# GDCRNATools: integrative analysis of protein coding genes, long non-coding genes, and microRNAs in GDC

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# 1 Introduction

GDCRNATools is an R package which provides a standard, easy-to-use and comprehensive pipeline for downloading, organizing, and integrative analyzing RNA expression data in the GDC portal with an emphasis on deciphering the lncRNA-mRNA related ceRNA regulatory network in cancer.

Competing endogenous RNAs (ceRNAs) are RNAs that indirectly regulate other transcripts by competing for shared miRNAs. Although only a fraction of long non-coding RNAs has been functionally characterized, increasing evidences show that lncRNAs harboring multiple miRNA response elements (MREs) can act as ceRNAs to sequester miRNA activity and thus reduce the inhibition of miRNA on its targets. Deregulation of ceRNAs network may lead to human diseases.

The Genomic Data Commons (GDC) maintains standardized genomic, clinical, and biospecimen data from National Cancer Institute (NCI) programs including The Cancer Genome Atlas (TCGA) and Therapeutically Applicable Research To Generate Effective Treatments (TARGET), It also accepts high quality datasets from non-NCI supported cancer research programs, such as genomic data from the Foundation Medicine.

Many analyses can be perfomed using GDCRNATools, including differential gene expression analysis (limma(Ritchie et al. 2015), edgeR(Robinson, McCarthy, and Smyth 2010), and DESeq2(Love, Huber, and Anders 2014)), univariate survival analysis (CoxPH and KM), competing endogenous RNA network analysis (hypergeometric test, Pearson correlation analysis, regulation similarity analysis, sensitivity Pearson partial correlation(Paci, Colombo, and Farina 2014)), and functional enrichment analysis(GO, KEGG, DO). Besides some routine visualization methods such as volcano plot, scatter plot, and bubble plot, etc., three simple shiny apps are developed in GDCRNATools allowing users visualize the results on a local webpage. All the figures are plotted based on ggplot2 package unless otherwise specified.

This user-friendly package allows researchers perform the analysis by simply running a few functions and integrate their own pipelines such as molecular subtype classification, weighted correlation network analysis (WGCNA)(Langfelder and Horvath 2008), and TF-miRNA co-regulatory network analysis, etc. into the workflow easily. This could open a door to accelerate the study of crosstalk among different classes of RNAs and their regulatory relationships in cancer.

# 2 GDCRNATools package installation

The R software for running GDCRNATools can be downloaded from The Comprehensive R Archive Network (CRAN). The GDCRNATools package can be installed from Github.

devtools::install\_github(repo='Jialab-UCR/GDCRNATools')

library(GDCRNATools)

### 3 Data download

Two methods are provided for downloading Gene Expression Quantification (HTSeq-Counts), Isoform Expression Quantification (BCGSC miRNA Profiling), and Clinical (Clinical Supplement) data:

• Manual download

Step1: Download GDC Data Transfer Tool on the GDC website

Step2: Add data to the GDC cart, then download manifest file and metadata of the cart

Step3: Download data using gdcRNADownload() function by providing the manifest file

Automatic download

Download GDC Data Transfer Tool, manifest file, and data automatically by specifying the project.id and data.type in gdcRNADownload() function for RNAseq and miRNAs data, and in gdcClinicalDownload() function for clinical data

Users can also download data from GDC using the API method developed in TCGAbiolinks(Colaprico et al. 2016) or using TCGA-Assembler(Zhu, Qiu, and Ji 2014)

### 3.1 Manual download

### 3.1.1 Installation of GDC Data Transfer Tool gdc-client

Download GDC Data Transfer Tool from the GDC website and unzip the file

### 3.1.2 Download manifest file and metadata from GDC Data Portal

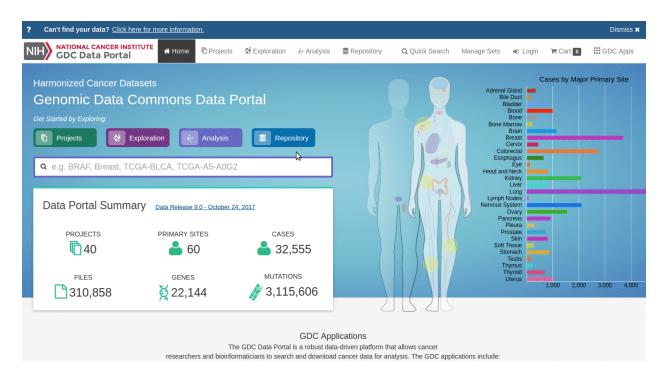


Figure 1:

### 3.1.3 Download data

### 3.2 Automatic download

• gdcRNADownload() will download HTSeq-Counts data if data.type='RNAseq' and download BCGSC miRNA Profiling data if data.type='miRNAs'. project.id argument is required to be provided.

• gdcClinicalDownload() download clinical data in .xml format automatically by simply specifying the project.id argument.

### 3.2.1 Download RNAseq/miRNAs data

#### 3.2.2 Download clinical data

# 4 Data organization

### 4.1 Parse metadata

Metadata can be parsed by either providing the metadata file that is downloaded in the data download step, or specifying the project.id and data.type in gdcParseMetadata() function to obtain information of data in the manifest file to facilitate data organization and basic clinical information of patients such as age, stage and gender, etc. for data analysis.

### 4.1.1 Parse metadata by providing the metadata file

```
####### Parse RNAseq metadata #######
metaMatrix.RNA <- gdcParseMetadata(metafile='TCGA-PRAD/TCGA-PRAD.RNAseq.metadata.2017-11-23T17-23-59.js
####### Parse miRNAs metadata ######
metaMatrix.MIR <- gdcParseMetadata(metafile='TCGA-PRAD/TCGA-PRAD.miRNAs.metadata.2017-11-23T17-33-55.js</pre>
```

