RWRMTN: a tool for predicting disease-associated microRNAs based on a microRNA-target gene network

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Additional file 1

User Manual & Case studies

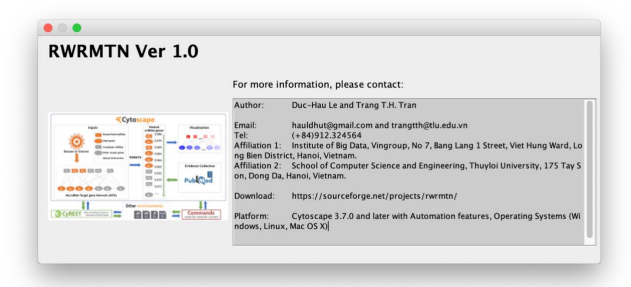


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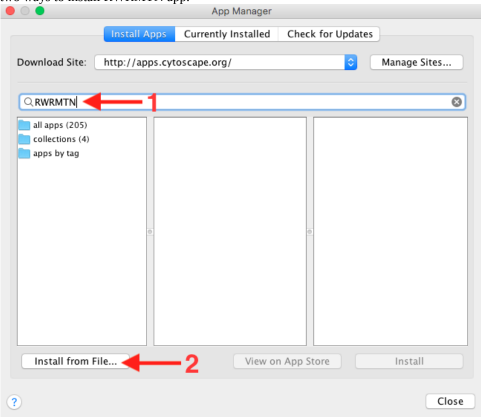
I. Setup

1. Install Cytoscape

- RWRMTN1.0 can only run on **Cytoscape 3.6 (or later)** platform, **which has Automation features**, therefore user should download this version at http://cytoscape.org/
- Cytoscape need JRE to run, therefore download JRE version 8.x or later from http://www.oracle.com/technetwork/java/index.html and install it.
- Install Cytoscape to the root folder (e.g., /Applications/Cytoscape_v3.6.0).

2. Install RWRMTN app

There are two ways to install RWRMTN app.

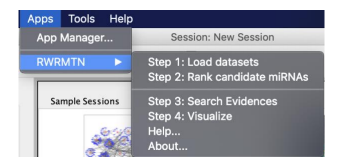


- Method 1: Automatically install from Cytoscape Appstore: Select menu App → AppManager in Cytoscape. Then type RWRMTN in search box to install directly from Appstore of Cytoscape.
- O Method 2: Manual install:
 - Download RWRMTN_v1.0.jar file from https://sourceforge.net/projects/rwrmtn/
 - Then, install it by going to **Apps** → **App Manager**.... After that, choose **Install from file...**, then browse the downloaded RWRMTN_v1.0.jar file.

Note that: RWRMTN_v1.0 can work on Windows, Ubuntu and Mac OS. The following manual was prepared when running RWRMTN on Mac OS.

II. Overview of RWRMTN

After installing, RWRMTN will be automatically loaded in the App menu of Cytoscape



The main tasks (Rank candidate miRNAs, Evidence Search and Visualization) of RWRMTN are completed after four steps:

- Step 1: Load data sets (miRNA-target gene interactions and known disease-miRNA associations)
- **Step 2**: Rank candidate miRNAs (including 4 sub-steps)
 - 1. Select a disease of interest
 - 2. Input candidate miRNAs to rank
 - 3. Parameters setting (for advanced users)
 - o 4. Rank
- **Step 3**: Search Evidences
- Step 4: Visualize

These steps can be performed, and results of each step can be exposed

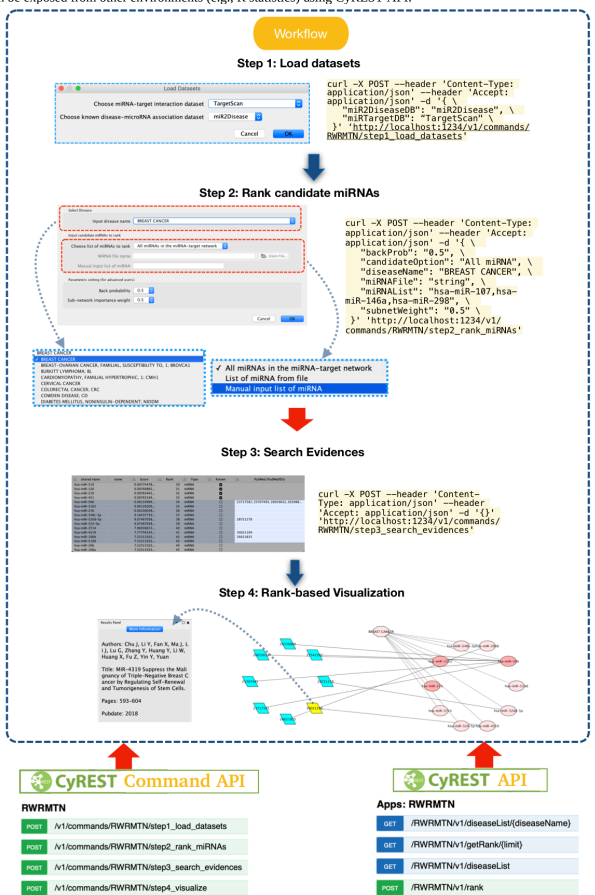
- Using Cytoscape menu

Beside the Cytoscape GUI, new upgraded automation feature of Cytoscape allows functions of Cytoscape and apps called via REST API. Therefore, we can call the functions of Cytoscape and apps in workflows in other environments such as R, Python, etc... Thus, RWRMTN functions can be used by

- CyREST Command API
- Cyrest Api

III. Case study: Prediction of breast cancer-associated miRNAs

In this section, we demonstrate the use of RWRMTN in predicting novel breast cancer-associated miRNAs by following workflow. The workflow can be done using Cytoscape menu or CyREST Command API. In addition, intermediate results can be exposed from other environments (e.g., R statistics) using CyREST API.



