04-Enrichment

R Markdown - Enrichment anlalysis using EnrichR R package

Summary

- I have already shortlisted the genes in my Tazeen_GroupCompAnalysis.Rmd file based on p-value cut
 off of 0.01
- I also have created a clean list of gene names in the same file above
- However, I will be redoing the filtering and gene name clean up to ensure I get the same results, and also to make sure the gene names are given to the EnrichR function in the correct/expected format
- I won't be writing additional files though for step 1 and 2 as the outputs are already there and I just need to check the content and formats
- Then I will connect to the EnrichR database and run Enrichment

Import T-test results so that we can short list further

Filtering/shortlisting Short list results based in p-value cut off (pvalue ≤ 0.01) This is already done but redoing just to be sure

```
pValueCutOff <- 0.01
which1 <- ((as.numeric(result1$Pvalue) <= pValueCutOff))
table(which1) #146 genes with p <= 0.01.

## which1
## TRUE
## 157

resultShort <- result1[which1, ] #short listed genes</pre>
```

Clean gene names

Note - the gene names are in the form of "reporterid_genename". So need to split this.

```
funcSplit <- function(rep_gene) {</pre>
    rep_gene_split <- unlist(strsplit(x = rep_gene,</pre>
                                        split = "|",
                                        fixed = TRUE))
    gene <- rep_gene_split[2]</pre>
    return(gene)
}
geneListSplit <- apply(X = as.matrix((resultShort$Feature)),</pre>
                        MARGIN = 1, FUN = funcSplit )
head(geneListSplit) #cleaned gene names
## [1] "TNKS"
                 "EXOC7"
                            "NDUFA10" "FAM91A1" "KYNU"
                                                             "SAMD15"
#remove duplicates
geneListSplit1 <- unique(geneListSplit)</pre>
# remove NA value
geneListFinal <- na.omit(geneListSplit1) #156 genes</pre>
head(geneListFinal)
                            "NDUFA10" "FAM91A1" "KYNU"
## [1] "TNKS"
                  "EXOC7"
                                                             "SAMD15"
#print number of unique genes
length(geneListFinal)
## [1] 156
```

Load Databases for Enrichr R package, and check connection

```
## 3
                          Pathways in cancer 2/531 0.0098313553
                                                                       0.055711013
## 4
                      VEGF signaling pathway 1/59 0.0175720140
                                                                       0.074681059
                                                                        0.076796241
## 5
                    Chronic myeloid leukemia
                                              1/76 0.0225871297
## 6
                   Th17 cell differentiation 1/107 0.0316774286
                                                                       0.079200112
##
    Old.P.value Old.Adjusted.P.value Odds.Ratio Combined.Score
                                    0 153.30000
                                                     1335.73937 SPI1; RUNX1
## 1
               0
## 2
               0
                                       52.11579
                                                      344.72067 SPI1:RUNX1
## 3
               0
                                                      85.03847 SPI1; RUNX1
                                    0
                                        18.39792
## 4
               0
                                    0
                                        68.74483
                                                      277.82863
                                                                        KDR
## 5
               0
                                    0
                                                                      RUNX1
                                        53.11733
                                                      201.33461
## 6
               0
                                        37.52453
                                                      129.54033
                                                                     RUNX1
#List of databases for which enrichment analysis will be run
dblist1 <- read.csv(file = "input/2023-EnrichR-Databases.txt",</pre>
                    header = F, stringsAsFactors = F)
head(dblist1)
```

```
## V1
## 1 KEGG_2021_Human
## 2 WikiPathway_2021_Human
## 3 GO_Biological_Process_2023
## 4 Reactome_2022
## 5 BioPlanet_2019
## 6 ClinVar_2019
```

Call function to run Enrichment

```
# set output file name
outputFileName1 <- paste("output/Tazeen_Moderate_Incipient", "_EnrichR.xlsx", sep="")
#Load R script into the environment
source(file = "functionEnrichment.R")
#call function to run Enrichment
functionEnrichment(dblist1, geneListFinal, outputFileName1)

## Uploading data to Enrichr... Done.
## Querying KEGG_2021_Human... Done.</pre>
```

```
## Parsing results... Done.
## Uploading data to Enrichr... Done.
## Querying WikiPathway_2021_Human... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
## Querying GO_Biological_Process_2023... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
## Querying Reactome_2022... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
## Querying BioPlanet_2019... Done.
```

```
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying ClinVar 2019... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying Transcription Factor PPIs... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
     Querying TRANSFAC_and_JASPAR_PWMs... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying TargetScan_microRNA... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying miRTarBase_2017... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying DisGeNET... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying OMIM_Disease... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying Jensen DISEASES... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying Chromosome_Location... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying VirusMINT... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying Virus-Host_PPI_P-HIPSTer_2020... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying HMDB_Metabolites... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
     Querying dbGap... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying MSigDB_Hallmark_2020... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying ProteomicsDB_2020... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying GWAS_Catalog_2023... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying InterPro_Domains_2019... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying CCLE Proteomics 2020... Done.
```

```
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying Proteomics_Drug_Atlas_2023... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying PheWeb_2019... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
     Querying Pfam_Domains_2019... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying ChEA_2022... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying SILAC_Phosphoproteomics... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
   Querying Azimuth_2023... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
    Querying MAGNET_2023... Done.
## Parsing results... Done.
## Uploading data to Enrichr... Done.
     Querying GeDiPNet_2023... Done.
## Parsing results... Done.
```

#NEED INTERNET CONNECTION

Note - you will need internet connection to complete the above step.