BB512/BB612 - Week VIII

OR.A

```
suppressPackageStartupMessages(library(enrichR))
selected_genes <- c("PPARA", "FABP4", "PLTP", "PPARG", "RXRA", "ATRX", "TP53", "KRAS", "MYC")
dbs <- listEnrichrDbs()</pre>
head(dbs)
##
     geneCoverage genesPerTerm
                                                      libraryName
## 1
            13362
                                              Genome_Browser_PWMs
                            275
## 2
            27884
                           1284
                                        TRANSFAC_and_JASPAR_PWMs
## 3
             6002
                             77
                                       Transcription_Factor_PPIs
                           1370
            47172
                                                        ChEA_2013
## 5
            47107
                            509 Drug_Perturbations_from_GEO_2014
##
            21493
                           3713
                                         ENCODE_TF_ChIP-seq_2014
##
                                                           link numTerms
  1 http://hgdownload.cse.ucsc.edu/goldenPath/hg18/database/
                                                                      615
                     http://jaspar.genereg.net/html/DOWNLOAD/
                                                                      326
##
## 3
                                                                      290
## 4
               http://amp.pharm.mssm.edu/lib/cheadownload.jsp
                                                                      353
## 5
                              http://www.ncbi.nlm.nih.gov/geo/
                                                                      701
## 6
                 http://genome.ucsc.edu/ENCODE/downloads.html
                                                                      498
##
                                       appyter categoryId
## 1 ea115789fcbf12797fd692cec6df0ab4dbc79c6a
## 2 7d42eb43a64a4e3b20d721fc7148f685b53b6b30
                                                         1
## 3 849f222220618e2599d925b6b51868cf1dab3763
                                                         1
## 4 7ebe772afb55b63b41b79dd8d06ea0fdd9fa2630
                                                         7
## 5 ad270a6876534b7cb063e004289dcd4d3164f342
                                                         7
## 6 497787ebc418d308045efb63b8586f10c526af51
                                                         7
dbs[grepl("KEGG", dbs$libraryName), ]
##
                                                                                 link
       geneCoverage genesPerTerm
                                      libraryName
## 11
               4128
                               48
                                        KEGG_2013 http://www.kegg.jp/kegg/download/
                                        KEGG_2015 http://www.kegg.jp/kegg/download/
## 60
               3800
                               48
## 94
               7010
                               87
                                        KEGG_2016 http://www.kegg.jp/kegg/download/
## 148
               7802
                               92 KEGG_2019_Human
                                                                https://www.kegg.jp/
## 149
               8551
                               98 KEGG 2019 Mouse
                                                                https://www.kegg.jp/
  173
               8078
                              102 KEGG_2021_Human
##
                                                                https://www.kegg.jp/
       numTerms
                                                   appyter categoryId
## 11
            200 eb26f55d3904cb0ea471998b6a932a9bf65d8e50
                                                                     7
## 60
            179 e80d25c56de53c704791ddfdc6ab5eec28ae7243
                                                                     7
            293 43f56da7540195ba3c94eb6e34c522a699b36da9
                                                                     7
## 94
            308 3477bc578c4ea5d851dcb934fe2a41e9fd789bb4
                                                                     7
## 148
                                                                     7
## 149
            303 187eb44b2d6fa154ebf628eba1f18537f64e797c
## 173
            320
```