### 4.1.2 Parse metadata by specifying project.id and data.type

```
write.meta = TRUE)
metaMatrix.RNA[1:6,1:6]
##
                                                                file name
## TCGA-2A-A8VL-01A d5b5e519-5ce4-4147-a500-25b2f442152d.htseq.counts.gz
## TCGA-2A-A8VO-01A ed22ecf9-2215-4bc4-a660-f8dbd2e2d15c.htseq.counts.gz
## TCGA-2A-A8VT-01A 4dd0008c-f544-438d-8802-e02dbf6c4a3e.htseq.counts.gz
## TCGA-2A-A8VV-01A 8f92f24b-90c7-46d0-b69d-e1d87a135a40.htseq.counts.gz
## TCGA-2A-A8VX-01A 6f421ec2-c74a-4719-b447-fab39c619d3b.htseq.counts.gz
## TCGA-2A-A8W1-01A 5d408d1f-e5e4-4901-b1a8-bf803e633117.htseq.counts.gz
##
                                                  file_id
                                                               patient
## TCGA-2A-A8VL-01A 2b760030-1cad-4554-931e-bac9205b56ca TCGA-2A-A8VL
## TCGA-2A-A8VO-01A 8c4da184-f1e1-4439-b501-8c3f88ba0d23 TCGA-2A-A8VO
## TCGA-2A-A8VT-01A 32eb6261-4d8b-4f8e-9435-cdbdc4766a3d TCGA-2A-A8VT
## TCGA-2A-A8VV-01A 8a880dc3-c706-4a16-9e1a-39adfdb7c72a TCGA-2A-A8VV
## TCGA-2A-A8VX-01A 2bacf214-6dfb-44e3-b5ab-ec51c343b24e TCGA-2A-A8VX
## TCGA-2A-A8W1-01A f6c810f2-344e-4e8e-8e7a-01c18ec72074 TCGA-2A-A8W1
                                        submitter_id
                             sample
## TCGA-2A-A8VL-01A TCGA-2A-A8VL-01 TCGA-2A-A8VL-01A
## TCGA-2A-A8VO-01A TCGA-2A-A8VO-01 TCGA-2A-A8VO-01A
## TCGA-2A-A8VT-01A TCGA-2A-A8VT-01 TCGA-2A-A8VT-01A
## TCGA-2A-A8VV-01A TCGA-2A-A8VV-01 TCGA-2A-A8VV-01A
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01 TCGA-2A-A8VX-01A
## TCGA-2A-A8W1-01A TCGA-2A-A8W1-01 TCGA-2A-A8W1-01A
                             entity_submitter_id
## TCGA-2A-A8VL-01A TCGA-2A-A8VL-01A-21R-A37L-07
## TCGA-2A-A8VO-01A TCGA-2A-A8VO-01A-11R-A37L-07
## TCGA-2A-A8VT-01A TCGA-2A-A8VT-01A-11R-A37L-07
## TCGA-2A-A8VV-01A TCGA-2A-A8VV-01A-11R-A37L-07
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01A-11R-A37L-07
## TCGA-2A-A8W1-01A TCGA-2A-A8W1-01A-11R-A37L-07
###### Parse miRNAs metadata ######
metaMatrix.MIR <- gdcParseMetadata(project.id = 'TCGA-PRAD',</pre>
                                   data.type = 'miRNAs',
                                   write.meta = TRUE)
metaMatrix.MIR[1:6,1:6]
##
                                                                                      file_name
## TCGA-2A-A8VL-01A a8ad4b62-68e8-4d56-893e-e247a3099d94.mirbase21.isoforms.quantification.txt
## TCGA-2A-A8VO-01A 22302d39-da19-4bfd-b4d8-aa951b9451a1.mirbase21.isoforms.quantification.txt
## TCGA-2A-A8VT-01A de5cc4c2-2709-4bbe-8777-9d8e9cd56246.mirbase21.isoforms.quantification.txt
## TCGA-2A-A8VV-01A f3402505-e2c1-4720-a9f2-f39105ad0327.mirbase21.isoforms.quantification.txt
## TCGA-2A-A8VX-01A c2a2d423-e481-4821-9970-5e93d7d4442b.mirbase21.isoforms.quantification.txt
## TCGA-2A-A8W1-01A b1a5f1a4-a95a-4770-a234-709c4e9da1fe.mirbase21.isoforms.quantification.txt
                                                 file_id
                                                               patient
## TCGA-2A-A8VL-01A a0b6cbc1-43fa-4bed-83e8-917794158b98 TCGA-2A-A8VL
## TCGA-2A-A8VO-01A addea5e5-5b25-417c-bbb2-00438b8da4c6 TCGA-2A-A8VO
## TCGA-2A-A8VT-01A 7a337162-08ee-4600-96f5-79fed7b68898 TCGA-2A-A8VT
## TCGA-2A-A8VV-01A fc64fdd9-b679-4a97-bf5e-d757b64b252c TCGA-2A-A8VV
## TCGA-2A-A8VX-01A 8387f768-8d31-4ffa-88ae-dae0ef11b2fb TCGA-2A-A8VX
## TCGA-2A-A8W1-01A cb18f79c-41d4-4bb9-af6a-28e35b6a4470 TCGA-2A-A8W1
##
                             sample
                                        submitter id
## TCGA-2A-A8VL-01A TCGA-2A-A8VL-01 TCGA-2A-A8VL-01A
```

```
## TCGA-2A-A8VO-01A TCGA-2A-A8VO-01 TCGA-2A-A8VO-01A
## TCGA-2A-A8VT-01A TCGA-2A-A8VT-01 TCGA-2A-A8VT-01A
## TCGA-2A-A8VV-01A TCGA-2A-A8VV-01 TCGA-2A-A8VV-01A
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01 TCGA-2A-A8VX-01A
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01 TCGA-2A-A8VX-01A
## TCGA-2A-A8VI-01A TCGA-2A-A8VI-01 TCGA-2A-A8VI-01A
## TCGA-2A-A8VL-01A TCGA-2A-A8VL-01A-21R-A37H-13
## TCGA-2A-A8VT-01A TCGA-2A-A8VT-01A-11R-A37H-13
## TCGA-2A-A8VV-01A TCGA-2A-A8VV-01A-11R-A37H-13
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01A-11R-A37H-13
## TCGA-2A-A8VX-01A TCGA-2A-A8VX-01A-11R-A37H-13
## TCGA-2A-A8VI-01A TCGA-2A-A8VX-01A-11R-A37H-13
## TCGA-2A-A8VI-01A TCGA-2A-A8VI-01A-11R-A37H-13
```

### 4.2 Filter samples

### 4.2.1 Filter duplicated samples

Only one sample would be kept if the sample had been sequenced more than once by gdcFilterDuplicate(). ####### Filter duplicated samples in RNAseq metadata #######

```
metaMatrix.RNA <- gdcFilterDuplicate(metaMatrix.RNA)

## Removed 3 samples
####### Filter duplicated samples in miRNAs metadata #######</pre>
```

```
metaMatrix.MIR <- gdcFilterDuplicate(metaMatrix.MIR)</pre>
```

## Removed 4 samples

#### 4.2.2 Filter non-Primary Tumor and non-Solid Tissue Normal samples

Samples that are neither Primary Tumor (code: 01) nor Solid Tissue Normal (code: 11) would be filtered out by gdcFilterSampleType().

####### Filter non-Primary Tumor and non-Solid Tissue Normal samples in RNAseq metadata ######
metaMatrix.RNA <- gdcFilterSampleType(metaMatrix.RNA)</pre>

```
## Removed 1 samples
```

####### Filter non-Primary Tumor and non-Solid Tissue Normal samples in miRNAs metadata ######
metaMatrix.MIR <- gdcFilterSampleType(metaMatrix.MIR)</pre>

## Removed 1 samples

### 4.3 Merge data

- gdcRNAMerge() merges raw counts data of RNAseq to a single expression matrix with rows are *Ensembl* id and columns are samples. Total read counts for 5p and 3p strands of miRNAs can be processed from isoform quantification files and then merged to a single expression matrix with rows are miRBase v21 identifiers and columns are samples.
- gdcClinicalMerge() merges clinical data to a dataframe with rows are *patient id* and columns are *clinical traits*. If key.info=TRUE, only those most commonly used clinical traits will be reported, otherwise, all the clinical information will be reported.