1. Run RWRMTN using Cytoscape menu and CyREST Command API

Step 1: Load datasets

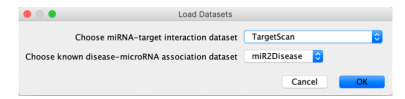
Load datasets for miRNA-target interaction network and known disease-miRNA associations:

- miRNA-target gene interaction dataset (*miRTargetDB*): choose built-in dataset **TargetScan or miRWalk** or your own dataset to build miRNA-target interaction network.
- Known disease-miRNA association dataset (miR2DiseaseDB): choose built-in dataset miR2Disease or HMDD.

Step 1 can be performed by two ways:

- Cytoscape menu: Apps →RWRMTN→ Step 1: Load datasets

Here, a miRNA-target interaction dataset TargetScan (Lewis, et al., 2003) and a known disease-miRNA association dataset miR2Disease (Jiang, et al., 2009) were used.



To build miRNA-target interaction network and load known disease-miRNA associations:

- 1. Choose miRNA-target interaction dataset.
- 2. Choose known disease-microRNA association dataset
- 3. Click **OK** to load datasets.

- CyREST command API: Help → Automation → CyREST Command API. Choose RWRMTN in the SwaggerUI.

 RWRMTN
 Show/Hide List Operations | Expand Operations |

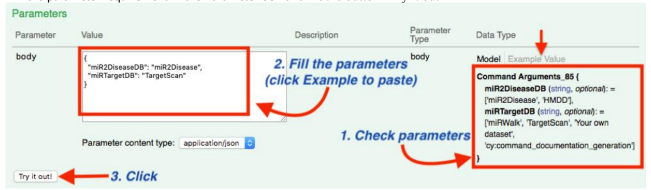
 POST
 /v1/commands/RWRMTN/step1_load_datasets
 Step 1: Load Datasets

 POST
 /v1/commands/RWRMTN/step2_rank_miRNAs
 Step 2: Rank candidate miRNAs

 POST
 /v1/commands/RWRMTN/step3_search_evidences
 Step 3: Search Evidences

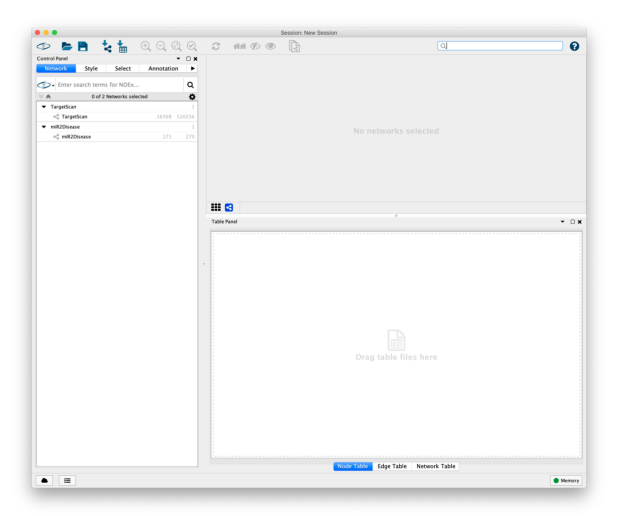
 POST
 /v1/commands/RWRMTN/step4_visualize
 Step 4: Visualize

Fill the parameter requirement in the Parameter box and hit the button "Try it out"



(For more details on using automation features (CyREST API and CyREST Command API) of Cytoscape, visit the site: https://github.com/cytoscape/cytoscape-automation/wiki/App-Developers:-Cytoscape-Command-Best-Practices)

- This will load corresponding datasets into Network tab of Cytoscape (see following figure).



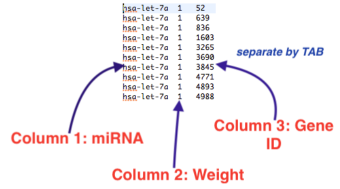
Note: a miRNA-target interaction dataset TargetScan (Lewis, et al., 2003) and a known disease-miRNA association dataset miR2Disease (Jiang, et al., 2009) were used.

For CyREST command API, return successful message:

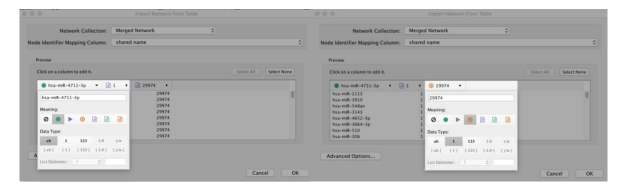
```
{
    "data": {
        "message": "Load Heterogeneous Network successfully"
     },
        "errors": []
}
```

Note that:

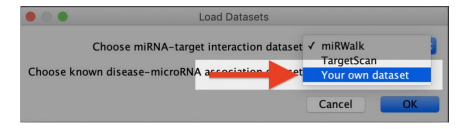
- For miRNA-target datasets: We pre-installed 2databases miRWalk (database of experimentally validated miRNA-target interactions) and TargetScan (a dataset containing predicted miRNA-target interactions). If you want to use your own dataset, it must follow the following format, and be imported into Cytoscape beforehand.



Your own dataset can be imported into Cytoscape by clicking menu $File \rightarrow Import \rightarrow Network \rightarrow File$. Set column 1 as source node and column 3 as target node.



Then, it will appear in miRNA network option:



For known disease-miRNAs dataset: We pre-installed 2 datasets miR2Disease and HMDD (an up-to-date human disease-miRNA association database).

