131] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[136] 4.124658e-09 7.420103e-01 9.953000e-01 9.953000e-01 9.953000e-01
[141] 9.953000e-01 9.953000e-01 9.953000e-01 4.766116e-02 9.953000e-01
[146] 9.953000e-01 7.420103e-01 9.953000e-01 9.953000e-01 9.953000e-01
[151] 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[156] 9.953000e-01 7.591141e-01 7.420103e-01 5.857503e-01 9.953000e-01
[161] 5.857503e-01 9.953000e-01 9.953000e-01 9.953000e-01 9.953000e-01
[166] 9.953000e-01 9.953000e-01 9.364901e-01 9.953000e-01 9.953000e-01
[171] 9.953000e-01 9.953000e-01 9.364901e-01 9.953000e-01 9.953000e-01
[176] 9.953000e-01 9.953000e-01 9.364901e-01 9.953000e-01 9.953000e-01
[181] 1.467431e-24 9.953000e-01 1.467431e-24 9.953000e-01 9.953000e-01
[186] 9.953000e-01
```

Then we can see which *P*-values are really significant, e.g., less than 0.05. In addition to this, it will allow us to evaluate which miRNAs really regulate target genes, e.g.,

> result\$P1[,1][p.adjust(result\$P1[,2],method="BH")<0.05]</pre>

```
[1] hsa-let-7a-5p hsa-let-7b-5p hsa-let-7c-5p hsa-let-7d-5p
[5] hsa-let-7e-5p hsa-let-7f-5p hsa-miR-98-5p hsa-miR-196a-5p
[9] hsa-let-7g-5p hsa-let-7i-5p hsa-miR-196b-5p hsa-miR-490-3p
[13] hsa-miR-4458 hsa-miR-4500
186 Levels: hsa-let-7a-5p hsa-let-7b-5p hsa-let-7c-5p ... hsa-miR-99b-5p
```

 x_gene is the transfection expreiments of let-7a, it is reasonable that only a few miRNAs including let-7a have significant *P*-values.

Appendix:Arguments

Functionality of MiRaGE changes dependent upon the values of arguments. In this section, we will try to explain how the functionality of MiRaGE changes.

species

This specifies target species. Considered miRNAs are based upon miRBase². Rel. 20. At the moment, supported species are human ("HS") and mouse ("MM"). MiRaGE downloads corresponding target gene table (named TBL2) from MiRaGE Server. Default is "MM".

²http://www.mirbase.org

ID

This specifies gene ID. Default is "refseq". If ID is not "refseq", MiRaGE downloads corresponding gene id conversion table (called ID) from RefSeq to specified gene ID from MiRaGE Server. Supported gene IDs are,

	com	mon
	ID	description
1	ensembl_gene_id	Ensembl Gene ID
2	$ensembl_transcript_id$	Ensembl Transcript ID
3	ensembl_peptide_id	Ensembl Protein ID
4	ensembl_exon_id	Ensembl Exon ID
5	ccds	CCDS ID
6	embl	EMBL (Genbank) ID
7	entrezgene	EntrezGene ID
8	merops	MEROPS ID
9	pdb	PDB ID
10	protein_id	Protein (Genbank) ID
11	refseq_peptide	RefSeq Protein ID [e.g. NP_001005353]
12	rfam	Rfam ID
13	$rfam_transcript_name$	Rfam transcript name
14	ucsc	UCSC ID
15	unigene	Unigene ID
16	uniprot_sptrembl	UniProt/TrEMBL Accession
17	uniprot_swissprot	UniProt/SwissProt ID
18	$uniprot_swissprot_accession$	UniProt/SwissProt Accession
19	uniprot_genename	UniProt Gene Name
20	$uniprot_genename_transcript_name$	Uniprot Genename Transcript Name
21	wikigene_name	WikiGene Name
22	wikigene_id	WikiGene ID
23	efg_agilent_sureprint_g3_ge_8x60k	Agilent SurePrint G3 GE 8x60k probe
24	$efg_agilent_wholegenome_4x44k_v1$	Agilent WholeGenome 4x44k v1 probe
25	$efg_agilent_wholegenome_4x44k_v2$	Agilent WholeGenome 4x44k v2 probe
26	codelink	Codelink probe
27	phalanx_onearray	Phalanx OneArray probe
28	smart	SMART ID
29	pfam	PFAM ID
30	tigrfam	TIGRFam ID
31	interpro	Interpro ID

	human		
	ID	description	
1	hgnc_id	HGNC ID(s)	
2	hgnc_symbol	HGNC symbol	
3	$hgnc_transcript_name$	HGNC transcript name	
4	affy_hc_g110	Affy HC G110 probeset	
5	affy_hg_focus	Affy HG FOCUS probeset	
6	affy_hg_u133_plus_2	Affy HG U133-PLUS-2 probeset	
7	affy_hg_u133a_2	Affy HG U133A_2 probeset	
8	affy_hg_u133a	Affy HG U133A probeset	
9	affy_hg_u133b	Affy HG U133B probeset	
10	affy_hg_u95av2	Affy HG U95AV2 probeset	
11	affy_hg_u95b	Affy HG U95B probeset	
12	affy_hg_u95c	Affy HG U95C probeset	
13	affy_hg_u95d	Affy HG U95D probeset	
14	affy_hg_u95e	Affy HG U95E probeset	