### 4.3.1 Merge RNAseq/miRNAs data

```
###### Merge RNAseq data ######
rnaMatrix <- gdcRNAMerge(metadata = metaMatrix.RNA,</pre>
                         path
                                   = 'TCGA-PRAD/RNAseq/',
                         data.type = 'RNAseq')
## ########### Merging RNAseq data ##############
## ### This step may take a few minutes ###
## Number of samples: 547
## Number of genes: 60483
rnaMatrix[1:6,1:6]
##
                   TCGA-2A-A8VL-01 TCGA-2A-A8VO-01 TCGA-2A-A8VT-01
## ENSG00000000003
                              2867
                                                               3140
                                              1667
## ENSG0000000005
                                                                  0
                                6
                                                 0
## ENSG00000000419
                              1354
                                               888
                                                               1767
## ENSG0000000457
                                               580
                               956
                                                               2163
## ENSG0000000460
                               119
                                                91
                                                                305
## ENSG0000000938
                               159
                                               171
                                                                228
                   TCGA-2A-A8VV-01 TCGA-2A-A8VX-01 TCGA-2A-A8W1-01
##
## ENSG0000000003
                              3996
                                              4869
                                                               2172
## ENSG0000000005
                               44
                                                                  0
                                                1
## ENSG0000000419
                              1408
                                              1171
                                                               1593
## ENSG0000000457
                              1494
                                               908
                                                                794
## ENSG0000000460
                               175
                                                121
                                                                166
## ENSG0000000938
                               172
                                                64
                                                                161
###### Merge miRNAs data ######
mirMatrix <- gdcRNAMerge(metadata = metaMatrix.MIR,
                         path = 'TCGA-PRAD/miRNAs/',
                         data.type = 'miRNAs')
## ############ Merging miRNAs data #############
## Number of samples: 546
## Number of miRNAs: 2588
mirMatrix[1:6,1:6]
##
                   TCGA-2A-A8VL-01 TCGA-2A-A8VO-01 TCGA-2A-A8VT-01
## hsa-let-7a-5p
                            130022
                                             77195
                                                             170937
## hsa-let-7a-3p
                               133
                                                                 91
                                                 10
## hsa-let-7a-2-3p
                                18
                                                                 13
                             68276
                                                              36009
## hsa-let-7b-5p
                                             19131
## hsa-let-7b-3p
                                78
                                                 30
                                                                 55
## hsa-let-7c-5p
                             43015
                                             22490
                                                              14099
##
                   TCGA-2A-A8VV-01 TCGA-2A-A8VX-01 TCGA-2A-A8W1-01
## hsa-let-7a-5p
                            247370
                                             73705
                                                              50261
## hsa-let-7a-3p
                               104
                                                59
                                                                 39
## hsa-let-7a-2-3p
                                13
                                                 3
                                                                  4
## hsa-let-7b-5p
                             58349
                                             17404
                                                               6663
## hsa-let-7b-3p
                                73
                                                19
                                                                 18
## hsa-let-7c-5p
                             36248
                                              9694
                                                              11759
```

#### 4.3.2 Merge clinical data

```
###### Merge clinical data ######
clinicalDa <- gdcClinicalMerge(path = 'TCGA-PRAD/Clinical/', key.info = TRUE)</pre>
## ############ Merging Clinical data ##############
clinicalDa[1:6,5:10]
##
                 clinical_stage clinical_T clinical_N clinical_M
## TCGA-EJ-5510
                                         T1c
                                                      NA
                                                                  MO
                              NA
## TCGA-HC-8260
                              NA
                                          NA
                                                      NA
                                                                 MO
## TCGA-Y6-A8TL
                              NA
                                         T2a
                                                      NA
                                                                  NA
                                                      NA
## TCGA-V1-A8X3
                              NA
                                         T1c
                                                                 MO
## TCGA-VP-A87J
                              NA
                                         T2a
                                                      NA
                                                                 MO
## TCGA-KK-A6DY
                              NA
                                         T<sub>1</sub>c
                                                      NA
                                                                  MO
##
                 gleason_grading gleason_score
## TCGA-EJ-5510
                             7433
                                               7
                                               7
## TCGA-HC-8260
                              734
## TCGA-Y6-A8TL
                              633
                                               6
                                               7
## TCGA-V1-A8X3
                              734
## TCGA-VP-A87J
                              734
                                               7
                                               7
## TCGA-KK-A6DY
                              734
```

### 4.4 TMM normalization and voom transformation

It has repeatedly shown that normalization is a critical way to ensure accurate estimation and detection of differential expression (DE) by removing systematic technical effects that occur in the data(Robinson and Oshlack 2010). TMM normalization is a simple and effective method for estimating relative RNA production levels from RNA-seq data. Voom is moreover faster and more convenient than existing RNA-seq methods, and converts RNA-seq data into a form that can be analyzed using similar tools as for microarrays(Law et al. 2014).

By running gdcVoomNormalization() function, raw counts data would be normalized by TMM method implemented in edgeR(Robinson, McCarthy, and Smyth 2010) and further transformed by the voom method provided in limma(Ritchie et al. 2015). Low expression genes (logcpm < 1 in more than half of the samples) will be filtered out by default. All the genes can be kept by setting filter=TRUE in the gdcVoomNormalization().

```
####### RNAseq data #######
rnaExpr <- gdcVoomNormalization(counts = rnaMatrix, filter = FALSE)
rnaExpr[1:6,1:6]</pre>
```

```
##
                   TCGA-2A-A8VL-01 TCGA-2A-A8VO-01 TCGA-2A-A8VT-01
## ENSG0000000003
                           5.891004
                                           5.469541
                                                            5.675430
## ENSG0000000005
                          -2.894134
                                          -6.233930
                                                           -6.941348
## ENSG0000000419
                           4.808971
                                           4.561298
                                                            4.846146
## ENSG0000000457
                           4.307047
                                           3.947222
                                                            5.137803
## ENSG0000000460
                           1.306293
                                           1.281770
                                                            2.313680
## ENSG0000000938
                           1.722839
                                           2.188135
                                                            1.894702
##
                   TCGA-2A-A8VV-01 TCGA-2A-A8VX-01 TCGA-2A-A8W1-01
## ENSG0000000003
                         6.3329382
                                          6.6613451
                                                            5.612615
## ENSG00000000005
                         -0.1558497
                                         -5.0032503
                                                           -6.472525
## ENSG0000000419
                         4.8283607
                                          4.6059284
                                                            5.165458
## ENSG0000000457
                         4.9138640
                                          4.2391299
                                                            4.161378
## ENSG0000000460
                          1.8237441
                                          1.3365997
                                                            1.906853
```

```
## ENSG0000000938
                          1.7988694
                                          0.4230145
                                                            1.862865
###### miRNAs data ######
mirExpr <- gdcVoomNormalization(counts = mirMatrix, filter = FALSE)
mirExpr[1:6,1:6]
##
                   TCGA-2A-A8VL-01 TCGA-2A-A8VO-01 TCGA-2A-A8VT-01
## hsa-let-7a-5p
                          14.676762
                                          14.246607
                                                           15.773276
## hsa-let-7a-3p
                           4.749056
                                           4.411257
                                                            4.905866
## hsa-let-7a-2-3p
                           1.897814
                                            1.402695
                                                            2.145054
## hsa-let-7b-5p
                                          12.234040
                                                           13.526256
                          13.747462
## hsa-let-7b-3p
                           3.982981
                                           2.941115
                                                            4.184582
## hsa-let-7c-5p
                          13.080929
                                          12.467406
                                                           12.173523
##
                   TCGA-2A-A8VV-01 TCGA-2A-A8VX-01 TCGA-2A-A8W1-01
## hsa-let-7a-5p
                          15.705812
                                          14.8712423
                                                           14.456561
## hsa-let-7a-3p
                           4.496858
                                          4.5965754
                                                            4.143175
## hsa-let-7a-2-3p
                           1.544386
                                          0.5091125
                                                            1.009320
## hsa-let-7b-5p
                          13.621931
                                          12.7889304
                                                           11.541459
## hsa-let-7b-3p
                           3.989171
                                          2.9871598
                                                            3.048848
## hsa-let-7c-5p
                          12.935132
                                          11.9447084
                                                           12.360934
```

### 5 Differential gene expression analysis

gdcDEAnalysis(), a convenience wrapper, provides three widely used methods limma(Ritchie et al. 2015), edgeR(Robinson, McCarthy, and Smyth 2010), and DESeq2(Love, Huber, and Anders 2014) to identify differentially expressed genes (DEGs) or miRNAs between any two groups defined by users. Note that DESeq2(Love, Huber, and Anders 2014) maybe slow with a single core. Multiple cores can be specified with the nCore argument if DESeq2(Love, Huber, and Anders 2014) is in use. Users are encouraged to consult the vignette of each method for more detailed information.