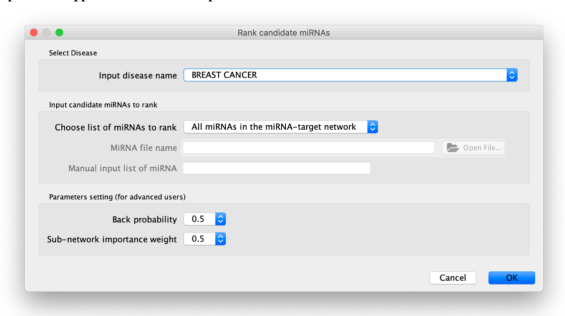
Step 2: Rank candidate miRNAs

This step includes 4 sub-steps:

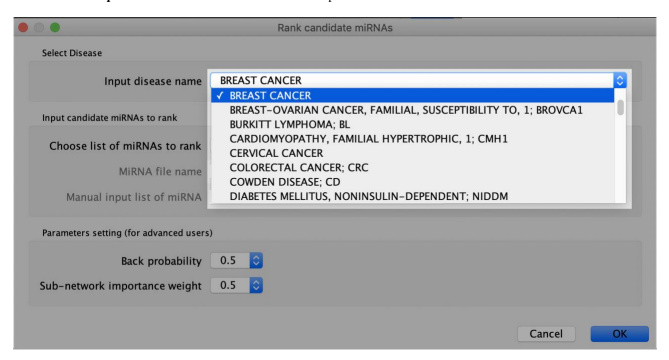
- Select a disease: For example, Breast cancer (OMIM ID: 114480) is selected.
- Input candidate miRNAs to rank: there are 3 options to choose
 - o All miRNAs in the miRNA-target network
 - o List of miRNAs from file: input the file.
 - Manual input list of miRNAs.
- Parameters setting (for advanced users)
 - o Back probability (default setting is 0.5)
 - o Sub-network importance weight (default setting is 0.5)
- Rank

Step 2 can be performed by two ways:

Cytoscape menu: Apps →RWRMTN→ Step 2: Rank candidate miRNAs



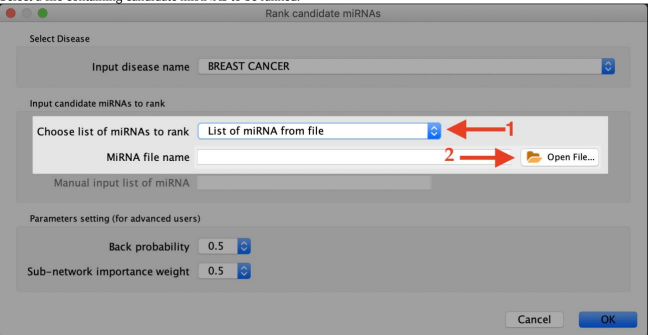
Step 2.1: Choose Disease name from the dropdown list:



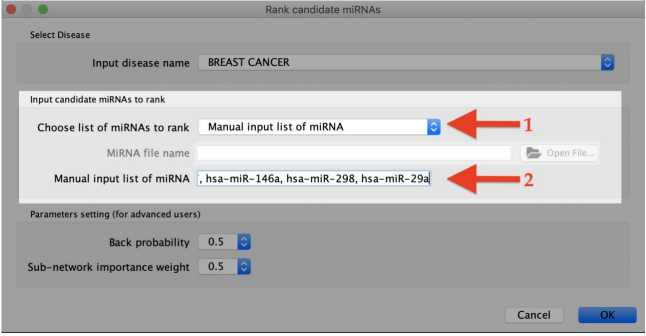
o **Step 2.2**: Input candidate miRNAs to rank:

Default setting is "All miRNAs in the network". If you want to choose specific list of miRNAs, you can select other options.

Select a file containing candidate miRNAs to be ranked.



Directly input list of candidate miRNAs

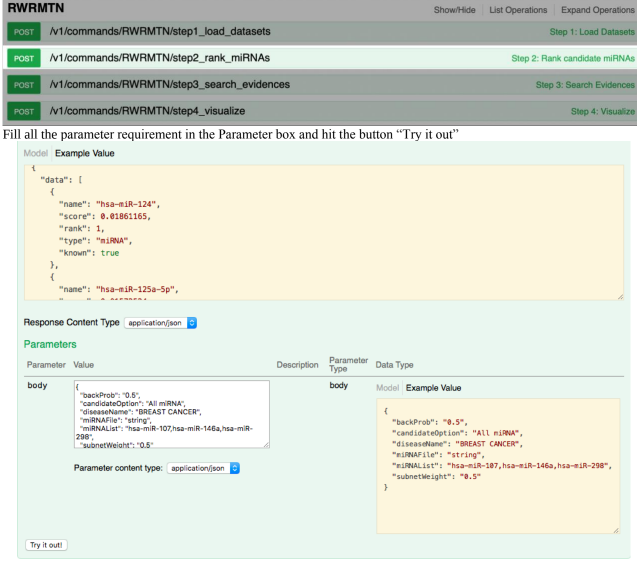


Step 2.3: Parameters setting (for advanced users)

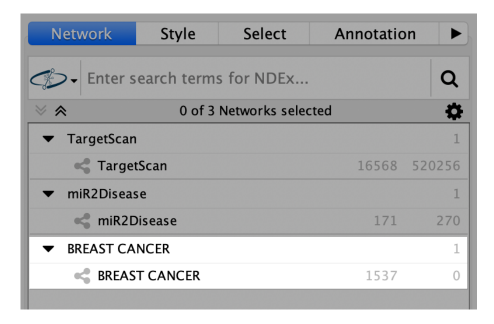
Choose to set Back probability and Sub-network importance weight in case of using different values other than default values.

o Step 2.4: Rank (Click OK)

- CyREST command API: Help → Automation → CyREST Command API



Step 2 will generate list of ranked miRNAs displayed in a network named by the selected disease (i.e. BREAST CANCER)



The following table lists known miRNAs associated with the selected diseases and the inputted candidate miRNAs

