'bimir' R package manual

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1. Introduction

'bimir' is an R package developed to generate biclusters of cell condition-specific microRNA targets from large mRNA transcriptome fold change (FC) matrix (a.k.a., FC table; it contains log2 FC values of 20,639 human genes under 5,158 experimental conditions). Besides the primary purpose of this package, it can be applied to create any constant biclusters allowing small number of noises using the progressive bicluster extension (PBE) algorithm. This package is downloadable from the Github repository (https://github.com/ysora/bimir) and will not be uploaded to CRAN or Bioconductor due to the large size of FC table (~140 MB)

2. Installation

- 1. Download the bimir source package ('bimir_1.0.tar.gz').
- 2. Install and load the package by typing following lines.
 - install.packages('devtools')
 - library(devtools)
 - install github('ysora/bimir')

3. Quick start with an example

Let's generate biclusters of 'hsa-miR-1-3p' where its target genes are up-regulated under common experimental conditions. To do this, type following lines.

- > FC = load FCtable() # To load FC table
- ➤ MP = getMIRprofile(miRNA = 'hsa-miR-1-3p', FCtable = FC, FCcutoff = log2(1.3)) # To generate MIR profile.
- > PBE MERGE(MIR profile = MP, mir = 'hsa-miR-1-3p', biclust.path = './', FCcutoff = log2(1.3))
- Then check the directory assigned to 'biclust.path' and see the biclusters (biclust_up_*.txt; fig. 1) and corresponding experimental condition (Experimental_condition_up_*.txt; fig. 2) and gene list (Targetlist_up_*.txt; fig. 3).
- **Down-bicluster:** To generate biclusters of targets commonly down-regulated under multiple conditions, just modify the 'FCcutoff' parameter to be negative value (e.g., -log2(1.3)).
- ➤ The available miRNAs are listed in the 'getmiRNAlist()' function.

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1		ANP32E		CHSY1		FAM72A	FN1		FNDC38	LYZ	NAMPT	PHLDAT		SULF1	TPM4	TWSG1	ZNF281
	10616_3	1.004				1.0		0.629	0.636					1.721			
	10714_6	1.096					48	-0.123	0.722					1.005		51 0.764	
	10714.7	1.116			2.159	1.4		-0.099	0.715				1.204	0.972			
	10791_1	0.777			2.18	0.7		-0.112	0.665					0.72			
6 P	11151,6	0.725		1.316		3.2		1.584	1.332					2.62			
	12606_2	1.806		2.713		2.1		1.478	2.241					2.22			
8 p	12606_3	1.791	1.96	2.482	1.953	2.4	76	1.667	2.878		4.635	0.509	3.11	1.86	0.3	16 1.64	
	13367_2	0.797	0.816	0.692	1.479	0.5	91	0.584	0.468	2.224	1.376	1.192	1.018	0.700	0.6	07 0.857	
	13367_5	2.073			5.01		78	1.212	1.197					3.18			
	14580_2	1.23		1.809	3.608	1.5		0.883	0.754				2.252	0.764			
	15471_1	0.846			2.749	1.		2.937	0.482					4.24			
	16515,1	0.961		0.648	0.922	1.7		1.905	0.452					2.380			
	16879_1	0.833			2.668	1.1		0.766	0.534					0.425			
	1,53481	1.965		0.805		1.6	64	2.93	0.65		2.997	2.724		1.38			
16 p	19728_2	0.715	0.445	1.145	0.043	0.9	14	1.132	0.399	0.775	1.589	2.101	0.826	1.14	0.4	17 1.466	
	19728_3	0.578	1.513	2.349	1.064	2.3	82	2.192	0.913	3.23	2.133	2.735	0.736	2.04	1.3	14 1.096	
18 p	19728_4	1.076	0.874	2.321	2.604	1.8	65	2.25	1.203	2.325	3.62	1.229		1.4	1.6	17 2.523	
19 p	25638_10	0.895			5.138	4.1		3.331	1,477			1.482		2.37			
	25638_11	0.568		0.505	4.8	3.		2.659	1.322					1.533			
	25638_9	-0.091		0.847	5.311	2.3		3.144	1.618					1.89		175 1.494	
	28914_2	0.611		0.609	1.159	0.9		0.409	1.302					1.64			
23 p	28914_3	0.733		0.587	1,523	0.6	75	1.235	1,198	1,943	1.684	0.734	1.458	2.56			
	29431_1	0.692	1,776	1.11	0.549	1,	67	1.916	0.435	1.8			1.081	2.44	1.2	18 0.693	
25 p	44337_1	0.425	0.189	7.046	5.595	5.3	09	3.371	1,404	2.651	0.718	0.774	0.975	1.81	1.1	69 1.297	0.98
26 p	7696_1	0.396	0.752	0.951	1.317	1.6	24	1.482	0.979					0.527	0.0	53 1.017	
27 p	7696,2	0.544	0.626	1.095	1.319	1.9	23	1.48	1.073	0.695	1.814	1.261	0.529	0.453	0.6	91 1.014	
28 p	8056_2	1.106	1.775	0.817	0.613	1.1	36	0.635	0.842	1.687	3.01	1.907	2.519	1.575	1.6	61 0.874	1.3
	8056_3	0.49		0.801	0.837	0.3		1.015	0.908					1.98			
	9600_4	2.976		2.498	1.325	0.4	66	2.42	1,584			2.963		1.1		99 3.134	
	9686,2	0.638		1.109	1.458	0.4	43	0.553	0.504	1,176	0.81	1.293	1.01	1.235	0.4	1.087	
32 p	9106,3	1.272	1.307	1.505	2,109	1.4	06	0.507	0.697			1.781	1.323	2.26	0.1	81 1.532	0.631
23 p	9731.5	0.436	-0.016	0.544	2,307	0.4	Q7	0.539	0.402	2.771	3.451	0.862	1.181	3.625	1.5	44 1.90	0.50