### 5.1 DE analysis

```
DEGAll <- gdcDEAnalysis(counts</pre>
                                    = rnaMatrix,
                                    = metaMatrix.RNA$sample_type,
                        group
                        comparison = 'PrimaryTumor-SolidTissueNormal',
                                    = 'limma')
                        method
DEGA11[1:6,]
##
                    symbol
                                               logFC
                                                       AveExpr
                                     group
                                                                       t
## ENSG00000187699 C2orf88 protein coding -2.657180 1.5056478 -19.46636
## ENSG0000176928
                     GCNT4 protein_coding -2.248112 0.5798701 -18.39206
                      CA14 protein_coding -2.630802 0.4748363 -17.57925
## ENSG0000118298
## ENSG0000103485
                      QPRT protein_coding -2.147259 1.9897483 -17.32704
## ENSG0000109667
                    SLC2A9 protein_coding -1.869863 1.6079446 -17.21612
## ENSG0000164764
                    SBSPON protein_coding -2.333725 2.5270242 -17.17468
##
                         PValue
                                          FDR.
                                                     В
## ENSG00000187699 1.453473e-64 2.259715e-60 136.1299
## ENSG00000176928 3.303402e-59 2.567900e-55 123.6779
## ENSG00000118298 3.348976e-55 1.735551e-51 114.6210
## ENSG00000103485 5.729814e-54 2.227035e-50 111.9647
## ENSG00000109667 1.990189e-53 6.188294e-50 110.6756
```

## 5.2 Report DE genes/miRNAs

All DEGs, DE long non-coding genes, DE protein coding genes and DE miRNAs could be reported separately by setting geneType argument in gdcDEReport(). Gene symbols and biotypes based on the Ensembl 90 annotation are reported in the output.

```
### All DEGs
deALL <- gdcDEReport(deg = DEGAll, gene.type = 'all')

#### DE long-noncoding
deLNC <- gdcDEReport(deg = DEGAll, gene.type = 'long_non_coding')

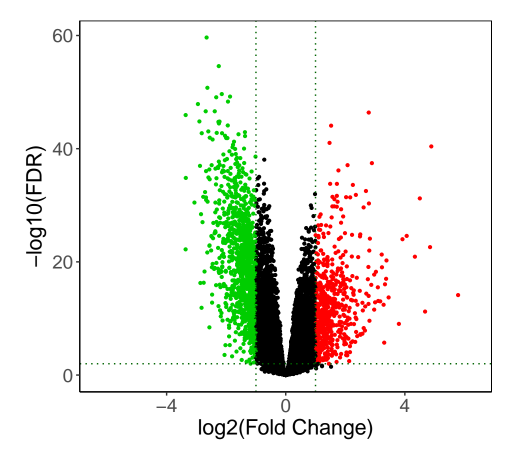
#### DE protein coding genes
dePC <- gdcDEReport(deg = DEGAll, gene.type = 'protein_coding')</pre>
```

### 5.3 DEG visualization

Volcano plot and bar plot are used to visualize DE analysis results in different manners by gdcVolcanoPlot() and gdcBarPlot() functions, respectively. Hierarchical clustering on the expression matrix of DEGs can be analyzed and plotted by the gdcHeatmap() function.

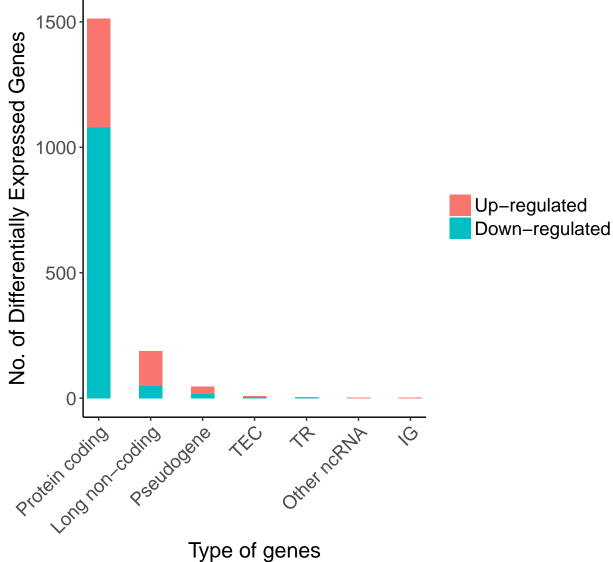
### 5.3.1 Volcano plot

```
gdcVolcanoPlot(DEGAll)
```



# 5.3.2 Barplot

```
gdcBarPlot(deg = deALL, angle = 45, data.type = 'RNAseq')
```



# Type of gene

### 5.3.3 Heatmap

Heatmap is generated based on the heatmap.2() function in gplots package.

```
degName = rownames(deALL)
gdcHeatmap(deg.id = degName, metadata = metaMatrix.RNA, rna.expr = rnaExpr)
```

# 6 Competing endogenous RNAs network analysis

Three criteria are used to determine the competing endogenous interactions between lncRNA-mRNA pairs:

- The lncRNA and mRNA must share significant number of miRNAs
- Expression of lncRNA and mRNA must be positively correlated
- Those common miRNAs should play similar roles in regulating the expression of lncRNA and mRNA

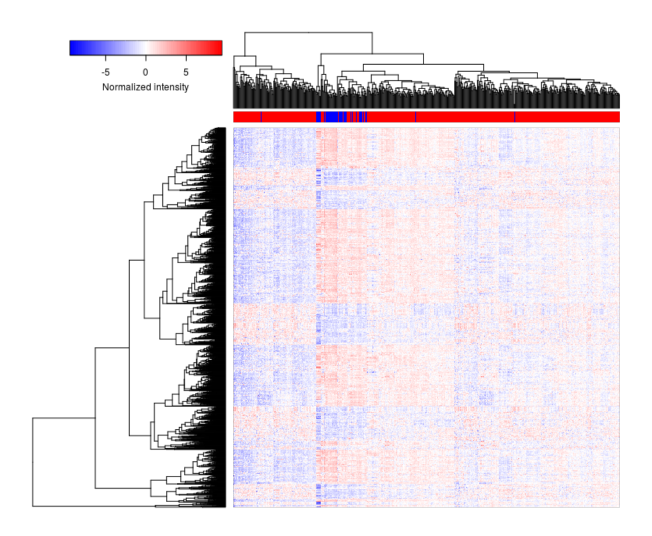


Figure 2:

### 6.1 Hypergeometric test

Hypergenometric test is performed to test whether a lncRNA and mRNA share many miRNAs significantly.

A newly developed algorithm **spongeScan**(Furió-Tarí et al. 2016) is used to predict MREs in lncRNAs acting as ceRNAs. Databases such as **starBase v2.0**(J.-H. Li et al. 2014), **miRcode**(Jeggari, Marks, and Larsson 2012) and **mirTarBase release 7.0**(Chou et al. 2017) are used to collect predicted and experimentally validated miRNA-mRNA and/or miRNA-lncRNA interactions. Gene IDs in these databases are updated to the latest Ensembl 90 annotation of human genome and miRNAs names are updated to the new release miRBase 21 identifiers. Users can also provide their own datasets of miRNA-lncRNA and miRNA-mRNA interactions.