🚠 shared name	name 🚠 Score	🚣 Rank	# Type	- Known
hsa-miR-124	0.00912531	1 1	miRNA	lacksquare
hsa-miR-27a	0.00857614	4 2	miRNA	
hsa-miR-128	0.00840957	7 3	miRNA	lacksquare
hsa-miR-200c	0.00838510	0 4	miRNA	
hsa-miR-200b	0.00838510	D 5	miRNA	
hsa-miR-429	0.00838123	3 6	miRNA	lacksquare
hsa-miR-125b	0.00835355	5 7	miRNA	lacksquare
hsa-miR-125a-5p	0.00835355	5 8	miRNA	lacksquare
hsa-miR-17	0.00832807	7 9	miRNA	lacksquare
hsa-miR-20a	0.00832807	7 10	miRNA	
hsa-let-7a	0.00826798	3 11	miRNA	
hsa-miR-206	0.00818675	5 12	miRNA	
hsa-miR-200a	0.00817063	3 13	miRNA	
hsa-miR-141	0.00817063	3 14	miRNA	
hsa-miR-204	0.00814230	D 15	miRNA	
hsa-miR-373	0.00805362	2 16	miRNA	
hsa-miR-516a-3p	0.00803928	3 17	miRNA	
hsa-miR-155	0.00802905	5 18	miRNA	
hsa-miR-7	0.00794792	2 19	miRNA	
hsa-miR-205	0.00794064	4 20	miRNA	
hsa-miR-221	0.00790413	3 21	miRNA	
hsa-miR-222	0.00790413	3 22	miRNA	
hsa-miR-127-5p	0.00790080	D 23	miRNA	
hsa-miR-335	0.00786437	7 24	miRNA	
hsa-miR-146a	0.00784430	D 25	miRNA	
hsa-miR-146b-5p	0.00784430	D 26	miRNA	
hsa-miR-21	0.00781843	3 27	miRNA	
hsa-miR-520c-5p	0.00778787	7 28	miRNA	
hsa-miR-10b	0.00778371	1 29	miRNA	
hsa-miR-510	0.00774478	30	miRNA	
hsa-miR-126	0.00766892	2 31	miRNA	
hsa-miR-210	0.00765442	2 32	miRNA	lacksquare
hsa-miR-451	0.00765194	4 33	miRNA	lacksquare
hsa-miR-506	0.00154994	4 34	miRNA	
hsa-miR-3163	0.00126209	9 35	miRNA	
hsa-miR-27b	0.00100038	36	miRNA	
hsa-miR-548c-3p	9.16437739	9 37	miRNA	
hsa-miR-520d-5p	9.07467058	3 38	miRNA	
hsa-miR-524-5p	9.07467058	3 39	miRNA	
hsa-miR-3714	7.96936672	2 40	miRNA	
hsa-miR-4319	7.77794334	4 41	miRNA	
hsa-miR-106b	7.52313163	3 42	miRNA	
hsa-miR-519d	7.52313163	3 43	miRNA	

And JSON result is exposed in CyREST Command API

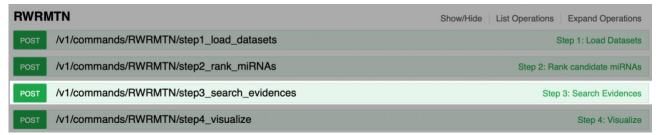
```
Request URL
 http://localhost:1234/v1/commands/RWRMTN/step2_rank_miRNAs
Response Body
  {
    "data": [
      {
        "name": "hsa-miR-124",
        "score": 0.009125315577015921,
        "rank": 1,
        "type": "miRNA",
        "known": true
        "name": "hsa-miR-27a",
        "score": 0.008576144025852219,
        "rank": 2,
        "type": "miRNA",
        "known": true
      {
        "name": "hsa-miR-128",
        "score": 0.008409577922964604,
        "rank": 3,
Response Code
 200
```

Step 3: Search Evidences

By selecting option "All miRNAs in the miRNA-target network" in Step 2, all miRNAs in the selected miRNA-target gene network are ranked and displayed in Node Table of the network (in Network Tab of Cytoscape) which has the same name as the disease of interest (i.e., BREAST CANCER).

To find the evidences in literature (PubMed), you need to select highly ranked miRNAs by highlighting rows in the network (i.e., BREAST CANCER) and choose between these two ways:

- Cytoscape menu: Select menu Apps → RWRMTN → Step 3: Search Evidences
- Cyrest command API: Help \rightarrow Automation \rightarrow Cyrest Command API.



Hit the button "Try it out" without any parameters (remember to highlight rows first).

<u>Note</u>: If you receive the alert that none row is selected. You need to show the column "selected" in Cytoscape table and set value to **true** once to trigger the function "selected" column to work.

The result of Step 3 is a list of PubMed IDs of the publications containing evidences about associations between selected miRNAs and the disease of interest displayed in . For more information (e.g., paper title, author list, publication date, etc..), refer to Step 4 or use command API of step 3.

& shared name	name	- Score		Type	# Known	PubMed (PudMedIDs)
hsa-miR-510		0.00774478	30	miRNA		
hsa-miR-126		0.00766892	31	miRNA		
hsa-miR-210		0.00765442	32	miRNA	□ □	
hsa-miR-451		0.00765194	33	miRNA		
hsa-miR-506		0.00154994	34	miRNA		23717581, 25707493, 26059632, 263988
hsa-miR-3163		0.00126209	35	miRNA		
hsa-miR-27b		0.00100038	36	miRNA		
hsa-miR-548c-3p		9.16437739	37	miRNA		
hsa-miR-520d-5p		9.07467058	38	miRNA		28721278
hsa-miR-524-5p		9.07467058	39	miRNA		
hsa-miR-3714		7.96936672	40	miRNA		
hsa-miR-4319		7.77794334	41	miRNA		30021199
hsa-miR-106b		7.52313163	42	miRNA		26621835
hsa-miR-519d		7.52313163	43	miRNA		
hsa-miR-20b		7.52313163	44	miRNA		
hsa-miR-106a		7.52313163	45	miRNA		