dbs[grepl("GO", dbs\$libraryName),]

```
##
       geneCoverage genesPerTerm
                                                   libraryName
## 15
              12753
                                   GO_Molecular_Function_2015
##
  20
              13236
                                   GO_Cellular_Component_2015
## 21
              14264
                               58
                                   GO_Biological_Process_2015
##
  46
               7682
                               78
                                   GO_Biological_Process_2013
##
  47
                                   GO_Cellular_Component_2013
               7324
                              172
##
  48
               8469
                              122
                                   GO Molecular Function 2013
##
  112
              10427
                               38
                                   GO Cellular Component 2017
## 113
              10601
                               25
                                   GO Molecular Function 2017
## 114
              13822
                                   GO Biological Process 2017
                              143 GO_Cellular_Component_2017b
## 115
               8002
## 116
              10089
                               45 GO_Molecular_Function_2017b
                               49 GO_Biological_Process_2017b
## 117
              13247
  130
              14433
                                   GO_Biological_Process_2018
## 131
               8655
                               61
                                  GO_Cellular_Component_2018
                                   GO_Molecular_Function_2018
##
  132
              11459
                                   GO_Biological_Process_2021
##
  176
                               33
              14937
## 177
              11497
                                   GO_Cellular_Component_2021
## 178
                                   GO_Molecular_Function_2021
              11936
##
                                                               link numTerms
## 15
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                         1136
  20
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                          641
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                         5192
##
  21
##
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                          941
  46
##
  47
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                          205
       http://www.geneontology.org/GO.downloads.annotations.shtml
                                                                          402
##
  48
##
  112
                                      http://www.geneontology.org/
                                                                          636
## 113
                                      http://www.geneontology.org/
                                                                          972
## 114
                                      http://www.geneontology.org/
                                                                         3166
                                                                          816
## 115
                                      http://www.geneontology.org/
## 116
                                      http://www.geneontology.org/
                                                                         3271
## 117
                                      http://www.geneontology.org/
                                                                        10125
## 130
                                      http://www.geneontology.org/
                                                                         5103
## 131
                                      http://www.geneontology.org/
                                                                          446
## 132
                                      http://www.geneontology.org/
                                                                         1151
## 176
                                      http://www.geneontology.org/
                                                                         6036
## 177
                                      http://www.geneontology.org/
                                                                          511
##
  178
                                      http://www.geneontology.org/
                                                                         1274
                                         appyter categoryId
##
       f531ac2b6acdf7587a54b79b465a5f4aab8f00f9
  15
                                                           7
                                                           7
## 20
       e1d004d5797cbd2363ef54b1c3b361adb68795c6
       bf120b6e11242b1a64c80910d8e89f87e618e235
                                                           7
## 21
       5216d1ade194ffa5a6c00f105e2b1899f64f45fe
                                                           7
##
  46
##
       fd1332a42395e0bc1dba82868b39be7983a48cc5
                                                           7
       7e3e99e5aae02437f80b0697b197113ce3209ab0
                                                           7
                                                           7
  112 8fed21d22dfcc3015c05b31d942fdfc851cc8e04
  113 b4018906e0a8b4e81a1b1afc51e0a2e7655403eb
                                                           7
                                                           7
## 114 d9da4dba4a3eb84d4a28a3835c06dfbbe5811f92
                                                           7
## 115 ecf39c41fa5bc7deb625a2b5761a708676e9db7c
   116 8d8340361dd36a458f1f0a401f1a3141de1f3200
                                                           7
                                                           7
  117 6404c38bffc2b3732de4e3fbe417b5043009fe34
## 130 b5b77681c46ac58cd050e60bcd4ad5041a9ab0a9
```

```
7
## 131 e9ebe46188efacbe1056d82987ff1c70218fa7ae
                                                            7
## 132 79ff80ae9a69dd00796e52569e41422466fa0bee
## 176
                                                            3
## 177
                                                            3
## 178
                                                            3
res <- enrichr(selected_genes, databases = c("KEGG_2021_Human", "GO_Biological_Process_2017"))
## Uploading data to Enrichr... Done.
     Querying KEGG_2021_Human... Done.
     Querying GO Biological Process 2017... Done.
## Parsing results... Done.
head(res$KEGG_2021_Human)
##
                                    Term Overlap
                                                    P.value Adjusted.P.value
## 1
                         Thyroid cancer
                                            5/37 2.0490e-12
                                                                   2.2948e-10
## 2
                PPAR signaling pathway
                                            5/74 7.5260e-11
                                                                   4.2146e-09
## 3
                                           5/157 3.4347e-09
                                                                   1.2823e-07
                            Hepatitis C
## 4 Thyroid hormone signaling pathway
                                           4/121 1.5687e-07
                                                                   4.3924e-06
                         Gastric cancer
                                           4/149 3.6210e-07
## 5
                                                                   8.1110e-06
## 6
                         Bladder cancer
                                            3/41 6.6589e-07
                                                                   1.2430e-05
     Old.P.value Old.Adjusted.P.value Odds.Ratio Combined.Score
## 1
               0
                                            779.65
                                                           20983.2
               0
                                      0
                                            360.91
                                                            8412.7
## 2
## 3
               0
                                      0
                                            163.15
                                                            3179.7
               0
## 4
                                      0
                                            135.89
                                                            2129.1
## 5
               0
                                      0
                                            109.50
                                                            1624.0
               0
## 6
                                      0
                                            262.54
                                                            3733.9
##
                            Genes
## 1
        RXRA; MYC; PPARG; KRAS; TP53
## 2 FABP4; RXRA; PPARG; PPARA; PLTP
## 3
        RXRA; MYC; KRAS; PPARA; TP53
## 4
              RXRA; MYC; KRAS; TP53
## 5
              RXRA; MYC; KRAS; TP53
## 6
                    MYC; KRAS; TP53
head(res$GO_Biological_Process_2017)
##
                                                                                       Term
## 1 positive regulation of transcription from RNA polymerase II promoter (GO:0045944)
## 2 negative regulation of transcription from RNA polymerase II promoter (GO:0000122)
## 3
                       negative regulation of sequestering of triglyceride (GO:0010891)
## 4
                                negative regulation of cholesterol storage (GO:0010887)
## 5
             peroxisome proliferator activated receptor signaling pathway (GO:0035357)
## 6
                                                response to gamma radiation (GO:0010332)
##
                P. value Adjusted. P. value Old. P. value Old. Adjusted. P. value
     Overlap
## 1
       6/712 1.5278e-07
                               2.3070e-05
                                                      0
## 2
       5/473 8.4392e-07
                               6.3716e-05
                                                      0
                                                                            0
## 3
         2/5 1.7987e-06
                                                      0
                                                                            0
                               9.0534e-05
         2/6 2.6974e-06
                                                      0
                                                                            0
## 4
                               1.0183e-04
## 5
         2/7 3.7755e-06
                                                      0
                                                                            0
                               1.1402e-04
         2/9 6.4693e-06
                               1.6281e-04
##
     Odds.Ratio Combined.Score
                                                           Genes
## 1
         54.632
                         857.41 RXRA; MYC; ATRX; PPARG; PPARA; TP53
## 2
         52.145
                         729.26
                                      RXRA; MYC; PPARG; PPARA; TP53
```