The figure and equation below illustrate how the hypergeometric test works

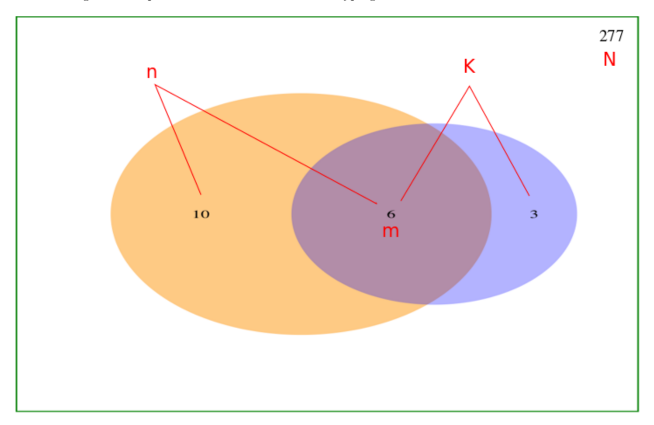


Figure 3:

$$p = 1 - \sum_{k=0}^{m} \frac{\binom{K}{k} \binom{N-K}{n-k}}{\binom{N}{n}}$$

here m is the number of shared miRNAs, N is the total number of miRNAs in the database, n is the number of miRNAs targeting the lncRNA, K is the number of miRNAs targeting the protein coding gene.

### 6.2 Pearson correlation analysis

Pearson correlation coefficient is a measure of the strength of a linear association between two variables. As we all know, miRNAs are negative regulators of gene expression. If more common miRNAs are occupied by a lncRNA, less of them will bind to the target mRNA, thus increasing the expression level of mRNA. So expression of the lncRNA and mRNA in a ceRNA pair should be positively correlated.

### 6.3 Regulation pattern analysis

Two methods are used to measure the regulatory role of miRNAs on the lncRNA and mRNA:

• Regulation similarity

We defined a measurement regulation similarity score to check the similarity between miRNAs-lncRNA expression correlation and miRNAs-mRNA expression correlation.

$$Regulation \ similarity \ score = 1 - \frac{1}{M} \sum_{k=1}^{M} \left[ \frac{|corr(m_k, l) - corr(m_k, g)|}{|corr(m_k, l)| + |corr(m_k, g)|} \right]^{M}$$

where M is the total number of shared miRNAs, k is the kth shared miRNAs,  $corr(m_k, l)$  and  $corr(m_k, g)$  represents the Pearson correlation between the kth miRNA and lncRNA, the kth miRNA and mRNA, respectively

• Sensitivity correlation

##

foldEnrichment

Sensitivity correlation is defined by Paci et al.(Paci, Colombo, and Farina 2014) to measure if the correlation between a lncRNA and mRNA is mediated by a miRNA in the lncRNA-miRNA-mRNA triplet. We take average of all triplets of a lncRNA-mRNA pair and their shared miRNAs as the sensitivity correlation between a selected lncRNA and mRNA.

Sensitivity correlation = 
$$corr(l, g) - \frac{1}{M} \sum_{k=1}^{M} \frac{corr(l, g) - corr(m_k, l)corr(m_k, g)}{\sqrt{1 - corr(m_k, l)^2} \sqrt{1 - corr(m_k, g)^2}}$$

where M is the total number of shared miRNAs, k is the kth shared miRNAs, corr(l, g),  $corr(m_k, l)$  and  $corr(m_k, g)$  represents the Pearson correlation between the long non-coding RNA and the protein coding gene, the kth miRNA and lncRNA, the kth miRNA and mRNA, respectively

The hypergeometric test of shared miRNAs, expression correlation analysis of lncRNA-mRNA pair, and regulation pattern analysis of shared miRNAs are all implemented in the gdcCEAnalysis() function.

```
ceOutput <- gdcCEAnalysis(lnc
                                        = rownames(deLNC),
                                       = rownames(dePC),
                           lnc.targets = 'starBase',
                           pc.targets = 'starBase';
                           rna.expr
                                       = rnaExpr,
                           mir.expr
                                        = mirExpr)
## Step 1/3: Hypergenometric test done!
## Step 2/3: Correlation analysis done!
## Step 3/3: Regulation pattern analysis done!
ceOutput <- ceOutput[order(ceOutput$regSim),]</pre>
ceOutput[1:6,]
                                 Genes Counts listTotal popHits popTotal
              lncRNAs
## 22 ENSG00000234456 ENSG00000163110
                                             2
                                                       2
                                                               36
                                                                       277
                                             2
   34 ENSG00000234456 ENSG00000043591
                                                       2
                                                               51
                                                                       277
  37 ENSG00000234456 ENSG00000119547
                                             2
                                                       2
                                                               71
                                                                       277
  45 ENSG00000234456 ENSG00000112984
                                             2
                                                       2
                                                                       277
                                                       2
  57 ENSG00000234456 ENSG00000184838
                                                               23
                                                                       277
## 65 ENSG00000228223 ENSG00000129514
                                             1
                                                               23
                                                                       277
```

miRNAs

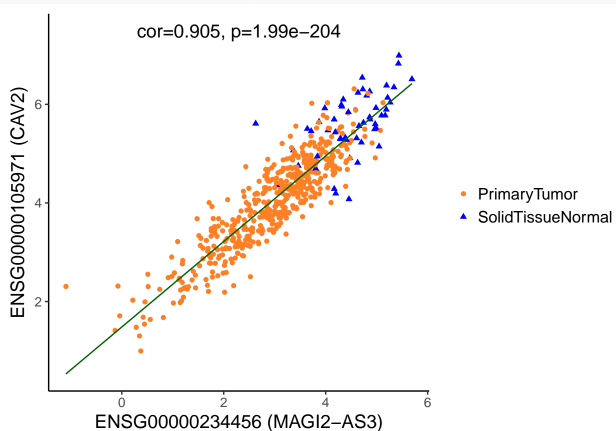
hyperPValue

```
## 22 7.6944444444444
                         0.016480929210485 hsa-miR-374b-5p,hsa-miR-374a-5p
## 34 5.43137254901961
                         0.0333542614974101 hsa-miR-374b-5p,hsa-miR-374a-5p
## 37 3.90140845070423
                         0.0650081096635797 hsa-miR-374b-5p,hsa-miR-374a-5p
## 45 92.333333333333 7.84806152880239e-05 hsa-miR-374b-5p,hsa-miR-374a-5p
## 57 12.0434782608696
                        0.00661853188929001 hsa-miR-374b-5p,hsa-miR-374a-5p
  65 6.02173913043478
                          0.159446450060168
                                                             hsa-miR-590-3p
             cor corPValue regSim
## 22 -0.3384664 1.0000000
                                  0.0006305722
## 34 -0.4330765 1.0000000
                                0 0.0012113510
## 37 -0.3662249 1.0000000
                                0 -0.0241567080
## 45 -0.4187818 1.0000000
                                0 -0.0297453228
## 57 -0.2210811 0.9999999
                                0 -0.0155415582
## 65 -0.3084449 1.0000000
                                0 -0.0061604421
```

### 6.4 ceRNAs visualization

### 6.4.1 Correlation plot

```
gdcCorPlot(gene1 = 'ENSG00000234456',
    gene2 = 'ENSG00000105971',
    rna.expr = rnaExpr,
    metadata = metaMatrix.RNA)
```