This result is exposed by using CyREST Command API

```
Request URL
 http://localhost:1234/v1/commands/RWRMTN/step3_search_evidences
Response Body
     "data": [
         "miRnaName": "hsa-miR-520d-5p",
         "PubMedIds": [
           "28721278"
         1.
         "info": {
           "28721278": {
             "pubdate": "2016",
             "authors": [
               "Ishihara Y",
               "Tsuno S",
               "Ping B",
               "Ashizaki T"
               "Nakashima M",
               "Miura K",
               "Miura Y",
               "Yamashita T",
               "Hasegawa J",
Response Code
```

As can be seen from the above screenshot, the result is returned as an array of JSON objects. Each object includes 3 pairs of key/value:

- "miRnaName" is the selected miRNA.
- "PubMedIds" lists the ids found in the PubMed which provided evidences of associations between selected miRNA and the disease of interest.
- "info" shows the detail information of each PubMed ID including publication date, authors, title and pages, which then can be seen by visualization functions of RWRMTN.

In this case study, four of ten highly ranked miRNAs are provided with evidences.

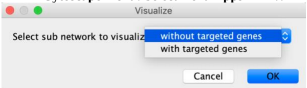
- "hsa-miR-506" supported by five studies (PubMed IDs: 23717581, 25707493, 26059632, 26398880 and 27542202). The study 23717581 showed that has-miR-506 regulates epithelial mesenchymal transition in breast cancer cell lines. Meanwhile, the study 26059632 proved notable inhibition of hsa-miR-506 over-expression to proliferation and metastasis of breast cancer cells. In addition, study 26398880 indicated that mechanism underlying miRNA-506 is a contributing factor in breast carcinogenesis (has-miR-506 was proven to be a tumor suppressor).
- "hsa-miR-520d-5p" supported by a study with PubMed ID 28721278. More specifically, it was reported that this miRNA upregulates the activation of BRCA1 (breast cancer 1, early onset) in the DNA repair process 35 days after transfection.
- "hsa-miR-4319" was showed in study PubMed ID 30021199 as a suppressor of the malignancy of triple-negative breast cancer by regulating self-renewal and tumorigenesis of stem cells.
- "has-miR-106b" was proven by the experiment carried on patient samples and cell lines in the study (PubMed ID 26621835)

Step 4: Visualize

Selected miRNAs can be visualized in a network based on the rankings. In addition, target genes, the disease of interest and detail information of PubMed IDs collected from Step 3 such as *paper title*, *author list*, *journal name* can be displayed aside in this network.

Choose between two options for visualization (Remember to highlight selected rows first)

Cytoscape menu: Select menu Apps → RWRMTN → Step 4: Visualize



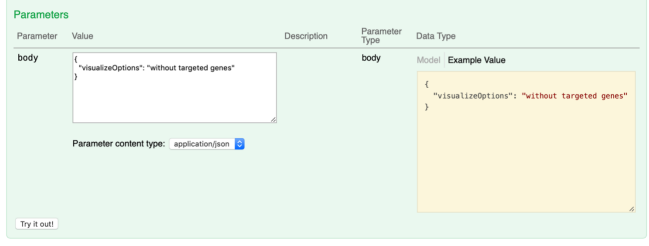
Visualization options:

- 1. Visualize without targeted genes of selected miRNAs
- 2. Visualize with targeted genes of selected miRNAs
- 3. Click **OK** to visualize.

CyREST command API: Help \rightarrow Automation \rightarrow CyREST Command API



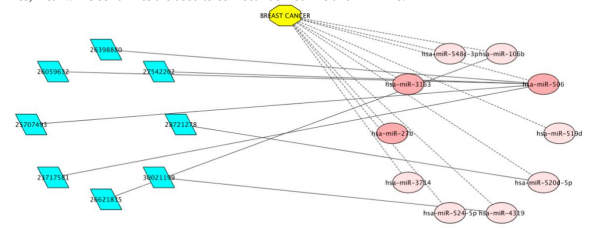
Fill the parameter requirement in the Parameter box and hit the button "Try it out"



Step 4 will create network view in two cases:

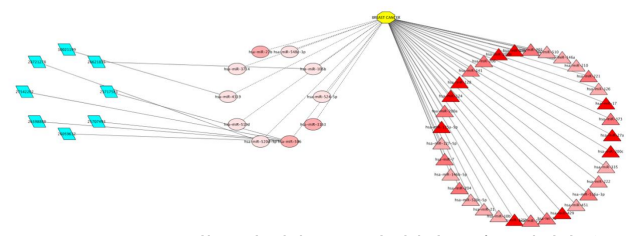
o Without targeted genes

Here, we selected top 10 ranked candidate miRNAs and then visualized based on their rankings. In addition, evidences (PubMed IDs) for each selected miRNAs were also visualized. Disease and candidate miRNAs are connected by long-dash lines, meanwhile solid lines are used to connect PubMed IDs and miRNAs.



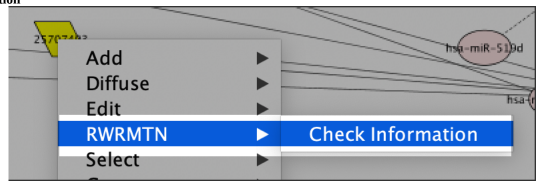
(Note: PubMed IDs, disease, and miRNAs are represented in parallelogram, octagon and ellipse shapes, respectively)

One may want to view the relationships with miRNAs known to be associated with the disease of interest (i.e., BREAST CANCER). Thus, we can additionally select known miRNAs to visualize

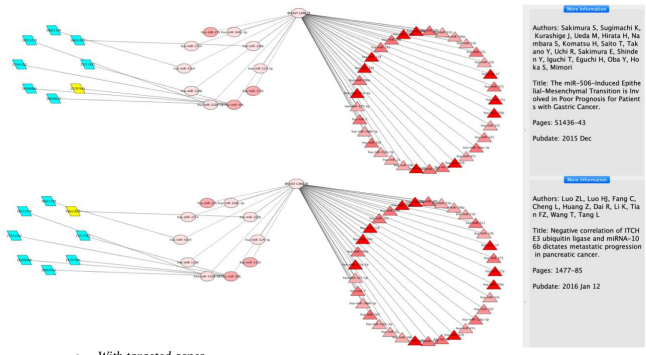


(Note: Known miRNAs are represented by triangles which are connected with the disease of interest by dot lines).