##	3	1903.619	25181.94	PPARG; PPARA
##	4	1427.643	18306.98	PPARG; PPARA
##	5	1142.057	14260.84	RXRA; PPARG
##	6	815.673	9746.03	MYC; TP53

plotEnrich(res\$KEGG_2021_Human)

Enrichment analysis by Enrichr Thyroid cancer PPAR signaling pathway Hepatitis C Thyroid hormone signaling pathway Gastric cancer Bladder cancer Transcriptional misregulation in cancer P value **Enriched terms** Pathways in cancer Lipid and atherosclerosis 1e-05 Endometrial cancer Chemical carcinogenesis 2e-05 Central carbon metabolism in cancer Non-small cell lung cancer 3e-05 Chronic myeloid leukemia

2

Gene count

3

plotEnrich(res\$GO_Biological_Process_2017)

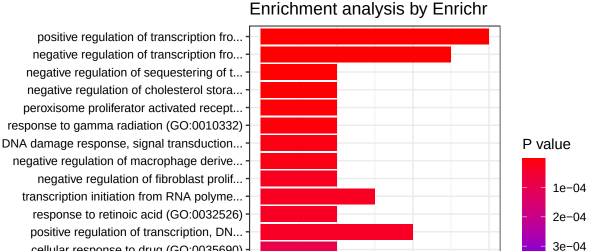
Non-alcoholic fatty liver disease

Longevity regulating pathway PI3K-Akt signaling pathway

Colorectal cancer Small cell lung cancer

Breast cancer

ò



Gene count

positive regulation of transcription, DN... cellular response to drug (GO:0035690) positive regulation of gene expression (... cellular response to UV (GO:0034644) regulation of circadian rhythm (GO:00427... protein homotetramerization (GO:0051289) cellular response to hypoxia (GO:0071456... Ras protein signal transduction (GO:0007... chromatin remodeling (GO:0006338)