### 6.4.2 Correlation plot on a local webpage by shinyCorplot

Typing and running gdcCorPlot() for each pair of lncRNA-mRNA is bothering when multiple pairs are being interested in. shinyCorPlot(), a interactive plot function based on shiny package, can be easily operated by just clicking the genes in each drop down box (in the GUI window). By running shinyCorPlot() function, a local webpage would pop up and correlation plot between a lncRNA and mRNA would be automatically shown.

```
shinyCorPlot(gene1 = rownames(deLNC),
    gene2 = rownames(dePC),
    rna.expr = rnaExpr,
    metadata = metaMatrix.RNA)
```

# Expression correlation of ce pairs

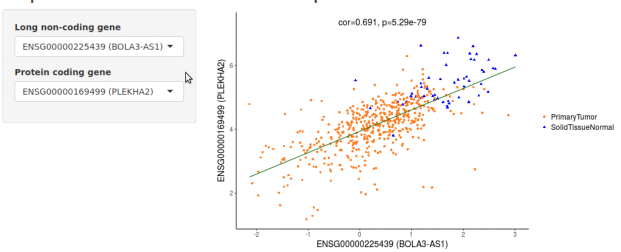


Figure 4:

### 6.4.3 Network visulization in Cytoscape

lncRNA-miRNA-mRNA interactions can be reported by the gdcExportNetwork() and visualized in Cytoscape.

```
ceOutput2 <- ceOutput[ceOutput$hyperPValue<0.01 & ceOutput$corPValue<0.01 & ceOutput$regSim != 0,]
edges <- gdcExportNetwork(ceNetwork = ceOutput2, net = 'edges')
edges[1:6,]</pre>
```

```
fromNode
                              toNode altNode1Name
## 1 ENSG00000234456 hsa-miR-374b-5p
                                        MAGI2-AS3
## 2 ENSG00000234456 hsa-miR-374a-5p
                                        MAGI2-AS3
## 3 ENSG00000245532
                       hsa-let-7i-5p
                                            NEAT1
## 4 ENSG00000245532
                      hsa-let-7e-5p
                                            NEAT1
## 5 ENSG00000245532
                      hsa-let-7g-5p
                                            NEAT1
## 6 ENSG00000245532
                      hsa-let-7f-5p
                                            NEAT1
```

```
nodes <- gdcExportNetwork(ceNetwork = ceOutput2, net = 'nodes')
nodes[1:6,]</pre>
```

```
##
                gene symbol type numInteractions
## 1 ENSG00000008300 CELSR3
                               рс
## 2 ENSG00000047597
                                                 2
                               рс
                                                 2
## 3 ENSG00000065320
                        NTN1
                               рс
                                                 2
## 4 ENSG00000065534
                        MYLK
                               рс
                                                 2
## 5 ENSG00000066468
                      FGFR2
                               рс
## 6 ENSG00000075651
                                                 1
                        PLD1
                               рс
```

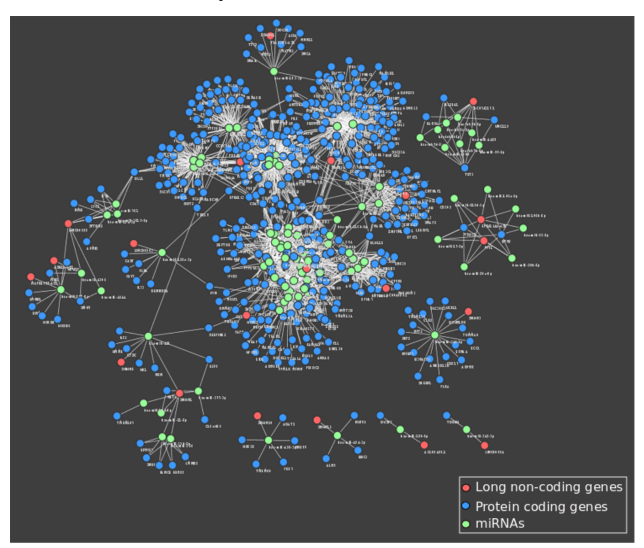


Figure 5:

# 7 Univariate survival analysis

Two methods are provided to perform univariate survival analysis: Cox Proportional-Hazards (CoxPH) model and Kaplan Meier (KM) analysis based on the survival package. CoxPH model considers expression value as

continous variable while KM analysis divides patients into high-expression and low-expression groups by a user-defined threshold such as median or mean. gdcSurvivalAnalysis() take a list of genes as input and report the hazard ratio, 95% confidence intervals, and test significance of each gene on overall survival.

### 7.1 CoxPH analysis

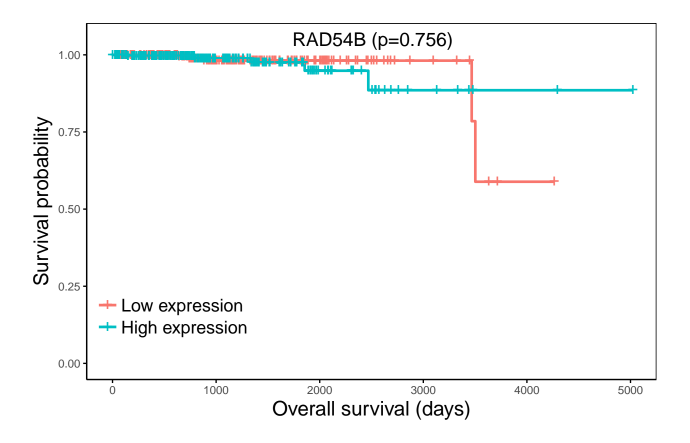
```
###### CoxPH analysis ######
survOutput <- gdcSurvivalAnalysis(gene</pre>
                                           = rownames(deALL),
                                  method
                                           = 'coxph',
                                  rna.expr = rnaExpr,
                                  metadata = metaMatrix.RNA)
head(survOutput[order(survOutput$pValue),])
##
                       symbol
                                    coef
                                                HR
                                                     lower95
                                                                upper95
## ENSG0000156804
                       FBX032 -0.9061689 0.4040693 0.2444365
                                                              0.6679526
## ENSG00000273478 AC099676.1 1.8426288 6.3131126 1.9864365 20.0637629
## ENSG0000069535
                         MAOB -0.4870443 0.6144398 0.4517982
                                                              0.8356304
## ENSG0000128298
                     BAIAP2L2 0.4950804 1.6406302 1.1837845
                                                              2.2737816
## ENSG00000255545 AP004608.1 0.7108727 2.0357671 1.2702956
                                                             3.2625066
## ENSG0000180447
                         GAS1 -0.6253213 0.5350895 0.3530306 0.8110367
##
                         pValue
## ENSG00000156804 0.0004100575
## ENSG00000273478 0.0017880566
## ENSG00000069535 0.0019053414
## ENSG00000128298 0.0029472842
## ENSG00000255545 0.0031344643
## ENSG00000180447 0.0032084588
```

### 7.2 KM analysis

### 7.3 KM analysis visualization

### 7.3.1 KM plot

KM survival curves are plotted using the gdckMPlot() function which is based on the R package survminer.



### 7.3.2 KM plot on a local webpage by shinyKMPlot

The shinyKMPlot() function is also a simply shiny app which allow users view KM plots of all genes of interests on a local webpackage conveniently.

```
shinyKMPlot(gene = rownames(deALL), rna.expr = rnaExpr, metadata = metaMatrix.RNA)
```

# 8 Functional enrichment analysis

One of the main uses of the GO is to perform enrichment analysis on gene sets. For example, given a set of genes that are up-regulated under certain conditions, an enrichment analysis will find which GO terms are over-represented (or under-represented) using annotations for that gene set and pathway enrichment can also be applied afterwards.