Fig 1. Example bicluster showing log2-FC values of miR-1 targets under multiple conditions

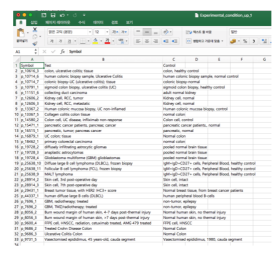


Fig 2. Experimental conditions (test and control conditions) for the example bicluster in figure 1

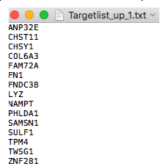


Fig 3. Gene list for the example bicluster in figure 1

4. Detailed function explanation

1. bicluster_extension_process

Extends biclusters allowing small noises

Usage: bicluster_extension_process(MIR_profile, seed_row, seed_col, zero_ratio)

Arguments

MIR_profile A binarized microRNA profile
seed_row List of a seed row index from MIR_profile
seed_col List of a seed column index from MIR_profile
zero_ratio Zero ratio to be allowed.

Value

Extended bicluster matrix

2. bicluster_trimming_process

Removes too noisy row or columns from bicluster.

Usage: bicluster_trimming_process(table, zero_ratio)

Arguments

table A matrix that will be reduced

zero_ratio Zero ratio allowed.

Value

Reduced bicluster matrix

3. getGenelist

Loads a vector of 20,639 human genes

Usage: getGenelist()

Value

A vector of 20,639 human genes

4. getMIRprofile

Returns MIR profile for input miRNA.

Usage: getMIRprofile(miRNA, FCtable = NULL, FCcutoff = log2(1.3))

Arguments

miRNA Human mature miRNA name (e.g., hsa-miR-1-3p). Available miRNAs can be referred by typing

'getmiRNAlist()'

FC matrix of 20,639 genes for 5,158 conditions. If NULL, it will be loaded from the local.

FCcutoff Fold change cutoff in log2 scale. Default = log2(1.3). To extract MIR profile for target down-

regulation in test condition, take negative value.

Value

A matrix of fold change value of sequence-based miRNA targets under selected cell conditions.

5. get_miR_target

Returns sequence-based miRNA targets predicted from three or more algorithms.

Usage: get_miR_target(miRNA)

Arguments

miRNA Human mature miRNA name (e.g., hsa-miR-1-3p). Available miRNAs can be referred by typing 'getmiRNAlist()'

Value

A vector of sequence-based miRNA targets

6. get_zero_rate

Returns the ratio of zero (noise) of a binarized matrix

Usage: get_zero_rate(target)

Arguments

Target a matrix

Value

The ratio of zero in a binarized matrix

7. getmiRNAlist

Loads human mature miRNA list

Usage: getmiRNAlist()

Value

A vector of 2,632 human mature miRNAs

8. load_FCtable

Loads a large table of fold change values for 20,639 human genes in 5,158 cell conditions.

Usage: load_FCtable()

Value

A fold change matrix of 5,158 rows and 20,639 columns.

9. make_biclust

Generates biclusters from MIR profile

Usage: make_biclust(MIR_PROFILE, FCcutoff, REPETITION, STEP_NUMBER, ZERORATE)

Arguments

MIR_PROFILE A matrix of MIR profile.

FCcutoff Binarization fold change cutoff in log2-scale for MIR_PROFILE. Target up-regulation

biclusters will be created for positive FC cutoff, and down-regulation biclusters will be

generated with negative FC cutoff.

REPETITION The number of running ensemble bicluster function.

STEP_NUMBER The number of extension process

ZERORATE Final zero rate allowed.

Value

List of seed and biclusters

10. merge_bicluster

Merges similar biclusters

Usage: merge_bicluster(MIR_profile, tree_cutoff = 0.5, biclusts, seedlist,

FCcutoff = log2(1.3), mir = "testmiRNA", ZERORATE)

Arguments

MIR_profile binarized microRNA profile

tree_cutoff Similarity cutoff

Biclusts list of extended biclusters

Seedlist list of seeds

FCcutoff Fold change cutoff (in log2 sclale) for binarization of MIR profile

Mir Name of miRNA (e.g., hsa-miR-1-3p)

ZERORATE Final zero ratio allowed

Value

Extended bicluster list

11. PBE

Progressive bicluster extension

Usage: PBE(MIR_profile, biclust_row, biclust_col, step_number, finalZR)

Arguments

MIR_profile Binarized microRNA profile

biclust_row List of index of row in seed biclust

biclust_col List of index of columns in seed biclust

step_number The number of extension process

finalZR final zero rate allowed.

Value

List of row and column symbols of extended bicluster

12. PBE_MERGE

Progressive extension and merging of biclusters

Usage: PBE_MERGE(MIR_profile, mir, biclust.path = "./", FCcutoff = log2(1.3),

REPETITION = 10, STEP_NUMBER = 10, ZERORATE = 0.1, tree_cutoff = 0.5)

Arguments

MIR_profile A matrix of microRNA profile where rows are cell conditions and columns are sequence-

based miRNA targets.

mir Name of microRNA

biclust.path Directory where result biclusters and corresponding experimental condition and gene lists

will be saved

FCcutoff Log2-fold change binarization cutoff for seed. Default = log2(1.3)

REPETITION The number of repetition that the ensemble function runs to extract seed biclusters. Default

= 10

STEP_NUMBER The number of extension process. Default = 10

ZERORATE Final zero rate allowed. Default = 0.1

tree_cutoff Similarity cutoff. Default = 0.5

Value

Extended and merged biclusters in the directory assigned to 'biclust.path'.