Enriched terms

FCS

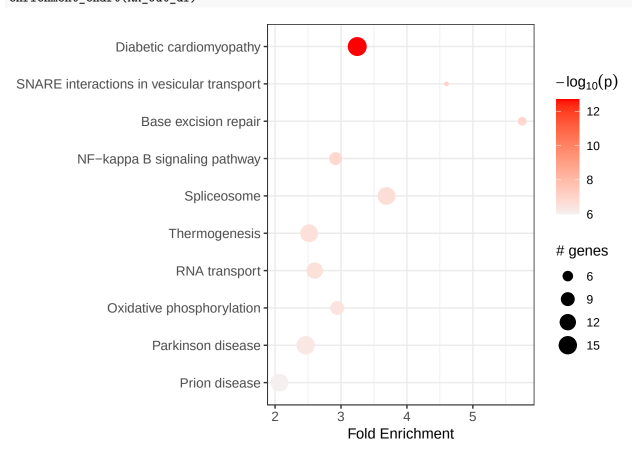


0

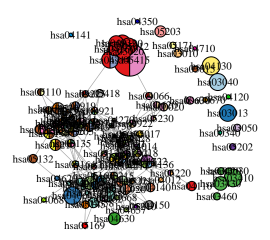
```
## $\`1221633 Meiotic Synapsis\`
    [1] "12189"
                      "13006"
                                  "15077"
                                               "15078"
                                                             "15270"
                                                                          "15512"
    [7] "16905"
                                                             "20843"
                      "16906"
                                  "19357"
                                               "20842"
                                                                          "20957"
##
##
   [13]
        "20962"
                      "21749"
                                  "21750"
                                               "22196"
                                                             "23856"
                                                                          "24061"
        "28113"
                      "50878"
                                  "56739"
                                               "57321"
                                                             "64009"
                                                                          "66654"
##
   [19]
   [25]
        "69386"
                      "71846"
                                  "74075"
                                               "77053"
                                                             "94244"
                                                                          "97114"
##
        "97122"
                      "97908"
                                               "140557"
                                                             "223697"
                                                                          "260423"
##
   [31]
                                  "101185"
##
   [37]
        "319148"
                      "319149"
                                  "319150"
                                               "319151"
                                                             "319152"
                                                                          "319153"
                                  "319156"
                                               "319157"
                                                                          "319159"
##
   [43]
        "319154"
                      "319155"
                                                             "319158"
   [49]
        "319160"
                      "319161"
                                  "319565"
                                                "320332"
                                                             "320558"
                                                                          "326619"
##
   [55] "326620"
                      "360198"
                                   "497652"
                                                "544973"
                                                             "625328"
                                                                          "667250"
##
##
   [61] "100041230" "102641229" "102641751" "102642045"
##
## $\`1368092_Rora_activates_gene_expression\`
                 "12753" "12894" "18143"
  [1] "11865"
                                              "19017"
                                                        "19883"
                                                                  "20787"
                                                                            "217166"
## [9] "328572"
```

```
head(exampleRanks, 2)
## 170942 109711
## -63.337 -49.748
fgseaRes <- fgsea(pathways = examplePathways,
                  stats = exampleRanks,
                  minSize = 15,
                  maxSize = 500)
## Warning in fgseaMultilevel(...): For some pathways, in reality P-values are less
## than 1e-10. You can set the `eps` argument to zero for better estimation.
head(fgseaRes[order(pval), ])
##
                          pathway pval
                                              padj log2err
                                                                 ES
                                                                       NES size
## 1: 5990979_Cell_Cycle,_Mitotic 1e-10 3.9067e-09
                                                        NA 0.55948 2.7691
                                                                            317
               5990980 Cell Cycle 1e-10 3.9067e-09
                                                        NA 0.53885 2.7059