### 8.1 GO, KEGG and DO analyses

gdcEnrichAnalysis() can perform Gene ontology (GO), Kyoto Encyclopedia of Genes and Genomes (KEGG) and Disease Ontology (DO) functional enrichment analyses of a list of genes simultaneously. GO and KEGG analyses are based on the R/Bioconductor packages clusterProfilier(Yu et al. 2012) and DOSE(Yu et al. 2015). Redundant GO terms can be removed by specifying simplify=TRUE in the gdcEnrichAnalysis() function which uses the simplify() function in the clusterProfilier(Yu et al. 2012) package.

# Kaplan Meier plot



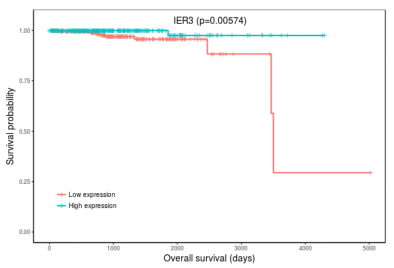


Figure 6:

```
enrichOutput <- gdcEnrichAnalysis(gene = rownames(deALL), simplify = TRUE)

## ### This step may take a few minutes ###

## Step 1/5: BP analysis done!

## Step 2/5: CC analysis done!

## Step 3/5: MF analysis done!

## Step 4/5: KEGG analysis done!

## Step 5/5: DO analysis done!

terms <- c()

for (category in c('GO_BP', 'GO_CC', 'GO_MF', 'KEGG', 'DO')) {
   terms <- c(terms, which(enrichOutput$Category==category)[1:3])
}

enrichOutput[terms,]</pre>
```

```
##
                                                                   Terms Counts
## 1
                                          GO:0006936~muscle contraction
                                                                             77
## 2
                           GO:2000027~regulation of organ morphogenesis
                                                                             47
                       GO:0051146~striated muscle cell differentiation
## 3
                                                                             56
## 63
                                        GO:0031012~extracellular matrix
                                                                             91
                                           GO:0043292~contractile fiber
                                                                             55
## 64
## 65
                                                  GO:0042383~sarcolemma
                                                                             38
## 79
                                   GO:0005539~glycosaminoglycan binding
                                                                             43
                                            GO:0015267~channel activity
                                                                             71
## 80
## 81
                 GO:0022803~passive transmembrane transporter activity
                                                                             71
## 91
                                  hsa05414~Dilated cardiomyopathy (DCM)
                                                                             25
## 92
                             hsa05410~Hypertrophic cardiomyopathy (HCM)
                                                                             22
## 93
      hsa05412~Arrhythmogenic right ventricular cardiomyopathy (ARVC)
                                                                             20
```

```
## 101
                                             DOID:10283~prostate cancer
                                                                             76
## 102
                              DOID:3856~male reproductive organ cancer
                                                                             76
## 103
                                                                             69
                                                      DOID: 423~myopathy
                                                  FDR foldEnrichment
##
                   BgRatio
       GeneRatio
                                 pValue
## 1
         77/1353 326/16447 9.881094e-18 4.882249e-14
                                                            2.871191
## 2
         47/1353 185/16447 1.527429e-12 2.515676e-09
                                                            3.088268
## 3
         56/1353 249/16447 2.588835e-12 2.741716e-09
                                                            2.733868
         91/1431 425/17563 5.476425e-18 2.650589e-15
## 63
                                                            2.627916
## 64
         55/1431 217/17563 1.472759e-14 2.376051e-12
                                                            3.110728
## 65
         38/1431 124/17563 3.068355e-13 3.320284e-11
                                                            3.761153
## 79
         43/1351 200/16514 3.287862e-09 2.689471e-06
                                                            2.628061
         71/1351 450/16514 5.409995e-08 1.475125e-05
## 80
                                                            1.928603
         71/1351 450/16514 5.409995e-08 1.475125e-05
## 81
                                                            1.928603
## 91
         25/607
                   89/7174 4.416347e-08 1.245410e-05
                                                            3.319882
## 92
          22/607
                   83/7174 8.615268e-07 1.139528e-04
                                                            3.132689
## 93
          20/607
                  72/7174 1.212263e-06 1.139528e-04
                                                            3.282995
## 101
         76/842 412/7577 3.902626e-06 2.993314e-03
                                                            1.659975
## 102
         76/842 422/7577 9.701641e-06 3.720579e-03
                                                            1.620639
## 103
          69/842 385/7577 2.987141e-05 5.727843e-03
                                                            1.612774
##
## 1
## 2
## 3
       ENSG00000164764/ENSG00000095713/ENSG00000197565/ENSG00000154736/ENSG00000183287/ENSG00000166833/
## 63
## 64
## 65
## 79
## 80
## 81
## 91
## 92
## 93
## 101
## 102
## 103
##
## 1
## 2
## 3
## 63
       SBSPON/CRTAC1/COL4A6/ADAMTS5/CCBE1/NAV2/MAMDC2/TGFBR3/DPT/GPLD1/FGFR2/NDP/FGF10/TINAGL1/CLU/COL1
## 65
## 79
## 80
## 81
## 91
## 92
## 93
## 101
## 102
## 103
##
       Category
## 1
          GO_BP
          GO BP
## 2
```

```
GO BP
## 3
## 63
           GO_CC
           GO CC
##
  64
           GO_CC
## 65
##
   79
           GO_MF
## 80
           GO MF
## 81
           GO_MF
            KEGG
## 91
## 92
            KEGG
## 93
            KEGG
## 101
              DO
## 102
              D0
## 103
              DO
```

### 8.2 Enrichment visualization

The output generated by gdcEnrichAnalysis() can be used for visualization in the gdcEnrichPlot() function by specifying type,category and numTerms arguments.

### 8.2.1 GO barplot

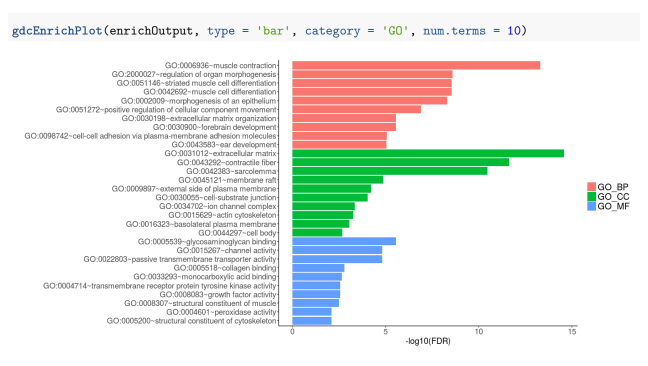


Figure 7:

### 8.2.2 GO bubble plot

```
gdcEnrichPlot(enrichOutput, type='bubble', category='GO', num.terms = 10)
```

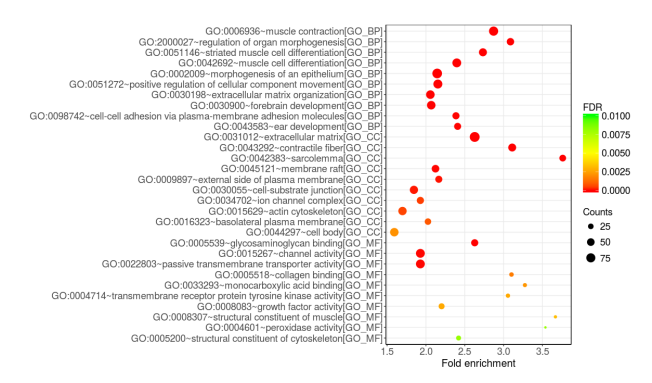


Figure 8:

