# miRGator v2.0 : an integrated system for functional investigation of microRNAs

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## ABSTRACT

miRGator is an integrated database of microRNA (miRNA)-associated gene expression, target prediction, disease association and genomic annotation, which aims to facilitate functional investigation of miRNAs. The recent version of miRGator v2.0 contains information about (i) human miRNA expression profiles under various experimental conditions, (ii) paired expression profiles of both mRNAs and miRNAs. (iii) gene expression profiles under miRNA-perturbation (e.g. miRNA knockout and (iv) known/predicted overexpression), miRNA targets and (v) miRNA-disease associations. In total, >8000 miRNA expression profiles, ~300 miRNA-perturbed gene expression profiles and  $\sim$ 2000 mRNA expression profiles are compiled with manually curated annotations on disease, tissue type and perturbation. By integrating these data sets, a series of novel associations (miRNAmiRNA, miRNA-disease and miRNA-target) is extracted via shared features. For example, differentially expressed genes (DEGs) after miRNA knockout were systematically compared against miRNA targets. Likewise, differentially expressed were miRNAs (DEmiRs) compared with disease-associated miRNAs. Additionally. miRNA profiles expression and disease-phenotype revealed miRNA pairs whose expression was regulated in parallel in various experimental and disease conditions. Complex associations are readily accessible using an interactive network visualization interface. The miRGator v2.0 serves as a reference database to investigate miRNA expression and function (http://miRGator.kobic.re.kr).

# INTRODUCTION

MicroRNAs (miRNAs) are important posttranscriptional regulators that are associated with various cell functions and human diseases. MiRNAs bind to complementary sequences in the 3' UTRs of target mRNAs resulting in negative regulation of gene expression (1,2). Significant efforts have been made to study the function of miRNAs and their target genes. However, the function of most miRNAs are still elusive and under active investigation. A number of miRNA-related databases have been developed. There are many miRNA target prediction algorithms available including TargetScan (3), PITA (4), miRanda (5), PicTar (6), miBridge (7) and many more. PhenomiR (8) and miR2Disease (9) provide information on diseaseassociated miRNAs from literature curation. FAME (10) uses a target prediction method to infer biological processes affected by human miRNAs. MMIA (11) and MAGIA (12) provide tools for gene (or miRNA) set analysis and target prediction using miRNA-miRNA expression profiles.

MiRGator is an integrated database of miRNAassociated gene expression, target prediction, disease association and genomic annotation, which aims to facilitate functional investigation of miRNAs. In this update version, we integrate a variety of publicly available data sets related to miRNA biology. Overall, more than 10000 miRNA/mRNA expression profiles are compiled and organized by manual curation according to miRNA, tissue type, disease and perturbation. MiRGator contains information about (i) human miRNA expression profiles under various experimental conditions, (ii) paired expression profiles of both mRNAs and miRNAs from the same sample, (iii) gene expression profiles under miRNAperturbation (e.g. miRNA knockout and overexpression), (iv) known/predicted miRNA targets and (v) miRNAdisease associations. In addition, a number of new

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The authors wish it to be known that, in their opinion, the first three authors should be regarded as joint First Authors.

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This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/ by-nc/2.5), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. features have been added to the previous version, such as association analysis among miRNAs, diseases and perturbations and an integrated network viewer. Here, an overview of the miRGator system, the collection and processing of data sets, and important features are briefly described. A more detailed usage and documentation are provided on-line (http://miRGator.kobic.re.kr).

### SYSTEM OVERVIEW

The miRGator v2.0 consists of three main modules: the data set browser, the association analysis and the gene set analysis (GSA)/miRNA set analysis (miRSA) modules. The overall system of miRGator v2.0 is shown in Figure 1. Each of these three modules has its distinct function but they are tightly integrated to facilitate functional investigation of miRNAs. Particularly, the expression data sets are manually annotated according to miRNA, target, tissue type, disease and perturbation, so that the user can easily focus on the subject of interest.

The data set browser provides diverse access routes to our comprehensive miRNA-related data sets by miRNA, tissue type, etc. The data are hierarchically organized in several tables with increasing details from top to bottom, e.g. from the full list of miRNAs to the actual expression values of each miRNA. The association analysis connects different miRNAs, diseases and perturbations by the overlap among miRNA/gene sets and by the similarity between miRNA expression/phenotype profiles. The association analysis module is also equipped with an integrated association network viewer for convenient exploration of association networks. Finally, the GSA/ miRSA module takes user-defined gene sets or miRNA sets and performs GSA or miRSA against various types of gene annotations [e.g. gene ontology (GO, 13), KEGG (14)] as well as the miRGator data sets.