                                                                            369
## 3:
          5990981_DNA_Replication 1e-10 3.9067e-09
                                                        NA 0.64400 2.6394
                                                                             82
## 4:
         5990987_Synthesis_of_DNA 1e-10 3.9067e-09
                                                        NA 0.64786 2.6429
                                                                             78
                                                                             98
## 5:
                  5990988_S_Phase 1e-10 3.9067e-09
                                                        NA 0.60131 2.5296
## 6:
          5990990_G1_S_Transition 1e-10 3.9067e-09
                                                        NA 0.62329 2.5738
##
                                   leadingEdge
## 1: 66336,66977,12442,107995,66442,12571,...
## 2: 66336,66977,12442,107995,66442,19361,...
## 3: 57441,17219,69270,12575,69263,17215,...
## 4: 17219,69270,12575,69263,17215,68240,...
## 5: 67849,17219,69270,12575,69263,71988,...
## 6: 20135,13555,17219,12575,12448,17215,...
\mathbf{PT}
suppressPackageStartupMessages(library(pathfindR))
head(RA_input)
##
     Gene.symbol
                    logFC adj.P.Val
## 1
         FAM110A -0.69394 3.4087e-06
## 2
         RNASE2 1.35350 1.0085e-05
## 3
          S100A8 1.54483 3.4664e-05
          S100A9 1.02809 2.2626e-04
## 4
## 5
          TEX261 -0.32360 2.2626e-04
        ARHGAP17 -0.69193 2.7081e-04
# RA_out_df <- run_pathfindR(RA_input, output_dir = tmp_dir)
RA_out_df <- RA_output</pre>
head(RA_out_df)
##
                                       Term_Description Fold_Enrichment occurrence
## 1 hsa05415
                                Diabetic cardiomyopathy
                                                                  3.2464
                                                                                 10
## 2 hsa04130 SNARE interactions in vesicular transport
                                                                  4.5990
                                                                                 10
## 3 hsa03410
                                   Base excision repair
                                                                  5.7488
                                                                                  1
## 4 hsa04064
                           NF-kappa B signaling pathway
                                                                  2.9186
                                                                                 10
## 5 hsa03040
                                            Spliceosome
                                                                  3.6888
                                                                                 10
## 6 hsa04714
                                          Thermogenesis
                                                                  2.5175
                                                                                 10
##
       support
                 lowest_p highest_p
```

```
## 1 0.0752907 1.9512e-13 1.9512e-13
## 2 0.0118360 8.0071e-08 8.0071e-08
## 3 0.0053191 1.2365e-07 1.2365e-07
## 4 0.0502592 1.2586e-07 1.2586e-07
## 5 0.0477457 2.2308e-07 4.4562e-05
## 6 0.0441982 2.5435e-07 2.5435e-07
                                                         Up regulated
## 1 NCF4, MMP9, NDUFA1, NDUFB3, UQCRQ, COX6A1, COX7A2, COX7C, GAPDH
## 2
## 3
                                                                POLE4
## 4
                                                                 LY96
                                                   SF3B6, LSM3, BUD31
## 5
                        NDUFA1, NDUFB3, UQCRQ, COX6A1, COX7A2, COX7C
## 6
                                                                      Down_regulated
##
## 1
                                   ATP2A2, MTOR, PDHA1, PDHB, VDAC1, SLC25A5, PARP1
## 2
                                                                 STX2, BET1L, SNAP23
## 3
                                                          MUTYH, APEX2, POLD2, PARP1
                                PRKCQ, CARD11, TICAM1, IKBKB, PARP1, UBE2I, CSNK2A2
## 4
## 5 SNRPB, SF3B2, U2AF2, PUF60, SNU13, DDX23, EIF4A3, HNRNPA1, PCBP1, SRSF8, SRSF5
                            ADCY7, CREB1, KDM1A, SMARCA4, ACTG1, ACTB, ARID1A, MTOR
enrichment_chart(RA_out_df)
```



RA_clu_df <- cluster_enriched_terms(RA_out_df, method = "fuzzy")</pre>



enrichment_chart(RA_clu_df, plot_by_cluster = TRUE, top_terms = 3)

