TRAPR : Total RNA-seq Analysis Package for R User's Guide

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1. Introduce TRAPR

RNA-Seq, is a standard technology for measuring gene expression at an unprecedented accuracy. Numerous Bioconductor packages have been developed for statistical analysis of RNA-Seq data. However, those tools focus on specific aspects of the data analysis pipeline and are hard to integrate appropriately with each other because of their ununified data structure and processing methods. They also lack visualization methods to confirm data integrity and process. Here, we present an R based RNA-Seq analysis pipeline TRAPR, an integrated tool including statistical analysis and visualization of RNA-Seq expression data. TRAPR provides various functions for data management, filtering of low quality data, normalization, transformation, statistical analysis, data visualization, and result visualization that allow researchers to build customized analysis pipelines TRAPR is written in R (as of version 2.15) and is available at http://www.snubi.org/software/trapr

2. Start TRAPR

- * Our manual was made based on window 7
- 2.1 Set up TRAPR package in your R
- 2.2 Previous set up packages

TRAPR must be needs some R packages(vsn, preprocessCore, edgeR, gridExtra, ggplot2, reshape2). We will show description of package setup method. For example, we explain vsn.



Fig1 Dependent packages installation

- > biocLite('vsn')
- > biocLite('preprocessCore')
- > biocLite('gridExtra')
- > biocLite('ggplot2')
- > biocLite('reshape2')

2.3 Loading TRAPR package

> library(TRAPR)

2.4 Format of input file

TRAPR input file consist of Tab-deliminated text file. Column is sample list. Row is gene list. Each cell is filled gene expression values(Fig2).

Gene	1017-NOR	1079-NOR	110-NOR	1199-NOR	1207-NOR	1264-NOR
1/2-SBSRNA	1.05316	1.12635	1.16598	0.903904	1.388	1.7264
A1BG	1.58885	1.53534	1.42561	1.45638	2.60126	1.46374
A1BG-AS1	0.813991	1.18167	0.549843	0.893526	1.82539	0.835946
A1CF	0	0	0	0.0512491	0	0.02684
A2LD1	1.47371	1.40145	1.66409	1.50761	0.89793	3.16842
A2M	677.94	788.356	555.677	536.844	299.616	789.546
A2ML1	0.0461678	0.0332678	0.0441994	0	0.0242496	0
A2MP1	0.161584	0	0.151889	0	0.204479	0.219432
A4GALT	6.09765	6.41299	2.60222	4.72684	4.70813	3.9633
A4GNT	0.306902	0	0.501595	1.12183	0.395527	0.266301
AA06	0	0	0	0	0	0
AAA1	0.130131	0	0	0	0	0
AAAS	8.66273	11.1746	12.4727	17.3145	20.0061	14.5962
AACS	4.82528	4.76136	5.15812	9.67915	6.27222	5.8261
AACSP1	0.163496	0.142302	0.516818	1.56209	0.420916	0.0664654
AADAC	14.6518	4.90271	9.24786	17.3143	3.16036	20.1152
AADACL2	1.91869	0	0.113368	0.964742	0	0.165812
AADACL3	0	0	0.047547	0	0	0
AADACL4	0.112633	0.131725	0	0.0943328	0.0909002	0
AADAT	4.43236	3.92109	5.01229	3.6641	3.4742	5.99568
AAGAB	12.5497	11.6327	11.1433	13.3532	9.16678	12.4696
AAK1	2.6064	2.49532	3.17003	2.4327	2.15502	2.50611

Fig2. input format of TRAPR

2.5 Sample data

Our sample data file name is 'sample.txt'. This file contained in TRAPR. The origin of sample.txt is breast cancer data from TCGA. It consist of 9 normal tissue cancer sample and 10 cancer sample using RNA-seq technology and It's value is FPKM(Fig3).

Gene
Cancer2
Cancer3
Cancer4
Cancer5
Cancer6
Cancer7
Cancer6
Normal1
Normal2
Normal3
Normal4
Normal6
N

Fig3 sample.txt in TRAPR

2.6 Start TRAPR analysis

2.6.1 Data Manipulation

Sample <- TRAPR.Data.ReadExpressionTable('sample.txt', sep = '\t', Exp1 = c(1:9), Exp2 = c(10:19), Tag = c('Normal', 'Cancer'))

Function loading sample file is TRAPR.Data.ReadExpressionTable. The code component explain showing Fig3. More Information in reference manual.





Fig4 sample loading code & example

Sample position (green color) is description for sample column position. Tag (blue color) is description for sample labels. Loaded sample data is saving 'Sample' value and after using function of filtering and normalization. More Information in reference manual.

2.6.2 Filter

lue name	Zero filer function	
Sample <- TRAP	R.Filter.ZeroValue(Sample)	
R Console		
<pre>> Sample = TRA Filter for Zet</pre>	APR.Filter.ZeroValue(Sample) ro values	
	appoint the for all appoint	
26 genes have	zero values for all samples	
26 genes have 15 genes have 12 genes have	zero values for Expl samples zero values for Exp2 samples	



> Sample <- TRAPR.Filter.ZeroValue(Sample)

Remove zero values from sample file. TRAPR provieds 6 kinds of filter condition ('Filter for Gene List', 'TRAPR.Filter.LowVariance', 'TRAPR.Filter.LowExpression', 'TRAPR.Filter.SampleDeletion', 'TRAPR.Filter.GeneDeletion'). If user want other function just typing replace 'TRAPR.Filter.ZeroValue'(read color). More Information in reference manual.

2.6.3 Normalization

′alue name ↑	Normalization function	Kinds of normalization method
nSample	<- TRAPR.Normalize(San	nple, Method = 'Quantile')
E P Consola		
> nSamp. Quantile	le = TRAPR.Normalize(e Normalization	Sample, Method='Quantile')
4		

Fig6 Normalization function code & example

> nSample = TRAPR.Normalize(Sample,Method='Quantile')

TRAPR provide 4 method for normalization ('UpperQuartile', 'Quantile', 'Median', 'Mean'). If you needs another function, just write method name replace 'Quantile'(purple color).

2.6.4 Data Visualization

TRAPR provide pre-processing (filtering + normalization) result visualization. Data visualization function include 5 kinds of plots(Fig7).



Fig7 Data visualization function code & example



Fig8 Result example plot of data visualization

> TRAPR.DataVisualization(nSample, type_of_plot = 'ALL', logged = FALSE)

2.6.5 Statistical Test

TRAPR provide 4 method for statistical test ('ttest', 'wilcoxon', 'edgeR', 'FC'). If you needs another function, just write method name replace ' ttest '(purple color). TRAPR also provide adjust test method('holm', 'hotchberg', 'hommel', 'bonferroni', 'BH', 'BY', 'fdr', 'none').



Fig9 Statistical Test function code & example

2.6.6 Result Visualization

TRAPR provide Result plot of after statistical test. We provide 2 kinds of plots (Heatmap, volcano plot)(Fig11).



Fig10 Result visualization function code & example



Fig11 Result example plot of result visualization

> TRAPR.ResultVisualization(tnSample, type_of_plot='ALL')

2.6.7 Out-print of DEG Result file

TRPR provide out-print your DEG result file in your directory. User just typing file name(blue color) in code.



Fig12 Out-print of DEG Result file function code & example

TRAPR.Data.DEGResulttoFile(tnSample, FileName = 'Result.txt')

3. Reference

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