# Processed human microRNA-overexpression data from GEO, and sequence information from TargetScan, and targetScore from TargetScore

Yue Li yueli@cs.toronto.edu

April 26, 2017

## **1** MicroRNA perturbation datasets

We collected 84 Gene Expression Omnibus (GEO) series corresponding to 6 platforms, 77 human cells or tissues, and 112 distinct miRNAs. To our knowledge, this is by far the largest miRNA-overexpression data compendium. To automate the data download and processing, we developed a pipeline written in R, making use of the function getGEO from *GEOquery* R/Bioconductor package (Davis and Meltzer [2007]). For each dataset, the pipeline downloads the raw or processed data (if available) and calculates (when necessary) the log fold-change (logFC) in treatment (miRNA transfected) vs (mock) control, taking into account the unique properties of each data. Next, we combined all of the logFC data columns into a single  $N \times M$  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.knn from *impute* R package (Troyanskaya et al. [2001]). For miRNA transfection data having multiple measurements (in different studies), we picked the one whose logFC correlate the most with the validated targets from mirTarBase Hsu et al. [2011] or average them if no validated target available.

```
$`GEO Series`
```

```
[1] 84
$Platform
[1] 6
$`Cell/Tissue`
[1] 77
```

# 2 TargetScan context score and PCT

TargetScan context score and PCT for all of the predicted sites (including conserved and nonconserved sites) downloaded from TargetScan website (http://www.targetscan.org/cgi-bin/ targetscan/data\_download.cgi?db=vert\_61)

```
> targetScanCS <- get_TargetScanHuman_contextScore()
> targetScanPCT <- get_TargetScanHuman_PCT()</pre>
```

```
> head(targetScanCS)
```

	Gene Symbol	Transcript ID	miRNA	3prime	pairing	local AU	positio
1	A1CF		nsa-miR-4711-3p				
2	A1CF	NM_138933 h	nsa-miR-4711-3p		-0.018	-0.095	-0.108
3	A1CF	NM_014576 h	nsa-miR-4711-3p		-0.018	-0.095	-0.108
4	A1CF	NM_001198820 h	nsa-miR-4711-3p		-0.018	-0.095	-0.108
5	Alcf	NM_001198819 h	nsa-miR-4711-3p		-0.018	-0.095	-0.108
6	Alcf	NM_001198818 h	nsa-miR-4711-3p		-0.018	-0.095	-0.108
	TA SPS	context+ score	context+ score	percent	tile		
1	0.003 0.017	-0.448			99		
2	0.003 0.017	-0.448			99		
3	0.003 0.017	-0.448			99		
4	0.003 0.017	-0.448			99		
5	0.003 0.017	-0.448			99		
6	0.003 0.017	-0.448			99		
> dim(targetScanCS)							
[1] 9569357 10							
> head(targetScanPCT)							
	miI	R Family Gene Sy	wmbol Transcript	ID PO	CT		
1			=	786 0.0			
2		c/23b-3p	A1BG NM_130	786 0.0	00		
7		297/4465		786 0.0	00		
8				786 0.0	00		
9	miR-103a/10	)7/107ab	A1BG NM_130	786 0.0	00		
10	miR-103a/10	)7/107ab	A1BG NM_130	786 0.0	)9		
			2				

SLC45A1 -0.018655797

#### **3** TargetScore

Encouraged by the superior performance of TargetScore (manuscript in peer-review), we applied TargetScore to all of the transfection data above. For further exploring miRNA targetome and their associations, we enclose the targetScores results in this package.

```
> targetScoreMatrix <- get_precomputed_targetScores()</pre>
> head(names(targetScoreMatrix))
[1] "hsa-miR-34b"
                    "hsa-miR-34c" "hsa-miR-205" "hsa-miR-124" "hsa-miR-1"
[6] "hsa-miR-181a"
> head(targetScoreMatrix[[1]])
               logFC targetScanCS targetScanPCT targetScore
SGIP1
         0.077526011
                              0.00
                                                   0.03489650
                                                0
AGBL4
         0.020639084
                              0.00
                                                0
                                                   0.03388637
NECAP2
         0.078650400
                              0.00
                                                0
                                                   0.03492518
CLIC4
         0.016043400
                             -0.03
                                                   0.24335149
                                                0
        -0.002303429
                              0.00
                                                   0.03417828
ADC
                                                0
```

0.03457975

0

We can reproduce targetScores using the above data as demonstrated in the following example (require *TargetScore* package). As a convenience function, we applied a wrapper function called getTargetScores that does the following: (1) given a miRNA ID, obtain fold-change(s) from logFC.imputed matrix or use the user-supplied fold-changes; (2) retrives TargetScan context score (CS) and PCT (if found); (3) obtain validated targets from the local mirTarBase file; (4) compute targetScore. We apply getTargetScores function using miRNA hsa-miR-1, which we know has all three types of data, namely logFC, targetScan context score, and PCT.

0.00

```
> library(TargetScore)
> library(gplots)
> myTargetScores <- getTargetScores("hsa-miR-1", tol=1e-3, maxiter=200)
> table((myTargetScores$targetScore > 0.1), myTargetScores$validated) # a v
> # obtain all of targetScore for all of the 112 miRNA
>
> logFC.imputed <- get_precomputed_logFC()
> mirIDs <- unique(colnames(logFC.imputed))
>
> # takes time
>
# targetScoreMatrix <- mclapply(mirIDs, getTargetScores)
>
> # names(targetScoreMatrix) <- mirIDs</pre>
```

## 4 Session Info

```
> sessionInfo()
R version 3.4.0 (2017-04-21)
Platform: x86_64-pc-linux-qnu (64-bit)
Running under: Ubuntu 16.04.2 LTS
Matrix products: default
BLAS: /home/biocbuild/bbs-3.5-bioc/R/lib/libRblas.so
LAPACK: /home/biocbuild/bbs-3.5-bioc/R/lib/libRlapack.so
locale:
 [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
 [3] LC TIME=en US.UTF-8
                                LC COLLATE=C
 [5] LC_MONETARY=en_US.UTF-8
                                LC MESSAGES=en US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                                LC_NAME=C
 [9] LC_ADDRESS=C
                                LC_TELEPHONE=C
[11] LC MEASUREMENT=en US.UTF-8 LC IDENTIFICATION=C
attached base packages:
[1] stats
              graphics
                       grDevices utils datasets methods
                                                                 base
other attached packages:
[1] TargetScoreData_1.12.0
loaded via a namespace (and not attached):
[1] compiler_3.4.0 tools_3.4.0
```

### References

- Sean Davis and Paul S Meltzer. GEOquery: a bridge between the Gene Expression Omnibus (GEO) and BioConductor. *Bioinformatics (Oxford, England)*, 23(14):1846–1847, July 2007.
- Sheng-Da Hsu, Feng-Mao Lin, Wei-Yun Wu, Chao Liang, Wei-Chih Huang, Wen-Ling Chan, Wen-Ting Tsai, Goun-Zhou Chen, Chia-Jung Lee, Chih-Min Chiu, Chia-Hung Chien, Ming-Chia Wu, Chi-Ying Huang, Ann-Ping Tsou, and Hsien-Da Huang. miRTarBase: a database curates experimentally validated microRNA-target interactions. *Nucleic acids research*, 39 (Database issue):D163–9, January 2011.
- O Troyanskaya, M Cantor, G Sherlock, P Brown, T Hastie, R Tibshirani, D Botstein, and R B Altman. Missing value estimation methods for DNA microarrays. *Bioinformatics (Oxford, England)*, 17(6):520–525, June 2001.