15	affy_hg_u95a	Affy HG U95A probeset	
16	affy_hugenefl	Affy HuGene FL probeset	
17	$affy_huex_1_0_st_v2$	Affy HuEx 1_0 st v2 probeset	
18	$affy_hugene_1_0_st_v1$	Affy HuGene 1_0 st v1 probeset	
19	affy_u133_x3p	Affy U133 X3P probeset	
20	$agilent_cgh_44b$	Agilent CGH 44b probe	
21	illumina_humanwg_6_v1	Illumina HumanWG 6 v1 probe	
22	illumina_humanwg_6_v2	Illumina HumanWG 6 v2 probe	
23	illumina_humanwg_6_v3	Illumina HumanWG 6 v3 probe	
24	illumina_humanht_12	Illumina Human HT 12 probe	

	mouse		
ID	description		
1	fantom	Fantom ID	
2	ipi	IPI ID	
3	mgi_id	MGI ID	
4	mgi_symbol	MGI symbol	
5	$mgi_transcript_name$	MGI transcript name	
6	affy_mg_u74a	Affy mg u74a probeset	
7	affy_mg_u74av2	Affy mg u74av2 probeset	
8	affy_mg_u74b	Affy mg u74b probeset	
9	$affy_mg_u74bv2$	Affy mg u 74 bv 2 probeset	
10	$affy_mg_u74c$	Affy mg u74c probeset	
11	$affy_mg_u74cv2$	Affy mg u74cv2 probeset	
12	affy_moe430a	Affy moe430a probeset	
13	$affy_moe430b$	Affy moe430b probeset	
14	$affy_moex_1_0_st_v1$	Affy MoEx probeset	
15	$affy_mogene_1_0_st_v1$	Affy MoGene probeset	
16	$affy_mouse 430_2$	Affy mouse430 2 probeset	
17	$affy_mouse430a_2$	Affy mouse430a 2 probeset	
18	affy_mu11ksuba	Affy mu11ksuba probeset	
19	affy_mu11ksubb	Affy mu11ksubb probeset	
20	illumina_mousewg_6_v1	Illumina MouseWG 6 v1 probe	
21	illumina_mousewg_6_v2	Illumina MouseWG 6 v2 probe	

Requirements for supporting any other gene IDs are welcomed.

method

This specifies how to treat replicates. if **method** is "mean", then averaged gene expression is attributed to each gene. If it is "mixed", they are used for statistical test as it is. This means, the number of target genes attributed to each miRNAs is as many as the number of replicates. If "one_by_one" is specified, all of combinations between the two groups, i.e.,

```
groupA.1 \times groupB.1, groupA.1 \times groupB.2, ..., groupA.N \times groupB.M.
```

are condicred. Thus, in this case, both P0 and P1 have $1 + N \times M$ columns, the later $N \times M$ includes *P*-value for each of combinations. Default is "mean".

\mathbf{test}

This specifies the statititical methods to evaluate significance of reglation of target genes. Supported are "ks" (Kolmogorov-Smirnov test), "t" (t-test), and "wilcox" (Wilcoxon test). These are performed by standard R functions, ks.test, t.test, and wilcox.test, respectively. Default is "ks".

conv

This specifies how well considered miRNAs must be conserved. Supported are "conserved", "weak_conserv" and "all". Baed upon TargetScan 7.2 ³, they correspond to broadly conserved, conserved, and others. For more detail, plese colusult with TargetScan. Default is "conserved".

Force download or not

species_force, ID_force, and conv_force spefify if target gene table, gene id conversion table, and miRNA conservation table are forced to be donwloaded. Dafult is T. If some of them have already been downloded and one would like to use it as it is, please specify they are F.

References

- [1] Y-h Taguchi, Jun Yasuda, 2010, Inference of Gene Expression Regulation via microRNA Transfection, ICIC2010, Proceedings, Springer, 6215, 672-679.
- [2] Y-h Taguchi, Jun Yasuda, 2012, MiRaGE: Inference of Gene Expression Regulation via MicroRNA Transfection II, ICIC2011, Proceedings, Springer, 6840,192-135
- [3] M. Yoshizawa, Y-h. Taguchi, Jun Yasuda, 2011, Inference of Gene Regulation via miRNAs During ES Cell Differentiation Using MiRaGE Method, Int. J. Mol. Sci., 12[12]:9265-9276
- [4] Taguchi, Y-h. (2013). Inference of Target Gene Regulation by miRNA via Mi-RaGE Server. Introduction to Genetics - DNA Methylation, Histone Modification and Gene Regulation. ISBN: 978-1477554-94-4. iConcept Press. Retrieved from http://www.iconceptpress.com/books/ IntroductionToGeneticsDNAMethylationHistoneModificationAndGeneRegulation/

³http://www.targetscan.org