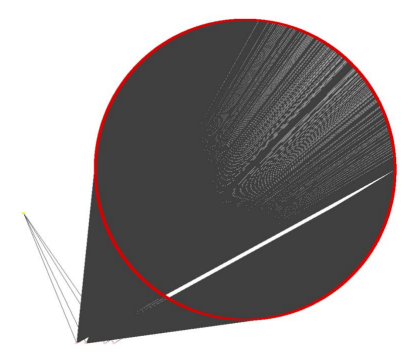
For detail information of each PubMed ID, right click on PubMed ID node and select menu **RWRMTN** → **Check Information**



The East panel will appear to display detail information of the selected PubMed ID.



• With targeted genes Select this option if user wants to see the relationships between the selected miRNAs and their target genes.



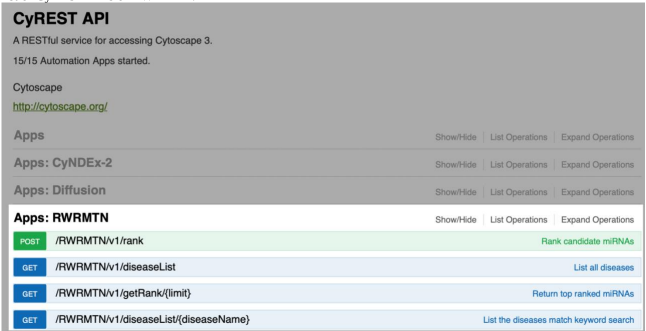
(Note: Each miRNA often targets to many genes)

2. Run RWRMTN by calling CyREST API

In this section, we first introduce some developed CyREST APIs which provides some helpful functions. Second, we demonstrate of their use in a workflow in R statistic environment.

Overview of CyREST APIs

To reveal all APIs, select menu **Help → Automation → CyREST API** to open Swagger UI of CyREST API. Here is the list of CyREST APIs of RWRMTN.



The detail of each API is available in Swagger Documentation. Here is short description.

GET /RWRMTN/v1/diseaseList List all diseases

This API returns list of all diseases (OMIM ID and disease name) available in the selected known disease-miRNA association database (i.e., miR2Disease). Based on this list, users can select a disease of interest.

```
Response Body

[
{
    "diseaseID": "MIM104300",
    "diseaseName": "ALZHEIMER DISEASE; AD"
},
    {
    "diseaseID": "MIM109800",
    "diseaseName": "BLADDER CANCER"
},
    {
    "diseaseName": "BURKITT LYMPHOMA; BL"
},
    {
    "diseaseID": "MIM114480",
    "diseaseName": "BREAST CANCER"
},
    {
    "diseaseName": "BREAST CANCER"
},
    {
    "diseaseID": "MIM114500",
    "diseaseID": "MIM114500",
    "diseaseName": "COLORECTAL CANCER; CRC"
```

GET /RWRMTN/v1/diseaseList/{diseaseName}

List the diseases match keyword search

This API provides a list of diseases whose names match the query parameter (e.g., cancer). This API help user narrow down list of diseases to the disease of interest (e.g., disease ID MIM114480 for BREAST CANCER). For example:

```
Request URL
 http://localhost:1234/RWRMTN/v1/diseaseList/cancer
Response Body
   [
       "diseaseID": "MIM109800",
       "diseaseName": "BLADDER CANCER"
       "diseaseID": "MIM114480",
      "diseaseName": "BREAST CANCER"
    },
       "diseaseID": "MIM114500",
      "diseaseName": "COLORECTAL CANCER; CRC"
    },
     {
      "diseaseID": "MIM133239",
       "diseaseName": "ESOPHAGEAL CANCER"
       "diseaseID": "MIM137215",
       "diseaseName": "GASTRIC CANCER, HEREDITARY DIFFUSE; HDGC"
```

POST /RWRMTN/v1/rank

Rank candidate miRNAs

This API will use RWRMTN to rank candidate miRNAs and return the result in JSON format. The request is POST request with input parameters as follows:

required:
Disease OMIM ID: String
List of miRNA: String

optional:
miRTargetDB: String
miR2DiseaseDB: String
backProb: String
subnetWeight: String

```
Model Example Value

RankParameters {
    diseaseOMIMID (string): Disease OMIM ID,
    listOfmiRNAs (string): List of miRNA to rank,
    miRTargetDB (string, optional):
    MicroRNA Dataset,
    miRZDiseaseDB (string, optional):
    Disease-miRNA Dataset,
    backProb (number, optional): Backprobability,
    subnetWeight (number, optional): Subnetwork importance weight
}
```

```
## Model Example Value

{
    "diseaseOMIMID": "MIM114480",
    "listOfmiRNAs": "All miRNAs",
    "miRTargetDB": "TargetScan",
    "miR2DiseaseBB": "miR2Disease",
    "backProb": 0.5,
    "subnetWeight": 0.5
}
```

The result is:

```
Request URL
 http://localhost:1234/RWRMTN/v1/rank
Response Body
      "rnaName": "hsa-miR-124",
      "rnaScore": 0.009125315577015921,
      "rnaRank": 1,
      "type": "miRNA",
      "known": true
      "rnaName": "hsa-miR-27a",
      "rnaScore": 0.008576144025852219,
       "rnaRank": 2,
       "type": "miRNA",
      "known": true
      "rnaName": "hsa-miR-128",
       "rnaScore": 0.008409577922964604,
      "rnaRank": 3,
      "type": "miRNA",
Response Code
 200
```

GET /RWRMTN/v1/getRank/{limit}

Return top ranked miRNAs

This API returns top ranked miRNAs by setting {limit} parameter.