#### DATA SOURCES AND PROCESSING

The core data set of miRGator v2.0 consists of seven different data types collected from various sources such as gene expression, miRNA-disease association and four major miRNA-target databases (TargetScan, PITA, miRanda and miBridge). The seven data types belong to one of the three broad categories miRNA set, gene set and miRNA profiles. Where appropriate, these data sets were



**Figure 1.** System overview of miRGator v2.0. There are three main modules. The data set browser allows access to all the data sets by miRNA, tissue, disease and perturbation. Within the association analysis module, miRNAs, diseases and perturbations can be associated by GSA, miRSA and miRPA. The resulting association network is visualized by an integrated association network viewer. In the GSA/miRSA module, the user may enter a list of miRNAs/genes and perform GSA/miRSA against miRNA sets or gene sets in the miRGator system.



Figure 2. The overall procedure of data collection and processing.

manually curated according to tissue, disease and perturbation (e.g. chemical treatment). The overall procedure of data collection and processing is shown in Figure 2.

The miRNA set category is basically a list of miRNAs. This category consists of known disease-miRNA associations from PhenomiR (8) and differentially expressed miRNAs (DEmiRs) from the gene expression omnibus (GEO) (15) and ArrayExpress (16). For DEmiRs, we collected miRNA expression data for various cancers and perturbations. In total, 154 miRNA expression data sets (8431 profiles) were downloaded from the GEO and ArrayExpress. The miRNA expression profiles were quantile normalized using LIMMA package (17). The miRNA profiles were hand-curated, resulting in the annotations of 26 human tissue types, 56 diseases and 83 perturbations.

The gene set category contains information for miRNA targets such as predicted miRNA targets, anti-coexpressed miRNA-gene pairs and miRNA-perturbed differentially expressed genes (DEGs) (i.e. the up-regulated genes under miRNA-knockout conditions and down-regulated genes under miRNA-overexpression conditions). The predicted miRNA targets were collected from TargetScan, PITA, miRanda and miBridge. The miRNA-perturbed gene expression profiles were obtained from GEO. We compiled 68 data sets of 613 gene expression profiles, where one or more miRNAs were knocked out/down or overexpressed. After filtering out inappropriate sets, the remaining 63 sets contain 595 profiles annotated with 88 miRNA perturbations (90 different miRNAs), 15 tissues and 18 diseases. For anti-coexpressed gene sets, we collected 22 data sets from GEO, where both miRNA and mRNA expression profiles were measured for the same

sample. The total number of the paired miRNA and mRNA profiles was 1265 and 1568, respectively. Pearson correlation between miRNA and mRNA expression values was calculated for each of the 22 data sets. Accordingly, a miRNA-mRNA pair may have up to 22 correlation values. We compiled anti-coexpressed miRNA-mRNA pairs, which show significant anti-correlation in at least two or more cases among the 22 data sets at *P*-value cut off <0.001. For miRNAs with more than 500 anti-correlated mRNAs above cut-off, we took only the top 500 mRNAs ranked by the sum of -log (P-value).

The last category is miRNA-related profile, which consists of miRNA expression profiles and phenotypic profiles. The miRNA expression profiles came from the same data used in the DEmiR set. Phenotypic profiles were obtained from the PhenomiR data describing up/ down regulation of miRNAs in various diseases. Additionally, various types of gene annotation information are integrated including GO and biological pathways (KEGG, BioCarta). The details of data source, type and statistics are shown in Table 1 and on-line.

### DATA SET BROWSER

From the data set browser, the user can access all the data sets complied in miRGator v2.0, which are organized according to miRNA, target, tissue type, disease and perturbation. Accordingly, the user may access the data set of interest using various annotation categories of choice. Several tables are sequentially displayed from top to bottom with increasing detail as the user chooses an item of interest. For example, the user can start from