### 8.2.3 KEGG/DO barplot

```
gdcEnrichPlot(enrichment = enrichOutput,
                                 = 'bar',
                   type
                                 = 'KEGG',
                   category
                                 = 'chocolate1',
                   bar.color
                  num.terms = 20)
                       hsa05414~Dilated cardiomyopathy (DCM)
                  hsa05410~Hypertrophic cardiomyopathy (HCM)
  hsa05412~Arrhythmogenic right ventricular cardiomyopathy (ARVC)
                            hsa04512~ECM-receptor interaction
                                    hsa04510~Focal adhesion
                                    hsa04360~Axon guidance
                   hsa04270~Vascular smooth muscle contraction-
                             hsa05205~Proteoglycans in cancer
                        hsa04022~cGMP-PKG signaling pathway
                             hsa00480~Glutathione metabolism
                                                                                    -log10(FDR)
```

Figure 9:

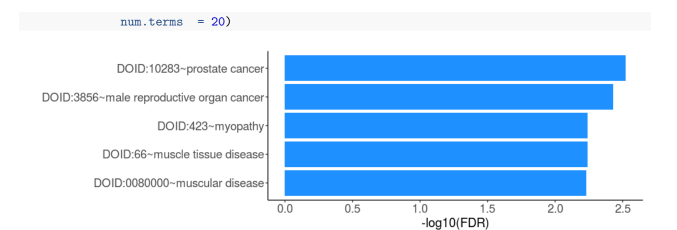


Figure 10:

### 8.2.4 KEGG/DO bubble plot

```
gdcEnrichPlot(=enrichOutput, category='KEGG',type = 'bubble', num.terms = 20)
```

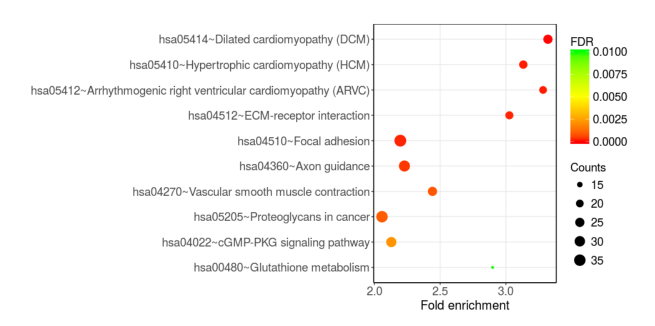


Figure 11:

gdcEnrichPlot(enrichOutput, category='DO', type = 'bubble', num.terms = 20)

#### 8.2.5 Pathview

Users can visualize a pathway map with pathview() function in the pathview(Luo and Brouwer 2013) package. It displays related many-genes-to-many-terms on 2-D view, shows by genes on BioCarta & KEGG pathway maps. Gradient colors can be used to indicate if genes are up-regulated or down-regulated.

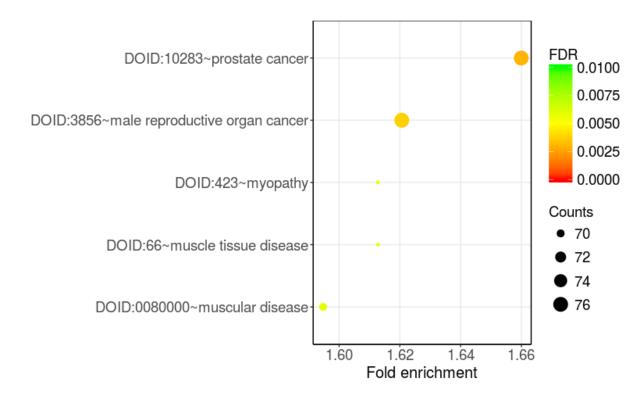


Figure 12:

### 8.2.6 View pathway maps on a local webpage by shinyPathview

shinyPathview() allows users view and download pathways of interests by simply selecting the pathway terms on a local webpage.

```
pathways <- as.character(enrichOutput$Terms[enrichOutput$Category=='KEGG'])
shinyPathview(deg, pathways = pathways, directory = 'pathview')</pre>
```

### 9 sessionInfo

```
sessionInfo()
```

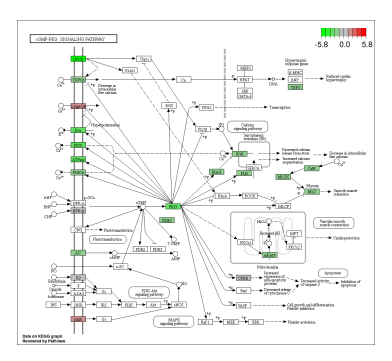


Figure 13:

### Pathview

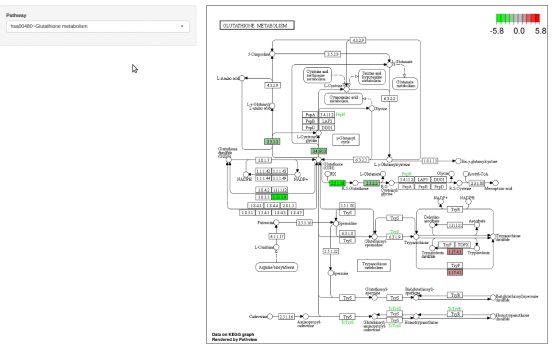


Figure 14:

```
## R version 3.3.1 (2016-06-21)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 16.04.1 LTS
##
## locale:
## [1] LC CTYPE=en US.UTF-8
                                   LC NUMERIC=C
## [3] LC TIME=en US.UTF-8
                                   LC COLLATE=en US.UTF-8
## [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
                                               datasets methods
                 graphics grDevices utils
                                                                   base
##
## other attached packages:
## [1] GDCRNATools_0.99.0
##
## loaded via a namespace (and not attached):
##
     [1] colorspace 1.3-2
                                    rjson 0.2.15
##
     [3] rprojroot_1.2
                                    qvalue_2.4.2
##
     [5] htmlTable 1.9
                                    XVector 0.12.1
##
     [7] GenomicRanges_1.24.3
                                    base64enc_0.1-3
                                    topGO 2.24.0
##
     [9] ggpubr_0.1.6
## [11] bit64_0.9-7
                                    AnnotationDbi 1.34.4
## [13] splines_3.3.1
                                    mnormt_1.5-5
## [15] GOSemSim_1.30.3
                                    geneplotter_1.50.0
## [17] knitr_1.17
                                    pathview_1.12.0
## [19] Formula_1.2-2
                                    jsonlite_1.5
## [21] km.ci_0.5-2
                                    broom_0.4.2
## [23] annotate_1.50.1
                                    cluster_2.0.6
## [25] GO.db_3.3.0
                                    png_0.1-7
##
  [27] graph_1.50.0
                                    shiny_1.0.5
## [29] httr_1.3.1
                                    backports_1.1.1
##
   [31] assertthat 0.2.0
                                    Matrix 1.2-11
## [33] lazyeval_0.2.1
                                    limma_3.28.21
## [35] acepack 1.4.1
                                    htmltools 0.3.6
## [37] tools_3.3.1
                                    bindrcpp_0.2
## [39] igraph_1.1.2
                                    gtable_0.2.0
## [41] glue_1.2.0
                                    reshape2_1.4.2
                                    dplyr 0.7.4
## [43] DO.db 2.9
## [45] Rcpp_0.12.13
                                    Biobase_2.32.0
## [47] Biostrings_2.40.2
                                    gdata_2.18.0
## [49] nlme_3.1-131
                                    psych_1.7.8
## [51] stringr_1.2.0
                                    mime_0.5
## [53] clusterProfiler_3.0.5
                                    gtools_3.5.0
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