For example: The following query returns top 10 ranked miRNAs

```
Request URL
 http://localhost:1234/RWRMTN/v1/getRank/10
Response Body
   [
       "rnaName": "hsa-miR-124",
      "rnaScore": 0.009125315577015921,
       "rnaRank": 1,
       "type": "miRNA",
       "known": true
     },
       "rnaName": "hsa-miR-27a",
       "rnaScore": 0.008576144025852219,
       "rnaRank": 2,
       "type": "miRNA",
       "known": true
     }.
       "rnaName": "hsa-miR-128",
       "rnaScore": 0.008409577922964604,
       "rnaRank": 3,
       "type": "miRNA",
Response Code
 200
```

Using RWRMTN in a workflow in R environment

In this case study, we used a dataset GSE19783 from GEO (Enerly, et al., 2011), which was created using Agilent-019118 Human miRNA Microarray 2.0 G4470B platform (GPL8227) and Agilent-014850 Whole Human Genome Microarray 4x44K G4112F (GPL6480) (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE19783). The study characterizes breast cancer subtypes from joint analysis of high throughput miRNA (using GPL8227) and mRNA (using GPL6480) Data.

In this case study, we explored the 799 miRNAs that were differentially expressed between the 64 wild-type samples (WT) and 36 *TP53* mutant samples via a workflow in R environment using CyREST API.

Briefly, here is the workflow:

- 1. Download the datasets
- 2. Perform differential expression analysis with limma package
- 3. Select a list of miRNAs, whose differential expression between cases and controls is statistically significant, as candidates
- 4. Rank the candidate miRNAs by RWRMTN via a CyREST API using a miRNA-target interaction dataset miRWalk (Dweep, et al., 2011) and a known disease-miRNA association dataset HMDD (Li, et al., 2014).

Before running the analysis, make sure RWRMTN and necessary packages are installed and they are functional:

Please run Check_CytoscapeConnection_LibraryInstallation_RWRMTN.R (download at https://sourceforge.net/projects/rwrmtn/) for checking connection with Cytoscape and whether necessary packages are installed.

Then, run the following source code in R (the source code can be found in CaseStudy_Final.R (download at https://sourceforge.net/projects/rwrmtn/)

```
11. ### Load series and platform data from GEO
12. gset <- getGEO("GSE19783", GSEMatrix =TRUE, AnnotGPL=FALSE)</pre>
13. if (length(gset) > 1){
14. idx <- grep("GPL8227", attr(gset, "names"))</pre>
15. }else{
16.
    idx <- 1
17. }
18.
19. gset <- gset[[idx]]</pre>
21. # make proper column names to match toptable
22. fvarLabels(gset) <- make.names(fvarLabels(gset))</pre>
23.
24. TP53Status<-gset$`tp53 mutation status:ch1`
25. # labeling for all samples
26. gsms<-""
27.
28. sml<-c()
29. for(i in 1:length(TP53Status)){
30. if(TP53Status[i]=="Mut"){
31.
        gsms<-paste0(gsms,"1")</pre>
32.
        sml[i]<-"Group1"</pre>
33.
      }else{
34.
      gsms<-paste0(gsms,"0")</pre>
        sml[i]<-"Group0"</pre>
35.
36.
37. }
38.
39. #Retrieve Expression Data From ESets
40. ex <- exprs(gset)
41. ex[which(ex <= 0)] <- NaN
42. # log2 transform
43. exprs(gset) <- log2(ex)
45. ### Differential expression analysis with limma package
46. # set up the data and proceed with analysis
47. fl <- as.factor(sml)
48. gset$description <- fl
49. #creates a design (or model) matrix
50. design <- model.matrix(~ description + 0, gset)</pre>
51. colnames(design) <- levels(fl)</pre>
52. fit <- lmFit(gset, design)</pre>
53. cont.matrix <- makeContrasts(Group1-Group0, levels=design)</p>
54. fit2 <- contrasts.fit(fit, cont.matrix)
55. fit2 <- eBayes(fit2, 0.01)
56. diffmiRNAlist <- topTable(fit2, adjust="fdr", number=nrow(fit2))</pre>
58. #Only select miRNAs whose differential expression between the two group (Group1 & Group0)
59. #is statistically significant (adj.P.val <=0.05) for ranking with RWRMTN
60. sigmiRNAlist <- subset(diffmiRNAlist,adj.P.Val<=0.05) #This returns 85 miRNAs
61. sigmiRNAlist <- subset(sigmiRNAlist, select=c("ID","adj.P.Val","P.Value"))</pre>
62. colnames(sigmiRNAlist)<-c("rnaName", "adj.P.Val", "P.Value")
63.
64. #Save statistically significant miRNAs standard output
65. write.table(sigmiRNAlist, file=stdout(), row.names=F, sep="\t")
66.
67.
68. ###Rank statistically significant miRNAs (candidate miRNAs) with RWRMTN
69. #Get miRNA list
70. lr<-sigmiRNAlist$rnaName
71. lor<-""
72. n<-length(lr)
73. for(i in 1:n){
74. lor<-paste(lor, lr[i],", ",sep='')
75.}
76.
77. #Select datasets (miRTargetDB, miR2DiseaseDB), the disease of interest (MIM114480: Breast
   cancer)
78. #and pass the candidata miRNAs list
79. login <- list(
```

```
80. diseaseOMIMID= "MIM114480", #OMIM ID of Breast cancer
81.
      listOfmiRNAs= lor,
82.
     miRTargetDB= "miRWalk",
83.
      miR2DiseaseDB= "HMDD"
84.)
85.
86. #Run Cytosacpe CyREST API
87. request_body_json <- toJSON(login)</pre>
88. res <- POST("http://localhost:1234/RWRMTN/v1/rank", body = login, encode="json")
89. y<-httr::content(res,"text", encoding = 'UTF-8')
90. get_prices_json <- fromJSON(y, flatten = TRUE)
91.
92. Output <- fromJSON((y))
93. #Remove miRNA with rank=0 (which are not available on the miRNA-target network)
94. rankedmiRNAlist <- Output[which(Output$rnaRank!=0),]
95. rankedmiRNAlist
96. write.csv(rankedmiRNAlist, file="rankedmiRNAlist.csv", row.names=F)
```