Data category	Data type	Description	Experiments (Profiles)	miRNAs	mRNAs	Diseases	Tissues	Perturbations	Data source
miRNA set	Disease- miRNAs	Disease associated	_	354 (up) 340 (down)	_	59	25	_	PhenomiR
	DEmiRs	DEmiRs under diseases/ perturbations	146 (8013)	689	_	47	21	128	GEO ArrayExpress
Gene set	miRNA targets	Predicted miRNA target genes	_	700	19069	-	_	_	miBridge, TargetScan, PITA, miRanda
	miRNA-	miRNA knockout	5 (92)	_	15708	4	4	16	GEO
	perturbed DEGs	miRNA overexpression	20 (205)	_	20887	12	12	27	GEO
	Coexpressed genes	Positive and negative coexpression between miRNA and mRNA from paired expression profiles	22 (2538)	685	27 830	11	13	_	GEO
miRNA profiles	Expression profiles	miRNA expression profiles under various conditions	146 (8013)	689	_	47	21	128	GEO ArrayExpress
	Phenotype profiles	Profiles of disease associations for each miRNA	_	354 (up) 340 (down)	_	59	25	_	PhenomiR

Table 1. Summary statistics of data sets in miRGator v2.0

the list of all miRNAs, click to show the list of miRNA profiles containing the selected miRNA, and then choose a specific miRNA experiment, where the actual expression values are displayed as a heatmap. MiRNA–disease association, and tissue-specific or disease related expression data can be accessed in a similar manner.

Particularly, miRGator v2.0 is useful to obtain integrated information on miRNA target reliability. The miRNA-target relationship may be supported by multiple, independent evidences such as target prediction, up-regulation on miRNA knockout, down-regulation on miRNA overexpression and miRNA-mRNA anticoexpression from paired expression experiments. These data supporting the miRNA-target relation are shown side-by-side simultaneously. For example, RAC1 (Entrez Gene ID: 5879) is down-regulated by overexpression of hsa-mir-155 in the GSE14477 data set. RAC1 is predicted as a target by two methods (PITA and miRanda) and also shows a strong negative correlation in two out of the 22 miRNA-mRNA paired expression data sets. This strongly suggests that RAC1 is a genuine target of hsa-mir-155, which needs further experimental validation.

# ASSOCIATION ANALYSIS AND ASSOCIATION NETWORK VIEWER

The data sets in miRGator belong to one of the three categories of gene set, miRNA set and miRNA-related profile. Within the data types of the same category, association relationships are established by the overlap among the gene/miRNA sets or by the similarity of profiles (Figure 3). For example, miRNA-perturbed DEGs can be linked to miRNA targets by GSA. Similarly, differentially expressed miRNAs (DEmiRs) can be associated to

disease-related miRNAs. In miRNA profile analysis (miRPA), a pair of miRNAs may be functionally related if their expression is regulated in parallel in various experimental and disease conditions as is the case with a pair of genes. Likewise, a pair of miRNAs with similar phenotypic or disease-related patterns is likely to be functionally associated. The resulting associations can be visualized by the association network viewer (bottom right in Figure 1). The network viewer is equipped with useful features such as save network, export image, highlighting nodes interactively, group nodes by disease, etc. and a detailed description is provided in the on-line documentation.

### miRSA AND GSA

The GSA/miRSA module allows the user to enter a list of genes and perform GSA/miRSA for statistical enrichment. Association between two sets of miRNAs is also possible in a similar manner to GSA, which we call miRSA. Most representative gene IDs are cross-referenced and automatically converted to the miRBase ID or NCBI entrez gene IDs. For miRSA, we use our own compilation of miRNA sets (i.e. disease-associated miRNAs and DEmiRs; data type A and B in Figure 2) in addition to the PhenomiR database. The GSA is applied for the predicted miRNA targets (data type C), miRNA-perturbed DEGs (data type D), anti-coexpressed genes (data type E). Additional categories include the GO, disease gene [Online Mendelian Inheritance in Man (OMIM, 18) and genetic association database (GAD, 19)], and pathway (KEGG and BioCarta) analyses. The user may enter a list of genes or miRNAs to check any relevance to miRNA biology or existing annotations. On log-in, the



Figure 3. The three different types of associations in the association analysis module. Depending on the data category, (A) miRSA, (B) GSA and (C) miRPA.

user-defined gene/miRNA sets may be saved on the system, which can be retrieved later for further analysis.

## CONCLUSION

The miRGator v2.0 integrates human miRNA-related expression patterns under various disease and perturbation conditions. The system is designed to facilitate functional investigation of miRNAs by integrating various evidences of miRNA targets and by revealing non-canonical associations among miRNAs, diseases, perturbations and user-defined gene/miRNA sets.

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