Here is the result:

```
rnaScore rnaRank type known
          rnaName
      hsa-miR-375 1.719388e-02 1 miRNA TRUE
2
      hsa-miR-107 4.059087e-03
                                     2 miRNA
                                              TRUE
3
      hsa-miR-15a 3.420814e-03
                                     3 miRNA
                                              TRUE
      hsa-miR-326 3.377178e-03
                                    4 miRNA
                                              TRUE
      hsa-miR-155 2.825815e-03
5
                                    5 miRNA
                                              TRUE
6
      hsa-miR-145 2.716448e-03
                                    6 miRNA
                                              TRUE
     hsa-miR-148b 2.686323e-03
                                    7 miRNA
                                              TRUE
                                   8 miRNA TRUE
9 miRNA TRUE
8
      hsa-miR-224 2.610611e-03
       hsa-let-7e 2.553925e-03
      hsa-miR-29c 2.545947e-03
hsa-miR-26b 2.467229e-03
10
                                   10 miRNA TRUE
11
                                    11 miRNA
                                              TRUE
                                   12 miRNA
      hsa-miR-30a 2.455210e-03
13
       hsa-let-7b 2.428472e-03
                                    13 miRNA
                                              TRUE
       hsa-let-7c 2.411595e-03
                                    14 miRNA TRUE
14
15
      hsa-miR-34b 2.396425e-03
                                   15 miRNA TRUE
      hsa-miR-10b 2.389018e-03
                                    16 miRNA
                                              TRUE
17
     hsa-miR-18a 2.352102e-03
                                    17 miRNA
                                              TRUE
                                    18 miRNA
18
      hsa-miR-328 2.344313e-03
                                              TRUE
19
      hsa-miR-143 2.331756e-03
                                    19 miRNA
                                             TRUE
2.0
      hsa-miR-214 2.313317e-03
                                   20 miRNA TRUE
                                    21 miRNA TRUE
21
      hsa-miR-152 2.283654e-03
     hsa-miR-135b 2.273221e-03
                                   22 miRNA TRUE
22
                                    23 miRNA TRUE
23
     hsa-miR-195 2.270405e-03
24 hsa-miR-125a-5p 7.286119e-04
                                    24 miRNA FALSE
25 hsa-miR-342-3p 5.469969e-04
                                   25 miRNA FALSE
       hsa-let-7a 4.714080e-04
                                    26 miRNA FALSE
2.6
27 hsa-miR-769-5p 4.352923e-04
                                    27 miRNA FALSE
28 hsa-miR-361-5p 3.802039e-04
                                   28 miRNA FALSE
  hsa-miR-142-3p 3.341059e-04
                                    29 miRNA FALSE
30 hsa-miR-34c-5p 2.341773e-04
                                   30 miRNA FALSE
                                    31 miRNA FALSE
31 hsa-miR-146b-5p 1.953439e-04
32
   hsa-miR-449a 1.556556e-04
                                    32 miRNA FALSE
33 hsa-miR-199a-5p 1.228826e-04
                                   33 miRNA FALSE
    hsa-miR-489 1.197518e-04
                                    34 miRNA FALSE
       hsa-miR-9 9.189293e-05
                                    35 miRNA FALSE
35
     hsa-miR-30a* 9.030027e-05
                                    36 miRNA FALSE
36
37
     hsa-miR-135a 8.225978e-05
                                    37 miRNA FALSE
                                   38 miRNA FALSE
    hsa-miR-181c 7.444931e-05
39
     hsa-miR-378* 7.208481e-05
                                    39 miRNA FALSE
     hsa-miR-30c 7.006724e-05
40
                                    40 miRNA FALSE
41
       hsa-let-7f 6.601124e-05
                                   41 miRNA FALSE
42 hsa-miR-483-3p 5.512170e-05
                                    42 miRNA FALSE
43 hsa-miR-342-5p 5.253228e-05
                                   43 miRNA FALSE
44 hsa-miR-501-3p 4.575108e-05
                                   44 miRNA FALSE
     hsa-miR-101 4.479886e-05
4.5
                                   45 miRNA FALSE
46 hsa-miR-574-3p 4.081944e-05
                                   46 miRNA FALSE
                                    47 miRNA FALSE
47
      hsa-miR-26a 3.510984e-05
48
      hsa-miR-103 3.103517e-05
                                   48 miRNA FALSE
49
      hsa-miR-378 2.822501e-05
                                    49 miRNA FALSE
50
      hsa-miR-99b 2.788255e-05
                                    50 miRNA FALSE
51 hsa-miR-590-5p 1.735067e-05
                                    51 miRNA FALSE
52 hsa-miR-362-5p 1.092474e-05
                                    52 miRNA FALSE
      hsa-miR-9* 2.842895e-06
5.3
                                    53 mirna false
54 hsa-miR-142-5p 1.520914e-06
                                    54 miRNA FALSE
55
     hsa-miR-101* 1.019211e-06
                                    55 miRNA FALSE
     hsa-miR-30e* 7.906392e-07
                                    56 miRNA FALSE
      hsa-miR-650 4.129138e-07
                                    57 miRNA FALSE
```

IV. Reference

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